A Physicochemical Approach to Design Bioactive Scaffolds for Tissue Engineering

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A Physicochemical Approach to Design Bioactive Scaffolds for Tissue Engineering

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General Introduction and Aims

Chapter 1

General Introduction and Aims
Abstract

Tissue engineering aims at developing biological substitutes to restore maintain or improve tissue function. Scaffold-based tissue engineering provides an alternative approach for tissue regeneration. This approach involves the combination of cells and/or biomolecules with scaffolds, which serve as a temporary substrate for cells adhesion and a physical support to guide new tissue formation. Nanofibrous scaffolds mimicking the structural features of natural extracellular matrix (ECM) can be appealing scaffold candidates for tissue engineering as they provide similar physical cues to the native environment of the targeted tissue to regenerate. This chapter discusses different strategies to fabricate nanofibrous scaffolds for tissue engineering. We first describe three major methods for nanofibrous scaffolds fabrication: molecular self-assembly, phase separation, and electrospinning. Then, approaches for surface modification of nanofibrous scaffolds including blending and coating, plasma treatment, wet chemical methods, and surface graft polymerization are presented. Finally, applications of nanofibrous scaffolds in tissue engineering are introduced and the synopsis of this thesis described.

Key words:

Electrospinning, self-assembly, phase separation, nanofibrous scaffolds, tissue engineering
1.1. Introduction

Tissue engineering aims at developing biological replacements through the integration of life science and engineering principles. In general, tissue engineering can be subdivided into three major approaches [1]: (1) injection or transplantation of isolated cells to a defect site or an injured tissue, (2) delivery of tissue-inducing biomolecules (e.g., growth factors, peptides, polysaccharides) to a targeted tissue, (3) growth and differentiation of specific cell types in three-dimensional (3D) scaffolds. Among these approaches, scaffold-based tissue engineering has become the most popular, due to the potential of current scaffold fabrication technologies to incorporate chemical, physical, and biological stimuli at different scales able to direct cell activity.

The extracellular matrix (ECM) usually provides structural support to the cells in addition to performing various other important functions. Many extracellular proteins have a fibrous structure with a diameter at the nanometer or sub-micrometer scales. For example, collagen, the most abundant ECM protein in our body, has a fibrous structure with a fiber diameter that ranges from 50 to 500 nm. For successful and functional engineering of tissues and organs in scaffold-based approaches, an artificial scaffold should mimic the structure and biological function of native ECM as much as possible both in terms of physical cues and chemical composition. Therefore, researchers have put much effort to develop nanofibrous scaffolds that mimic the fibrillar structure of native ECM. Currently, there are three basic techniques available for generating nanofibrous scaffolds: molecular self-assembly, phase separation, and electrospinning (Table 1). Although these techniques allow developing nanofibrous scaffolds that mimic ECM physical structure with a number of degradable and non-degradable synthetic polymers, additional modifications are desirable to present biological cues, which exhibit a positive effect on cell adhesion, proliferation, and differentiation. Furthermore, recent studies have shown that submicron and nanoscale topographies can also modulate cell activity, thus acting as “synthetic” biological cues [2]. Surface modifications of nanofibrous scaffolds include blending and coating, plasma treatment, wet chemical methods, and surface graft polymerization.
In this chapter, we illustrate nanofibrous scaffolds for tissue engineering. Processing technologies to achieve structural features resembling the ECM and approaches to improve cell-scaffold interactions will be introduced before discussing nanofibrous scaffolds for tissue engineering applications.
1.2. Methods for nanofibrous scaffolds fabrication

1.2.1 Molecular self-assembly

Molecular self-assembly has emerged as a useful approach for manufacturing scaffolds for tissue engineering [3]. Unlike electrospinning, self-assembly can be defined as a spontaneous process to form ordered and stable structure through a number of non-covalent interactions, such as hydrophilic, electrostatic, and van der Waals interactions [4, 5]. The structural features of self-assembled materials can be tuned by controlling the kinetics, molecular chemistry, and assembling environment (pH, solvent, light, salt addition and temperature). The key challenge in self-assembly is to design molecular building blocks that can undergo spontaneous organization into a well-defined pattern that mimic the structural features of biological systems [6]. Small building blocks including small molecules, nucleic acids, and peptides, can self-assemble into nanofibrous structures. Among these building blocks, peptide-amphipile (PA) units, which combine the functions of peptides with the characteristics of surfactants, have gained a lot of attention due to the versatility in their design for biological applications. The chemical structure of a representative PA molecule consists of four key structure features: a hydrophobic domain consisting of a long alkyl tail, a short peptide sequence capable of forming intermolecular hydrogen bond, charged groups for enhanced solubility in water, and a bioactive epitope. Self-assembly of PA in water is governed by at least three major forces: hydrophobic interaction of alkyl tails, electrostatic repulsions between charged groups, and hydrogen bonding among peptide segments. The final structure of assembled PA reflects a balance of each force contribution [7].

The structural characteristics of nanofibers and nanofibrillar systems resulting from molecular self-assembly can be tailored by controlling processing parameters. Bundles of aligned PA nanofibers were obtained, for example, by self-assembling PA within parallel channels [8]. By introducing hydrophilic amino acids in peptide segments, specific sequences of PA could self-assemble into nanobelts with fairly monodisperse widths on the order of 150 nm and lengths of up to 0.1 mm that can be used as cell culture systems and drug delivery carriers [9]. Nanofibrous scaffolds were also fabricated from self-assembly of β-sheet peptides containing phenylalanine for controlled release [9, 10]. These studies proposed that the position of aromatic moieties is a significant determinant for supramolecular self-assembling conformations. Self-assembled chitin fibers with diameters of 3 nm and 10 nm could be also obtained from dissolving in hexafluoro-2-propanol (HFIP) and LiCl/N,N-dimethylacetamide (DMAC) respectively [11]. These
observations demonstrated that nanofiber assembly occurs at a critical concentration and fiber morphology can be controlled by solution concentration and solution evaporation rate. Collagen-like nanofibers were self-assembled from collagen mimetic peptides by controlling triple-helical nucleation [12]. Young-Hag et al. fabricated macro/nano-porous collagen scaffolds comprising nanofibrous collagen filaments through 3D plotting in combination with self-assembly of collagen molecules [13]. Nanofibrous hydrogels, which were self-assembled from lysine-containing hexapeptides, had a stiffness of up to 40 kPa and were biocompatible [14]. In addition, this self-assembling hydrogel could serve as an ideal bioink for bioprinting.

Molecular self-assembly is a fairly new technique to develop nanofibrous scaffolds which have been studied for a variety of tissue engineering applications including nerve [15], bone [16], and cartilage [17] regeneration. However, several technical hurdles still need to be addressed: First, it has not been demonstrated how to control the pore size and pore structure, which are important to allow for cell proliferation and migration. Secondly, their degradation profiles have not been systematically addressed [18]. Furthermore, most of self-assembled scaffolds are mechanically weak and do not effectively sustain and transfer mechanical loadings to the cells and surrounding tissues.

1.2.2. Thermally induced phase separation

Another interesting method used for manufacturing nanofibrous scaffolds is phase separation, which is a thermodynamic process involving the separation of phases due to physical incompatibility. Specifically, a homogeneous polymer solution becomes thermodynamically unstable under certain temperature conditions, and will form a polymer-rich phase and a polymer-poor phase. When the solvent is removed, the polymer-rich phase will solidify to a 3D structure while the polymer-poor phase will become the void space. The process of developing nanofibrous scaffolds from phase separation typically consists of 5 major steps: (i) raw material dissolution, (ii) gelation, (iii) solvent extraction, (iv) freezing and (v) drying [19]. The pore morphology of nanofibrous scaffolds can be tuned by varying processing parameters including polymer concentration, phase separation temperature and solvent/non-solvent exchange [20]. Furthermore, thermally induced phase separation can be combined with other processing techniques (e.g. particulate leaching and solid free-form fabrication) to generate scaffolds with complex porous structure and well-defined pore morphology [21]. For example, sodium chloride particles with diameters of 200-450 µm were mixed with a warm poly (lactic-co-glycolic acid)/tetrahydrofuran (PLGA/THF) solution and then cooled to a preset gelation
temperature. The formed composite gels were extracted with cold ethanol and washed with distilled water to remove the solvent and leach the salt particles. The samples were freeze-dried, resulting in a nanofibrous scaffold with a macroporous structure left behind from the leached salt [22]. Nanofibrous poly(l-lactic acid) (PLLA) scaffolds have been developed by using phase separation to improve cell seeding, distribution and mass transport [23]. When compared with solid pore-walled PLLA scaffolds, nanofibrous scaffolds allowed approximately a 2-fold increase in adhesion of osteoblast cells. More recently, novel injectable functional nanofibrous hollow microspheres (FNF-HMS) were fabricated using phase separation [24]. Transforming growth factor-β1 mimicking peptide cytomodulin, and bone morphogenetic protein-2 mimicking peptide P24 were conjugated onto FNF-HMS separately. In vitro and in vivo studies demonstrated that the injectable cell carriers successfully induced rabbit bone marrow-derived mesenchymal stem cells differentiation toward chondrocytes and osteoblasts.

Thermally induced phase separation is a promising technique to develop nanofibrous scaffold with well-defined pore shape and pore size. Although this technique can be combined with other fabrication methods to control the final 3D structure, the drawbacks of this technique, such as little control over fiber orientation and diameter, long fabrication time, and lack of mechanical properties need to be addressed to further achieve a fine control of the resulting scaffolds at the macro, micro, and nano scales.

1.2.3. Electrospinning

Electrospinning is a versatile and well-established process capable of producing fibers with diameters down to the sub-micron or nanometer range [25]. A basic electrospinning setup (Figure 1A) includes a high voltage electric source with positive or negative polarity; a syringe pump with capillaries or tubes to carry the solution (or melt) from a syringe to the spinneret and a conducting collector [26]. Under the electric field, the pulling force overcomes the surface tension of the polymer solution (or melt) and create a charged jet that travels through the atmosphere allowing the solvent to evaporate, thus leading to the deposition of solid polymer fibers on the collector [26, 27]. The fiber formation and structure is affected by three general types of variable: solution properties (concentration, viscosity, conductivity, and surface tension), processing factors (applied potential, collection distance, emitting electrode polarity, and feed rate), and environmental parameters (temperature, relative humidity, and velocity of the surrounding air in the spinning chamber) [26, 28].
Figure 1. (A) Typical electrospinning set-up using a grounded static collector; (B) Configuration of the coaxial electrospinning setup used for preparing core-shell structured nanofibers.

For many applications, it is necessary to fabricate a scaffold made of aligned nanofibers, as the anisotropy in topography and structure can have a great impact on mechanical properties and cell behavior. Currently, a number of collecting devices have been developed to attain aligned electrospun fibers. These collecting devices can be divided into three main categories according to the type of forces involved [29]: (1) mechanical forces by using of a rotating mandrel; (2) electrostatic forces by using parallel electrodes; (3) magnetic forces by using parallel permanent magnets. Furthermore, electrospinning can be used for fabricating scaffolds with complex architectures such as stacked arrays and tubular conduits. Stacked arrays of nanofiber scaffolds can be achieved by multilayering electrospinning. Aligned nanofibers were stacked into multilayered films with a controllable hierarchical porous structure by depositing fibers on an insulating substrate (e.g. quartz) onto which an array of electrodes have been patterned [30]. Tubular conduits made of electrospun fibers can be fabricated by depositing fibers over a small diameter rod as a collector. These electrospun conduits are usually applied in vascular or neural tissue engineering since they mimic the structure of these tissues. For instance, electrospun fiber conduits were manufactured with a length of 12 cm and a thickness of 1 mm via electrospinning a mixture of collagen type I, elastin, and poly(D,L lactide-co-glycolide) on a rotating rod of diameter 4.75 mm [31]. Compliance tests demonstrated that the electrospun fiber conduit possesses a diameter change of
12~14% within the physiologic pressure range, which exhibited a compliance behavior similar to that of a native vessel with a diameter change of 9%. Burst pressure testing showed that the burst pressure for the electrospun fiber conduit was 1425 mmHg or nearly 12 times that of normal systolic pressure.

In order to tailor the structures of resultant fibers, the modification of electrospinning set-ups have been carried out not only on the fiber collector but also on the spinneret. Coaxial spinnerets have been designed by many researchers for various aims in electrospinning (Figure 1.1B). Using this technology, core-shell nanofibers can be achieved. By using a spinneret with double coaxial capillaries, two components can be fed through different coaxial capillary channels to generate composite nanofibers in the form of core-sheath structure [32-34]. Some difficult-to-process polymer solutions can be also electrospun to form an ultrafine core within the shell of spinnable polymers. Thus, when the polymer sheath is removed, the desired polymer nanofiber core is retained [35]. Using a similar concept, the core solution can be removed instead of the polymer sheath, resulting in hollow nanofibers. For example, an ethanol solution containing poly (vinyl pyrrolidone) (PVP) and tetraisopropyl titanate (Ti (OiPr)₄) was used as sheath solution, while mineral oil was used as core solution. After simultaneous ejection through the inner and outer capillaries, hollow nanofibers were formed through removal of the inner core mineral oil phase [36]. Shell-core and hollow nanofibers have shown their potential in tissue engineering and drug delivery due to their unique architecture and properties. Biologically active agents can be encapsulated inside a polymer shell to form a reservoir-type drug delivery device. Therefore, the polymer shell would offer a temporal protection for the bioactive agents and control their sustain release [32]. Tissue regeneration processes can be also facilitated via introduction of biomolecules (e.g. growth factors) into the core of shell-core nanofibrous scaffolds and, eventually, the release of multiple factors can be envisioned by adding different compounds in the shell and in the core fibers.

Electrospun fibers with high surface to volume ratio and structures resembling extracellular matrix (ECM) have shown great potential applications in tissue engineering and drug delivery [37]. Yet, a fine control over nanofiber population distribution is usually lacking and is so far a limit in obtaining standardized scaffolds. Recently, melt electrospinning writing has emerged as a new technology to overcome this problem. Dalton and his colleagues accurately deposited melt electrospun poly(ε-caprolactone) (PCL) fibers to create complex structures having filament diameters typically in the range of 5 to 30 µm [38]. The melt electrospun fibers could consistently be stacked in a manner
like melt-extrusion based additive manufacturing techniques. More recently, through optimization of spinning parameters (e.g. voltage, flow rate, and spinneret diameters), PCL scaffolds with submicron fiber diameter (817±165 nm) were produced using melt electrospinning [39].

1.3. Surface modification of nanofibrous scaffolds

Surfaces play a vital role in biology and medicine, as most biological reactions take place at the interface between biological systems [40]. In tissue engineering, the chemical and physical characteristics of the biomaterial surface strongly impact on cell behavior, such as migration, attachment, and proliferation [41]. Although various degradable and non-degradable synthetic polymers have been used as tissue engineering scaffolding materials, a shortcoming of these materials is their lack of biological recognition [6]. Currently, several techniques have been developed to modify the scaffold surface including physical coating and blending, plasma treatment, graft polymerization and wet chemical methods (Figure 2).
Figure 2. Surface modification techniques of nanofibrous scaffolds. (A) Blending and coating; (B) Plasma or wet chemical treatment; (C) Surface graft polymerization.

1.3.1. Physical Coating or Blending

Perhaps the most straightforward and convenient mean of modifying a polymer surface is blending functional molecules and active agents into the bulk polymer or just coating it on the polymer surface. Usually, two or more materials are physically blended together to attain a new biocomposite with superior surface and/or mechanical properties. PLLA nanofibers with hydroxyapatite (HA) particles exposed on their surface were achieved by electrospinning different blended solutions [42, 43]. These composite fibers not only promoted osteoblast adhesion and growth, but also exhibited superior tensile properties as compared to the pure PLLA fibers due to the internal ionic bonding between calcium
ions in HA particles and ester groups in PLLA [44]. Jun et al. [45] developed electrically conductive blends of Poly (L-lactide-co-caprolactone) (PLCL) and polyaniline (PANI) submicron fibers and investigated their influence on the induction of myoblasts into myotubes. Incorporation of PANi into PLCL fibers significantly increased the conductivity. In addition, the tensile strength and elongation at break of PLCL fibers increased as the concentration of PANi in the fibers decreased, while the Young's modulus exhibited an opposite trend. Silk fibroin/keratin films demonstrated enhanced mechanical properties after blending with gelatin. When further mixed with calcium peroxide, films sustained the release of high level of oxygen over two weeks *in vitro*, and promoted cellular growth [46]. Taken together, these examples show that blending modifies not only the surface properties of substrate but also their bulk properties.

Coating is another physical approach of functionalizing nanofibrous scaffolds. Although this technique is limited in terms of controllability and coatings maybe prone to detachment from scaffold, it remains one of the easiest and most convenient ways to functionalize biomaterials' surface. In general, a material containing desired functional properties is used to coat nanofibrous scaffolds, aiming at improving their surface biocompatibility and enhancing cell-scaffold interactions. In recent years, coating of functional molecules onto polymers was significantly developed by combining new techniques including coaxial electrospinning and layer-by-layer assembly (LbL). Collagen-coated PCL nanofibers were developed by a coaxial electrospinning technique, with collagen and PCL as inner and outer solutions respectively, and cultured with human dermal fibroblasts [47]. The results demonstrated that collagen-coated PCL nanofibers from coaxial electropinning showed higher efficiency in favoring cell proliferation compared to collagen-coated PCL nanofibers prepared by soaking the PCL matrix in a 10 mg/mL collagen solution. The mechanism behind LbL coatings is that positively and negatively charged macromolecules are alternatively introduced onto the substrate surface via strong electrostatic interaction [19]. An environmentally benign surface modification process for poly(ethylene terephthalate) (PET) films was demonstrated by fabricating composite coatings through layer-by-layer assembly with cellulose and chitin nanofibrils [48]. The modified PET films exhibit high light transparency, flexibility, surface-hydrophilicity, and specific nanoporous structures. β-tricalcium phosphate coating on poly(propylene fumarate) scaffolds showed advantage in terms of osteoconductivity and osteointegration of scaffolds [49]. Aligned and aminolyzed PLA nanofibrous scaffolds, which were coated with graphene oxide, promoted rat Schwann cells growth and
regulated their orientation[50]. Moreover, this kind of scaffolds induced rat pheochromocytoma 12 cells differentiation and neurite growth in the presence of nerve growth factor. Physical coating could also be used to modulate the scaffolds properties including mechanical property, degradation rate and surface topography. The mechanical strength, degradation rate, and cell behavior of mesoporous bioactive glass could be modulated through controlling the amount of poly(glycerolsebacate) (PGS) coating [51]. Biphasic calcium phosphate scaffolds were coated with PLLA/HA nanocomposites for bone regeneration. Such PLLA/HA coatings improved the compressive strength of scaffolds from 0.31 to 3.56 MPa [52].

1.3.2. Plasma-based surface modification

Plasma, also named the forth state of matter, is a gaseous mixture of particles containing charged particles, excited and neutral atoms and molecules, radicals and UV photons [53]. Generally, plasma can be subdivided into two categories according to the temperature of particles: thermal plasma and non-thermal plasma. Non-thermal plasma, which is composed of low temperature particles has been commonly employed to modify the surface of heat-sensitive materials such as biodegradable polymers [54]. By using this technique, diverse functional groups can be effectively incorporated on the target surface of biodegradable polyesters to improve surface adhesion and wetting properties. For example, typical plasma treatment with different gaseous atmospheres such as oxygen, air, and ammonia can introduce carboxyl groups or amine groups to the target surface [55].

After introduction of specific functional groups on the surface of substrates by plasma treatment, synthetic and natural macromolecules could further immobilize monomers (etc. extracellular matrix protein components) on their surface to enhance cellular adhesion and proliferation. This process is called plasma grafting. Nanofibrous scaffolds composed of PLLA and PLGA were fabricated by electrospinning [56]. Their surfaces were treated with oxygen plasma treatment and grafted with hydrophilic acrylic acid. Such surface-modified scaffolds were shown to improve fibroblast attachment and proliferation in vitro. Plasma polymerization, another plasma-based surface modification approach, is distinct from plasma grafting, since it coats the substrate instead of covalent binding species to a plasma-treated polymer surface [54, 57]. Under plasma condition, gaseous or liquid monomers are converted into reactive fragments, which can in turn recombine to a polymer film deposited onto the substrate exposed to the plasma. To gain more insights on plasma polymerization processes, various -C:H:N thin films were deposited on glass
under different NH$_3$/C$_2$H$_4$ gas ratios, power inputs $W$, and gas flow rates $F$ [58]. The results demonstrated that the film growth is determined by the ratio between the reaction parameter $W$ and $F$, and the energy dissipated at its surface during the deposition given by ion flux times mean ion energy per deposition rate. After treatment with oxygen plasma, PLLA electrospun meshes had an increase in surface roughness as well as surface hydrophilicity [59]. PCL electrospun fibrous scaffolds, which were modified by magnetron discharge plasma during sputtering of a titanium target, increased human endothelial hybrid cell line (EA-hy 926) adhesion and viability within the plasma treatment time of 6 min [60]. However, cell viability would decrease when the plasma treatment increased to 9 min. In another recent study, PCL electrospun scaffolds embedded with PLGA microsphere were treated with cold atmospheric plasma [61]. Human bone marrow-derived mesenchymal stem cells showed better cellular growth and total collagen production on this kind of scaffolds compared to the plasma untreated ones.

Plasma treatment, an effective and solvent-free technique, is commonly used for surface modification. The disadvantage of plasma treatment is that only localized surface areas can be treated (in depth from several hundred angstroms to 10 nm) without changing the bulk properties of the polymers [41]. Currently, most plasma-based surface modification is performed on two dimensional (2D) polymer substrates, and in some cases on 3D porous scaffold.

1.3.3. Wet chemical methods

The cellular response to a biomaterial may be enhanced in synthetic polymer formulations by mimicking the surface roughness created by the associated nano-structured ECM components of natural tissue. PLGA films with nano-structured surface features were developed by treating them in a concentrated NaOH solution [62]. The results from cell experiments indicated that NaOH treated PLGA enhanced vascular smooth muscle cell adhesion and proliferation compared to conventional PLGA. This surface modification method is not only used on 2D film surfaces but also on 3D constructs. Chen et al. [63] obtained PCL membranes with nanofibrous topology by coating the membrane surface with electrospun nanofibers. When the nanofibers-modified PCL was treated with 5M NaOH for 3h, a favorable 3T3 fibroblast cells morphology and strong cell attachment were observed on the modified surface, possibly due to the unique hydrophilic surface topography. Electrospun PLLA nanofibrous scaffolds were used as a matrix for mineralization of hydroxyapatite [64]. Carboxylic acid groups were introduced on the surface of PLLA scaffolds by hydrolysis in NaOH aqueous
solution. Since calcium ions can bind to the carboxylate groups on the fiber surface, a significant improvement of mineralization was observed on modified PLLA electrospun scaffolds. Hydrophobic polytetrafluoroethylene (PTFE) was wet-chemically treated and subsequently grafted with an endothelial cell specific peptide named arginine-glutamic acid-aspartic acid-valine (REDV). The immobilization of REDV peptide endowed the PTFE substrate a better cellular growth and antifouling properties [65]. Poly(ethylene oxide terephthalate)-poly(butylene terephthalate) rods were surface-modified by etching chloroform and implanted in the subcutaneous space of pigs to form a fibrocellular tissue capsule. The harvested tissue capsules remodeled towards a vessel after grafting bilaterally as carotid artery interposition graft [66]. PLA electrospun scaffolds were wet-chemically treated with polyallylamine to introduce amine groups and subsequently covalently grafted with epidermal growth factor [67]. The functionalized scaffolds supported neural stem-like cells proliferation, which kept viable even in the absence of the soluble growth factor in culture medium.

Although wet chemical methods have been widely applied to modify the surface wettability of biomaterials or to generate new functionalities, some major drawbacks should be noted. This modification technique can cause a partial degradation and scissions of the polymeric chains, resulting in a loss of mechanical properties and a faster degradation process [54]. In addition, the use of hazardous organic solvents might affect cell viability.

1.3.4. Surface graft polymerization

Among the surface modification techniques, surface graft polymerization has emerged as a simple, effective and versatile approach in the surface fictionalization of polymers for a wide variety of applications [19]. Grafting exhibits several advantages [19, 68]: Firstly, the ability to modify the polymer surface to possess desired properties through the choice of different monomers. Secondly, the ease and controllable introduction of graft chains with a high density and exact location of graft chains to the surfaces without changing the bulk properties. Finally, a stable chemical bond is formed between the nanofiber surface and the preformed polymer, which offer endurance of the functional component, in contrast to physically coated polymer chains.

Surface graft polymerization is often initiated with plasma discharge, ultraviolet (UV) light, gamma rays, and electron beams [68, 69]. Chua et al. investigated how a functional poly (ε-caprolactone-co-ethyl ethylene phosphate) (PCLEEP) nanofibrous scaffold with surface-galactose ligand influences rat hepatocytes behavior [70]. The modification of
nanofibrous scaffolds were achieved by conjugating galactose ligands to a poly (acrylic acid) spacer UV-grafted onto the fiber surface. The functionalized nanofibrous scaffolds exhibited the unique property of promoting hepatocyte aggregates within the mesh and around the fibers. Furthermore, hepatocyte functions are maintained on these functional nanofiber substrates, similar to a galactosylated-film substrate configuration.

The grafting of extra cellular protein like collagen, RGD peptides, and gelatin on nanofiber surfaces has become a popular method to develop biomimicking-tissue scaffolds. For example, the modification of electrospun polyethylene terephthalate (PET) nanofibers involved fiber treatment with formaldehyde to generate hydroxyls on the surface, followed by graft polymerization of methacrylic acid monomers initiated by Cerium (IV). Gelatin was then immobilized on the nanofibers by conjugation to the carboxylic acid moieties on their surface [71]. The grafting of gelatin enhanced the adhesion and spreading of cells on nanofibrous scaffolds. In another study, gelatin was grafted onto PLLA scaffolds via aminolysis of the PLLA fibers following by glutaraldehyde coupling [72].

Despite surface graft polymerization have been established as a universal method for surface modification, some issues arise from its mechanism to initiate grafting. An initiator, which is usually cytotoxic, is involved in grafting initiated by a chemical process. UV light induced grafting may cause polymer degradation, which will compromise the mechanical strength of the scaffold [73]. In addition, grafting induced through high energy irradiation causes scission of the base polymer, which can lead to forming block co-polymer [73, 74].

1.4. Applications of nanofibrous scaffolds in tissue engineering

The characteristics of nanofibrous scaffolds that render them promising candidates for tissue engineering include high surface area and porosity, as well as the similarity of their fibrous structure to the physical features of natural ECM [68, 75-77]. These features results in a more biologically compatible environment for cells to grow and perform their regular functions. Therefore, nanofibrous scaffolds have been widely used as scaffolds for tissue engineering such as neural, cartilage, vascular, and bone tissue engineering (Table 2).
### Table 2. Nanofibrous scaffold for Tissue Engineering applications

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<td>Poly-l-lactide (PLLA)</td>
<td><em>In vitro</em></td>
<td>Human aortic smooth muscle cells (HASMCs)</td>
<td>Vascular TE</td>
<td>Hu et al., 2010</td>
</tr>
<tr>
<td><strong>(C) Electrospinning nanofiber</strong></td>
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</tr>
<tr>
<td>Poly(l-lactic acid) (PLLA)</td>
<td><em>In vitro</em></td>
<td>neural stem cells (NSCs)</td>
<td>Neural TE</td>
<td>Yang et al., 2005</td>
</tr>
<tr>
<td>Poly(ε-caprolactone) (PCL)</td>
<td><em>In vitro</em></td>
<td>human mesenchymal stem cells (hMSCs)</td>
<td>Cartilage TE</td>
<td>Li et al., 2005</td>
</tr>
<tr>
<td>Gelatin-modified PET nanofibers</td>
<td>In vitro</td>
<td>endothelial cells</td>
<td>Vascular TE</td>
<td>Ma et al., 2004</td>
</tr>
<tr>
<td>PCL-collagen</td>
<td>Rabbit aortoiliac bypass model</td>
<td>vascular cells</td>
<td>Vascular TE</td>
<td>Tillman et al., 2009</td>
</tr>
<tr>
<td>Poly(ε-caprolactone) (PCL)</td>
<td><em>In vitro</em></td>
<td>Rabbit mesenchymal stem cells (MSCs)</td>
<td>Bone TE</td>
<td>Yoshimoto et al., 2003</td>
</tr>
<tr>
<td>Poly(ε-caprolactone) (PCL)</td>
<td>The omentum of rats</td>
<td>Rats mesenchymal stem cells (MSCs)</td>
<td>Bone TE</td>
<td>Shin et al., 2004</td>
</tr>
</tbody>
</table>
1.4.1. Nanofibrous scaffolds for neural tissue engineering

The nervous system coordinates the action of human and transmits signals between different parts of the body. However, the nervous system has a limited capacity to repair itself after an injury. As a result, patients who have injuries or traumas in the nervous system often suffer from the loss of sensory or motor function, and neuropathic pains [25, 78]. In order to facilitate the re-growth of nerves, many therapeutic approaches have been attempted. One of the most promising approaches is to adopt neural tissue engineering strategy that employs a scaffold or conduit to facilitate the nerve regeneration.

Yang et al. [79] studied aligned and random PLLA electrospun nanofibrous scaffolds for neural tissue engineering. Their result indicated that neural stem cell (NSCs) elongated and their neurites out grew along the direction of the fiber orientation of the aligned nanofibers (Figure 3). Furthermore, NSCs on aligned nanofibrous scaffolds showed higher rate of differentiation than on microfibers. Thus, the aligned PLLA nanofibrous scaffolds showed potential application in neural tissue engineering. Arginine-Alanine-Aspartate (RAD)16-I are the most commonly used peptides in self-assembled peptide scaffolds for neural cell culture [80-82]. Semino et al. [81] developed RAD16-I self-assembled peptide scaffolds as a 3D culture system for cell encapsulation. Primary rat hippocampal cells were cultured on top of the self-assembled nanofibrous scaffold. Immunochemistry assays showed that glial cells and neurons increasingly migrated into the peptide scaffolds to an approximate depth of 400-500 μm from the edge of the tissue after 1 week of culture. Furthermore, mitotic activity of neural cells was maintained for 3 days after migration, which was attributed to the presence of the nanofibrous scaffold environment resembling the native ECM. These results revealed that self-assemble peptide nanofibrous scaffolds are potential substrate for supporting neural tissue regeneration. A hybrid nanofibrous matrix with homogeneous fiber diameter of 20-30 nm was designed by self-assembly of PA with the ability of presenting laminin epitopes (IKVAV or YIGSR) and collagen molecules [83]. Granule cells and Purkinje cells, two major neuronal subtypes of the cerebellar cortex, were cultured on the hybrid scaffold. The result showed the ability to modulate neuron survival and maturation by easy manipulation of epitope density.
In addition to *in vitro* experiments, self-assembled nanofibrous scaffolds have been transplanted into animal models for the treatment of central nervous system injuries. Self-assembled RAD16-I peptide scaffold combined with adult rat Schwann cells and embryonic NPCs were transplanted into adult rats [15]. The results indicated that the scaffolds integrated well with the host tissue with no obvious gap between the implants and the injured sites. A noteworthy observation from these experiments was that large amount of host cells migrated into the scaffolds and extensive blood vessel formation was observed in the implants. Functionalized self-assembling peptides were incorporated into electrospun scaffolds via coaxial electrospinning [84]. The scaffold was fabricated from an emulsion of PCL/PLGA composite together with a functionalized self-assembling peptide as a shell, while PCL/PLGA composite as core demonstrated a better cell viability of mice neural stem cells in *vitro* and in *vivo* inflammatory response in a rodent spinal cord injury model. Silk-gold based artificial conduits, which were fabricated by electrospinning, promoted cellular adhesion and proliferation of Schwann cells *in vitro* [85]. Moreover, the rat sciatic nerve injury model demonstrated that the artificial conduits did not evoke any toxic or immunogenic responses.
1.4.2. Nanofibrous scaffolds for cartilage tissue engineering

Cartilage, a predominantly avascular, alymphatic, and aneural tissue, is composed of chondroblasts embedded within a dense extracellular matrix (ECM) [86]. Cartilage is classified in three types: elastic cartilage, hyaline cartilage and fibrocartilage. Cartilage damage resulting from developmental abnormalities, trauma, aging related degeneration and joint injury cause disability and extensive pain [78, 87]. Due to its limited capacity to self-regenerate, cartilage is an ideal candidate for tissue engineering. In cartilage tissue engineering, chondrocytes and mesenchymal stem cells (MSCs) are commonly used for cartilage repair. Electrospun PCL nanofibrous scaffolds were seeded with human bone marrow-derived MSCs to investigate their ability to support in vitro MSCs chondrogenesis [88]. The results demonstrated that PCL nanofibrous scaffolds in the presence of transforming growth factor-β1 (TGF-β1) induced the differentiation of MSCs to chondrocytes comparable to pellet cultures. Although no inductive property of PCL nanofibrous scaffolds was observed, these meshes had better mechanical properties than cell pellets making them suitable for cartilage tissue engineering. Kisiday et al. [89] investigated a self-assembling peptide hydrogel scaffold for cartilage regeneration. They combined the peptide KDK-12 with bovine chondrocytes and allowed to self-assemble into a hydrogel. Their results indicated that chondrocytes proliferated and maintained a chondrocytic phenotype in the hydrogel. Cells were able to produce cartilage-like ECM which was rich in type-II collagen and proteoglycans. PAs synthesized with a peptide binding sequence to TGF-β1 were design to form nanofibers via self-assembling for cartilage regeneration [17]. The study demonstrated that these scaffolds support the survival and promote the chondrogenic differentiation of human MSCs. In vivo experiments showed that these scaffolds promoted regeneration of articular cartilage in a rabbit model with or even in the absence of exogenous growth factor (Figure 4).
1.4.3. Nanofibrous scaffolds for vascular tissue engineering

Blood vessels perform a very important function of carrying and transporting blood to and from the heart. In addition, they are also important places for the exchange of water and other chemicals between blood and the tissue [19, 37]. Mo et al. [90] developed an aligned electrospun nanofibrous scaffold from biodegradable PLLA-CL (75:25) with the goal of developing constructs for vascular tissue engineering. The fabricated nanometric
fibers resembled the dimensions of natural ECM, possessed mechanical properties comparable to human coronary artery, and supported smooth muscle cell adhesion and proliferation.

A major reason for graft failing is that the graft surface is only partially covered by endothelial cells following a degenerative state. To overcome this problem, developed PET nanofibrous by electrospinning and modified its surface by grafting gelatin. Their study indicated that gelatin-modified PET nanofibers were favorable for spreading and proliferation of endothelial cells and maintained cell phenotype [71]. Tillman et al. [91] studied the in vivo stability of electrospun PCL-collagen scaffolds in vascular regeneration in a rabbit aortal-iliac bypass model. Their finding suggested that PCL-collagen scaffolds maintained enough mechanical strength for cell growth and other physiologic conditions.

1.4.4. Nanofibrous scaffolds for bone tissue engineering

Bone is a composite material made from an organic phase of collagen and glycoproteins and an inorganic phase primarily consisting of hydroxyapatite (HA) [92]. The organic phase provides bone resilient nature and the inorganic minerals is responsible for bone hardness [19, 93]. Bone may be lost caused by trauma, fractures, periodontitis, cancer, infectious disease, and osteoporosis [94]. Presently, many approaches have been developed for bone regeneration, such as autografts, metal implants, and allografts. However, all these methods have obvious drawbacks. For example, autografts present problems associated with a limited donor source as well as a secondary surgery site. Therefore, tissue engineering approaches are presently being investigated as a promising method for bone regeneration. Yoshimoto et al. [95] studied the potential of non-woven PCL scaffolds generated by electrospinning for bone tissue engineering. MSCs derived from bone marrow of neonatal rats were seeded on the nanofibers. The results demonstrated that MSCs migrated into the scaffolds and produced abundant ECM. Based on this study, Shin et al. [96] implanted MSCs along with PCL nanofibrous scaffolds into the omentum of rats. Their study indicated ECM formation throughout the scaffolds along with the evidence of mineralization and type I collagen synthesis. When HA is incorporated into nanofibrous scaffolds it does not only improve the mechanical properties, but also creates more biomimetic constructs. Either nanoparticles of HA, morphogenetic proteins (e.g. BMP-2) or both were incorporated in electrospun nanofibrous scaffolds of silk fibroin [97]. Scaffolds were seeded with human MSCs for 31 days in osteogenic medium. HA nanoparticles were associated with improved bone formation. Furthermore, when silk fibroin scaffolds were combined with both nanoparticles
of HA and BMP-2, they were associated with the highest observed mineralization.

Figure 5. Representative microcomputed tomography images of bone regeneration in the rabbit model at 6 weeks postimplantation. White arrows in the first column identify the defect site. Remaining b-tricalcium phosphate fragments are identified in the 2D transverse images of the bone graft substitute (BGS) scaffolds [98].

Porous three-dimensional tyrosine-derived polycarbonate (TyrPC) scaffolds were developed for bone regeneration using a combination of salt-leaching and phase separation techniques [98]. The TyrPC scaffolds treated with or without recombinant human bone morphogenetic protein-2 (rhBMP-2) were implanted in a rabbit calvarial critical-sized defect model for 6 weeks. The in vivo studies showed that TyrPC scaffolds treated with rhBMP-2 or coated with calcium phosphate alone promoted bone regeneration at 6 weeks post-implantation (Figure 5). Moreover, TyrPC polymeric scaffolds even without addition any biological agents induced similar bone regeneration to a commercially available bone graft substitute. Biomimetic nanofibrous scaffolds made
Chapter 1

of hydroxyapatite/collagen/chitosan (HAp/Col/CTS) showed higher capacity than control scaffolds (tissue culture plate, CTS, and HAp/CTS) for the osteogenesis of induced pluripotent stem cell-derived mesenchymal stem cells (iPSC-MSCs) in vitro [99]. Furthermore, mouse cranial defects were created to investigating the efficacy of biomimetic scaffolds for bone repair in vivo. After six weeks implantation, HAp/Col/CTS scaffolds showed nearly a 2-fold higher bone mineral density than other scaffolds. Mimicking the formation of vertebrate bone during early stage, Rajamani et al. developed nanofibrous collagen scaffolds containing catecholamines and Ca$^{2+}$ for bone regeneration [100]. The developed collagen composite scaffolds had outstanding mechanical properties with a Young's modulus close to cancellous bone. Human fetal osteoblasts displayed increased cell proliferation and differentiation in the collagen composite scaffolds compared to tissue culture plate or pristine collagen mats.

1.5. Scope and Outline of Thesis

Mimicking the architecture of ECM is one of the key challenges of tissue engineering. Amongst all the methods used to generate artificial ECM, nanofibrous scaffolds demonstrated the most promising results. Nanofibrous scaffolds, irrespective of their method of synthesis, are characterized by high surface area and enhanced porosity, which are highly desired for tissue engineering and drug delivery applications.

At present, there are three dominant approaches to generate nanofibrous scaffolds: molecular self-assembly, phase separation, and electrospinning. Among these approaches, electrospinning is the most widely studied approach since it is a simple and versatile technique that can produce fibers with a diameter from the micrometer down to nanometer range. Furthermore, it allows one to tailor many aspects of the resulting scaffolds [29, 101, 102]: (1) the fiber diameter can easily be controlled by varying the solution properties and the processing parameters; (2) the nanofibers can be collected with a rich variety of aligned/random structures using specially designed collectors; (3) they can be stacked and/or folded to form complex structures or architectures; (4) they can be obtained with a hollow or core-sheath structure by changing the configuration of the spinneret; (5) nanofibers with porous surface structure can be achieved by altering the parameters such as solvent and humidity. The potential of nanofibrous scaffolds for the regeneration of various tissues are now under extensive investigation. In spite of the great achievements behind the designing of nanofibrous scaffolds via electrospinning, there is still plenty of room for improvement. The further development of melt-electrospinning direct writing in the last few years has shown the advantage of creating
nanofiber patterns that can guide tissue organization. In fabricating smart scaffolds to
direct cell fate, more knowledge over chemical and physical stimuli that can be built-in the
biomaterial formulation of choice is desirable. In developing new techniques to fabricate
smart nanofibrous scaffolds, the configuration seen in native tissues should be taken into
consideration if we want to achieve the highest degree of biomimicry possible. Finally,
charge repulsion should be improved when aiming at clinically relevant large scaffold
fabrication.

To better restore and recover the damaged tissues or organs, there is a great need for
bioactive scaffolds which not only provide space and mechanical properties for cellular
growth and migration, but can also offer chemical and physical cues to control cell fate
and finally direct tissue formation. The work presented hereafter is aimed to provide a
small library of possible strategies for designing bioactive scaffolds for skeletal tissue
engineering through electrospinning-based techniques.

Human mesenchymal stem/stromal cells (hMSCs) have been chosen as a cell model,
because of their known potential as a cell source for regenerative medicine due to their
capacity to differentiate into a variety of cell type, including adipocytes, chondrocytes, and
osteoblasts[103]. The approaches to achieve this goal are developed as follows: (1)
investigate the effect of chemical composition of a scaffold and correlate its degradation
and wettability properties to the response of seeded cells; (2) fabricate novel flexible
ceramic fibrous mats with tailored surface roughness and study the influence of surface
roughness of electrospun scaffolds on cell faith; (3) modify the surface roughness of
electrospin fibers via regulating humidity during electrospinning; (4) fabricate buckled
fibrous meshes to mimic the crimped feature of collagen fibers in native tissue using
thermal shrinkable films; (5) and develop a scaffold mimicking the zonal organization of
articular cartilage using a new direct writing electrospun technique.

Scaffold-based tissue engineering is nowadays an established strategy for repairing
damaged tissues and organs. Therefore, there is a great need for new biodegradable
polymers, like polyesters, to fabricate scaffolds with enhanced bioactivity Such new
bioactive scaffolds should be characterized by having more flexibility in tuning their
physical (e.g. mechanical and hydrolysis properties) and biological (e.g. bioactive cues for
triggering stem cell differentiation) properties. When achieved, this will give the
unprecedented opportunity to steer cell differentiation through the rationale design of
synthetic functional materials, whereas today this is mostly done by biological molecules
in solution or coupled to synthetic materials. To this end, in chapter 2 electrospun
scaffolds were fabricated from a new copolyester family containing either butylene thiodiglycolate (BTDG) or butylene diglycolate (BDG) sequences by reactive blending. The synthesized polyesters displayed tunable mechanical properties and hydrolysis rate depending on the molecular architecture and on the kind of etheroatom introduced along the polymer backbone. The potential of these new scaffolds in skeletal tissue engineering was examined *in vitro* by culturing hMSCs on them in basic or differentiation media.

The main drawback in hard tissue regeneration of bioactive ceramic scaffolds is their low mechanical strength. Stabilized zirconia, especially the tetragonal 3 % yttrium-stabilized zirconia (YSZ) phase, has been considered as a bioinert ceramic material with high mechanical strength. In chapter 3, novel flexible nanofibrous YSZ scaffolds were fabricated by electrospinning. Their surface roughness could be tailored by varying the heat treatment methods. The perfromance of this new ceramic scaffold in bone tissue engineering was further investigated. It has been fully demonstrated that the surface roughness of 2D materials have influence on cellular response, such as cell proliferation, attachment, and differentiation. However, the biological environment for cells in a tissue or organ is a 3D structure instead of a 2D flat substrate. To investigate the effect of surface roughness of flexible ceramic 3D nanofibrous scaffolds on the skeletal differentiation of hMSCs, electrospun meshes with same surface chemistry, yet different surface roughness, were obtained by varying the relative humidity levels during electrospinning (Chapter 4).

The periodic waviness of native collagen fibrils is observed in nearly all human tissue, such as tendon, blood vessels, and valve leaflets. The features of such a waived or crimped structure ensure that loads are sustained both by the surrounding matrix and the fiber network at low levels of tensile strain. To get a better resemblance of natural collagen fibrils, in chapter 5, we exploit a simple but efficient and versatile method to produce crimp patterned electrospun fibers by using thermally shrinkable films. hMSCs were cultured on the wavy fibers to evaluate whether cell infiltration into the scaffolds could be also improved, which is nowadays still one of the major drawback of electrospun meshes. Given the resemblance of the obtained fibers with the crimp patterns typical of tendons and ligaments, we also investigated if the wavy patterns could regulate the TGF-β pathway, a key signaling pathway involved in tendon and ligament maintenance.

One of the limitations of traditional solution electrospinning technique lies in designing of scaffold’s architecture, which often results in a densely packed structure. However, the structure of most human tissue or organ is a more complex hierarchical structure. For
example, articular cartilage contains four main zones: the superficial, middle, deep and calcified cartilage zone. To overcome this problem, in chapter 6, a new technique for the fabrication of microfibrous structures in a precise and controllable way was developed, using a direct-write electrospinning process. Furthermore, a scaffold mimicking the zonal organization of articular cartilage was produced by this method and its performance on cartilage tissue formation was evaluated through the chondrogenic differentiation of seeded hMSCs in vitro.

Chapter 7 gives an overview of the obtained results and of future perspective in designing scaffold for skeletal tissue engineering.

Finally, chapter 8 discuss the valorisation potential of the research presented in this thesis.
Chapter 1

References

General Introduction and Aims


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General Introduction and Aims


Tailoring Chemical and Physical Properties of Fibrous Scaffolds from Block Copolyesters Containing Ether and Thio-ether Linkages for Skeletal Differentiation of Human Mesenchymal Stromal Cells

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Abstract

Bioactive scaffolds for tissue engineering call for demands on new materials which can enhance traditional biocompatibility requirements previously considered for clinical implantation. The current commercially available thermoplastic materials, such as poly(lactic acid) (PLA), poly(glycolic acid) (PGA), poly(ε-caprolactone) (PCL) and their copolymers, have been used to fabricate scaffolds for regenerative medicine. However, these polymers have limitations including lacking of broadly tuning mechanical and degradable properties, and activation of specific cell-scaffold interactions, which limit their further application in tissue engineering. In the present study, electrospun scaffolds were successfully fabricated from a new class of block poly(butylene succinate)-based (PBS-based) copolyesters containing either butylene thiodiglycolate (BTDG) or butylene diglycolate (BDG) sequences. The polyesters displayed tunable mechanical properties and hydrolysis rate depending on the molecular architecture and on the kind of heteroatom introduced along the polymer backbone. To investigate their potential for skeletal regeneration, human mesenchymal stromal cells (hMSCs) were cultured on the scaffolds in basic, osteogenic and chondrogenic media. Our results demonstrated that PBS-based copolyesters containing thio-ether linkages (i.e. BTDG segments) were more favorable for chondrogenesis of hMSCs than those containing ether linkages (i.e. BDG sequences). In contrast, PBS-based copolyesters containing ether linkages showed enhanced mineralization. Therefore, these new functional scaffolds might hold, potential for osteochondral tissue engineering applications.

Keywords:
Aliphatic copolyesters, ether and thioether linkages, electrospun scaffolds, human mesenchymal stromal cells, skeletal differentiation
2.1. Introduction

A promising strategy for repairing damaged tissues and organs is to design scaffolds able to support and eventually control cell activity, i.e. adhesion, proliferation and differentiation. Electrospinning is a simple technique capable of producing non-woven scaffolds made of continuous fibers with diameters ranging from few tens of nanometers up to several micrometers[1]. The resulting fibrillar networks mimic the morphology of native extracellular matrix (ECM), which serves to organize cells and provides signals for cellular responses [2].

Among the manifold of current commercially available thermoplastic materials used to fabricate three-dimensional (3D) scaffolds for regenerative medicine, aliphatic polyesters undoubtedly represent so far the most extensive studied class, since they combine good physico-chemical and mechanical properties with assessed biocompatibility[3]. Poly(lactic acid) (PLA), poly(glycolic acid) (PGA), poly(ε-caprolactone) (PCL) and their copolymers are typically used[4] and their hydrolytic and enzymatic degradation under physiological conditions has been extensively proven[3]. Although these polymers have found many applications in clinical practice, the wide tuning of their mechanical, physico-chemical and biological properties is somehow limited[4]. For instance, the copolymerization of PLA with PCL enables to decrease polymer elastic modulus, thus obtaining more deformable materials that, however, are highly hydrophobic and display a very slow degradation rate. Hydrophilicity and hydrolysis rate can be significantly enhanced through copolymerization of PLA with PGA but, in this case, the rigidity of the material persists. This hampers the fabrication of 3D scaffolds capable of adapting to the needs of the specific tissue to be regenerated. Therefore, new polyesters with more flexibility in tuning their properties attract considerable attention in regenerative medicine.

In the last few years, an increasing number of researchers proposed poly(butylene succinate) (PBS) as a possible alternative to the above mentioned aliphatic polyesters due to the following advantages: (i) simple synthetic strategy through melt polycondensation; (ii) low cost of raw materials (1,4-butandiol and succinic acid), which can be also obtained from renewable resources[5]; (iii) high melting temperatures among poly(alkylene dicarboxylate)s (T_m=113-115°C); (iv) good thermal stability (thus a wide processing temperature range); (v) proven biodegradability and biocompatibility[6]. Furthermore, PBS is already commercialized by different companies for various applications such as compost bags and mulching films, but also fishery, civil engineering and construction and for common household goods.
Many efforts have been recently devoted to target PBS modification aiming to achieve better performances with respect to the intended application. For biomedical applications, composites have been realized by incorporating organic or inorganic fillers such as chitosan, collagen, calcium phosphate or hydroxyapatite along the PBS macromolecular chain[6]. A plethora of PBS-based blends and copolymers have been also obtained[6]. Among other copolymerization techniques, reactive blending is a versatile and solvent free approach which permits to prepare block copolymers of different molecular architecture by simply varying the reaction time [7].

Recent studies have demonstrated that the properties of scaffold, such as scaffold elasticity, structure and organization, and chemistry, can direct stem cell differentiation into specific lineage[8-10]. In this study, electrospun scaffolds fabricated from a class of block PBS-based copolyesters containing either butylene thiodiglycolate (BTDG) or butylene diglycolate (BDG) sequences, obtained by reactive blending, are presented. In particular, the possibility to use the synthesized copolymers as an alternative to conventional polyesters was explored to achieve enhanced control over stem cell differentiation. In these systems, the addition of heteroatoms along the macromolecular chains has been demonstrated to tune chain flexibility, thus affecting polymer crystallinity and mechanical properties of aliphatic polyesters [11, 12]. Moreover, through this strategy surface hydrophilicity has been modified, due to the presence of electronegative oxygen and sulphur atoms[13-15]. All these factors significantly impact the biodegradation rate and biocompatibility of the final materials [16, 17]. Scaffolds with tailored chemical composition and physical properties have been prepared and characterized. Their hydrolytic degradation profile has been evaluated under physiological conditions. Finally, we show that these copolymers may have the potential to steer stem cell differentiation towards skeletal lineages.

2.2. Materials and Methods

2.2.1. Materials

Dimethylsuccinate (DMS), 1,4-butanediol (BD), diglycolic acid (DGA), thiodiglycolic acid (TDGA), and titanium tetrabutoxide (TBT) (Sigma-Aldrich) were reagent grade products. All the reagents were used as supplied with the exception of TBT which was distilled before use. Chloroform, dichloromethane (DCM), 2-chloroethanol (CE) and hexafluoro-2-propanol (HFIP) were purchased from Sigma-Aldrich and were used without any further purification.
2.2.2. Polymer synthesis

Starting homopolymers, i.e. PBS, poly(butylene diglycolate) (PBDG) and poly(butylene thiodiglycolate) (PBTDG) were synthesized in bulk by the usual two steps melt polycondensation, starting from 1,4-butanediol (BD) and DMS, DGA or TDGA, as reported elsewhere[11, 12]. Two blends, PBS/PBDG and PBS/PBTDG, were obtained by melt mixing for 10 min equimolar amounts of PBS with PBDG and PBTDG, respectively, according to the procedures previously described [11,12]. Poly(butylene succinate/diglycolate) (P(BS\textsubscript{11}BDG\textsubscript{11})) and poly(butylene succinate/thiodiglycolate) (P(BS\textsubscript{10}BTDG\textsubscript{10})) copolymers were obtained by melt mixing equimolar amounts of PBS with either PBDG (for 30 min) or PBTDG (for 45 min), respectively[11, 12]. The reactions were carried out at 225°C, under dry nitrogen atmosphere to prevent polymer degradation. Copolymer formation was catalysed by the residual titanium tetrabutoxide used for the synthesis of the parent homopolymers. The reactive blending provided two block copolymers (P(BS\textsubscript{11}BDG\textsubscript{11}) and P(BS\textsubscript{10}BTDG\textsubscript{10})) with similar block length.

2.2.3. Scaffold and film preparation

PBS, PBS/PBDG and PBS/PBTDG blends and P(BS\textsubscript{11}BDG\textsubscript{11}) and P(BS\textsubscript{10}BTDG\textsubscript{10}) copolymers were subjected to electrospinning to obtain 3D mats. Scaffolds were produced by using a customized electrospinning apparatus, comprised of a high-voltage power supply (SL 50 P 10/CE/230; Spellman), a syringe pump (KD Scientific 200 series), a glass syringe, a stainless steel blunt-ended needle (inner diameter = 0.84 mm) connected with the power supply electrode, and a grounded aluminium plate-type collector (7 cm × 7 cm). Polymer solution was dispensed through a Teflon tube to the needle which was placed vertically on the collecting plate at a measured distance (D). The scaffolds were produced at room temperature (RT) and a relative humidity of 40 ± 5%. The electrospun solution and the operating conditions for each polymer were selected after optimization experiments aimed at obtaining bead-free fibers and are reported in Table S1. Electrospun mats were kept under vacuum over P\textsubscript{2}O\textsubscript{5} at RT overnight to remove residual solvents.

Films (0.2 mm thick) of PBS, blends and copolymers were obtained by compression moulding the polymers between Teflon plates, with an appropriate spacer, at 140 °C for 1 min under a pressure of 2 ton m\textsuperscript{-2} (Carver C12, laboratory press).
2.2.4. Material characterization

Polymer structure and composition were determined by means of proton nuclear magnetic resonance (\(^1\)H-NMR) spectroscopy, whereas the distribution of the co-monomeric sequences along the polymer chain was evaluated by \(^{13}\)C-NMR spectroscopy. The measurements were carried out at RT by employing a Varian Inova 400-MHz instrument (Agilent Technologies, USA). Molecular weights were evaluated by gel permeation chromatography (GPC) at 30°C using a 1100 HPLC system (Agilent Technologies, USA) equipped with PLgel 5-μm MiniMIX-C column (Agilent Technologies, USA). A refractive index was employed as detector. Chloroform was used as eluent with a 0.3 mL/min flow and sample concentrations of about 2 mg/mL. A molecular weight calibration curve was obtained with polystyrene standards in the range of molecular weight 2000-100,000 g/mol. Thermogravimetric analysis (TGA) was carried out under nitrogen atmosphere using a Perkin Elmer TGA7 apparatus (gas flow: 30 mL/min) at 10 °C/min heating rate, up to 900 °C. Calorimetric measurements were conducted by using a Perkin Elmer DSC7 instrument. In the typical setup, the external block temperature control was set at -120 °C and weighed samples of c.a. 10 mg were heated up to 40 °C above fusion temperature at a rate of 20 °C/min. Scanning electron microscopic (SEM) observations were carried out using a Philips 515 SEM (FEI, Eindhoven, The Netherlands) at an accelerating voltage of 15 kV, on samples sputter coated with gold. The distribution of fibre diameters was determined through the measurement of approximately 150 fibres by means of an acquisition and image analysis software (EDAX Genesis, New York, USA), and the results were given as the average diameter ± standard deviation. Static contact angle measurements were performed on polymer films using a KSV CAM101 (KSV, Espoo, Finland) instrument at ambient conditions, by recording the side profiles of deionized water drops for image analysis. Five drops were observed on different areas for each film, and contact angles were reported as the average value ± standard deviation. Stress-strain measurements were performed with an Instron 4465 tensile testing machine equipped with a 100N load cell on rectangular films, according to ASTM D882-02 standard. Sample were 5 mm wide, the gauge length was 20 mm and the cross-head speed was 5 mm/min. Load-displacement curves were obtained and converted to stress-strain curves. At least six replicate specimens were run for each sample and results were provided as the average value ± standard deviation.
2.2.5. Hydrolytic degradation studies

Hydrolytic degradation studies were carried out on rectangular mats of PBS, blends and copolymers in triplicate. Prior to degradation experiments, each specimen was dried over P₂O₅ under vacuum at RT, and weighed to obtain the sample initial mass. Mats were individually immersed in phosphate buffered solution (0.1 M, pH = 7.4) and incubated in a SW22 Julabo shaking water bath at 37 °C and 50 rpm. The buffer solution was periodically changed to keep the pH constant during the entire time scale of the degradation experiments. At different time intervals, triplicate specimens for each sample were recovered from the bath, repeatedly washed with deionized water and dried over P₂O₅ under vacuum to constant weight. The mass loss was determined gravimetrically by comparing the dry weight remaining at a specific time with the initial sample weight. The molecular weight of each recovered sample was determined by GPC. ¹H-NMR analysis was performed to detect changes in the molecular composition during the hydrolysis experiments and differential scanning calorimetry (DSC) was used to evaluate the change in crystallinity degree.

2.2.6. Cell culture and seeding

Human mesenchymal stromal cells (hMSCs, donor No. 8001L) were provided from Texas A&M Health Science Center, which has supplied standardized preparations of MSCs to hundreds of laboratories under the auspices of an NIH/NCRR grant (P40 RR 17447-06). Briefly, human bone marrow aspirates were obtained under a protocol approved by an institutional review board. Mononuclear cells were separated from bone marrow aspirate using density centrifugation. They were plated to achieve adherent hMSCs and harvested when cells reached 60–80% confluence. These cells were called passage zero (P0) cells and were further expanded, harvested and frozen at passage one (P1). Prior to release, the frozen P1 cells were analyzed over three passages to assess successful expression of MSCs surface markers[18, 19]. To achieve sufficient cells for seeding onto scaffolds, P1 cells were expanded in proliferation medium (PM), consisting of α-MEM (Gibco), 10% fetal bovine serum (Lonza), 2 mM L-glutamin (Gibco), 0.2 mM ascorbic acid (Sigma-Aldrich), 100 U/mL penicillin and 100 mg/mL streptomycin (Gibco) and 1 ng/mL recombinant basic fibroblast growth factor (bFGF, Instruchemie). Cells were harvested for further sub-cultures when they became 80-90% confluence. Electrospun scaffolds were placed in cell culture plates and pressed with rubber O-rings (Eriks BV, The Netherlands) to prevent them from floating. The specimens were sterilized by immersion in 70% (v/v)
ethanol for 15 min for three times. Subsequently, scaffolds were washed for 1 min with PBS for three times and immersed in basic medium (BM) overnight before cell seeding. A density of 5×10^3 cells/cm^2 was used to seed hMSCs onto electrospun scaffolds. Cell-scaffold constructs were cultured for up to 28 days in BM consisting of PM without bFGF, osteogenic medium (OM) comprising BM supplemented with 1×10^{-6} M dexamethasone), mineralization medium (MM) composed of OM plus b-glycerol phosphate, and chondrogenic medium (CM) containing DMEM (Gibco,) supplemented with 1% ITS premix (Micronic BV), 50 μg/mL ascorbic acid (Sigma-Aldrich), 100 units/mL penicillin (Life technologies), 100 μg/mL streptomycin (Life technologies), 100 nM dexamethasone (Sigma-Aldrich), 40 μg/mL proline (Sigma-Aldrich), 100 μg/mL sodium pyruvate (Life technologies), 0.01 μg/mL TGFβ1 (R&D systems). Culture experiments were performed in an incubator with 5% CO_{2} humid atmosphere at 37 °C. Culture medium was refreshed every 2 days.

2.2.7. Metabolic activity

Cell culture medium was removed from cell-scaffold constructs and supplemented with a medium containing 10% (v/v) PrestoBlue™ reagent (Life technologies). The cell-scaffold constructs were incubated at 37 °C for 1 h in the dark. Fluorescence was measured at 590 nm using a multi-label plate reader (Perkin Elmer Victor3 1420, USA).

2.2.8. Cell morphology analysis

Cell morphology and attachment on scaffolds was determined by SEM using an XL 30 ESEM-FEG (Philips, The Netherlands). Briefly, samples were rinsed twice with PBS, fixed with 10% formalin for 2h. After washing with PBS two times, the samples were dehydrated with a gradient ethanol series, dried by using critical point drying (Balzers CPD 030) and sputtered with gold (Cressington) before SEM analysis.

2.2.9. Quantification of DNA

Samples were washed with PBS three times and stored at -80 °C overnight. After thawing, samples were lysated for 16 h with a Tris/EDTA buffer containing 1mg/mL proteinase K, 185 μg/mL iodoacetamine (Sigma-Aldrich) and 10 μg/mL Pepstatin A (Sigma-Aldrich). Subsequently, the quantification of the total amount of DNA was examined by using the CyQuant® DNA assay (Molecular Probes, Eugene, USA) as previously reported[20]. Lysate was pipetted in duplicate to a black 96-well plate, followed by addition of NaCl-EDTA buffer containing component B of the kit (20×) and RNase (1000×). The plate was
incubated for 1h at RT. Subsequently, Gr-dye solution was added and the samples were incubated for 15 min. The fluorescence signal was determined using a spectrophotometer (Victor3 1420, Perkin Elmer, USA) at an excitation wavelength of 480 nm and emission wavelength of 520 nm.

2.2.10. Alkaline phosphatase activity (ALP) assay

The protocol of measuring ALP activity was similar to a previous study[21]. Briefly, the cell-scaffold constructs were digested for 1 h at RT in a cell lysis buffer at a pH 7.8 composed of 0.1 M KH₂PO₄, 0.1 M K₂HPO₄ and 0.1% Triton X-100. Then, cell lysates were pipetted in duplicate to 96-well plate. After addition of CDP-star® (Roche Life Science), the lysate were incubated for 15 min at RT. The measurements were performed on a Victor3 1420 (Perkin Elmer, USA) plate reader. The value of ALP activity was normalized to DNA quantity per scaffold.

2.2.11. Glycosaminoglycan (GAG) assay

Cell-scaffold constructs were washed with PBS three times and stored at -80 °C for at least 24 h. After thawing, the constructs were digested for 16 h with Tris/EDTA buffer as mentioned above. The amount of GAG was determined spectrophotometrically after reaction with 16 mg of dimethylmethylened blue dye (DMMB, Sigma-Aldrich) in a 10 mM hydrochloric acid solution containing 3.04 g/L of glycine and 2.37 g/L of NaCl. The absorbance was measured on a micro plate reader (Multiskan GO, Thermo Fisher, USA). The amount of GAG was calculated using a standard of chondroitin sulfate (Sigma-Aldrich).

2.2.12. Histological staining

ALP staining was carried out by using an alkaline phosphatase kit (Sigma-Aldrich). Samples were fixed in 40% citrate working solution in acetone for 30 s and rinsed with deionized water three times. Alkaline dye mixture was added and incubated for 30 min at RT in the dark. Samples were finally rinsed thoroughly with deionized water three times. For alizarin red staining (ARS) assay, after fixation in 10% formalin for 1 h, samples were rinsed with PBS three times, washed twice with distilled water and stained with a freshly filtered 2% aqueous Alizarin red solution (pH 4.2) for 2 min. The excess stain was washed off thoroughly with distilled water. To quantify the orange-red coloration of ARS, the protocol was performed as described in a previous study [22, 23]. After removing excess water, samples were transferred to a microcentrifuge tube. A 10% acetic acid
solution was added and incubated for 30 min at RT with shaking. The slurry was centrifuged for 15 min at 20,000 g after vortexing for 30 s. Then, the supernatant was transferred to a new 1.5-mL microcentrifuge tube. A 10% ammonium hydroxide was utilized to neutralize the acid. Aliquots of the supernatant were read in triplicate at 405 nm in 96-well format using opaque-walled, transparent-bottomed plates (Fisher Lifesciences). All the images for ALP staining and ARS staining were taken using a stereomicroscope (Nikon SMZ800 with Q-imaging Retiga 1300 camera).

2.2.13. Quantitative PCR analysis

To analyze the gene expression of hMSCs on scaffolds, total RNA was isolated using a combination of TRIzol® with a NucleoSpinRNAII ISOLATON® kit (Bioke). Briefly, scaffolds at day 7 and day 28 were rinsed with PBS three times and 1 mL of TRIzol was added to each sample. After addition of chloroform, phase separation occurred by centrifugation at 12,000 g. The aqueous phase containing RNA was collected, mixed with equal volume of 75% (v/v) ethanol and loaded onto an RNA binding column of the kit. After a series of purification steps, RNA was re-suspended in RNase-free water and collected using a clean Eppendorf® tube by centrifugation at 12,000 g. The concentration and purity of RNA were measured by using a spectrophotometer (ND 1000, Nanodrop Technologies, USA). iScript (Bio-Rad, Veenendaal, the Netherlands) was used for the cDNA synthesis following the manufacturer’s instruction. Quantitative polymerase chain reaction (qPCR) was carried out using SYBR-green supermix (Bio-Rad). Cycling parameters for running qPCR were set as follows: initial denaturation at 95 °C for 10 min, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. The sequences of the analyzed primers are shown in Table S3. Data are expressed as fold change relative to controls, after normalizing to B2M.

2.2.14. Statistical analysis

All data points are expressed as mean ± standard deviation. Statistical difference was determined by using two-way analysis of variance (ANOVA), followed by a Tukey’s post-hoc test. A value of p < 0.05 was considered statistically significant.

2.3. Results and discussion

2.3.1. Electrospinning and sample characterization

PBS, PBDG and PBDTG displayed a similar chemical structure, all containing two ester groups per repeating unit along a saturated aliphatic chain (Figure 1). With respect to
PBS, differences can be observed in the acid sub-units: PBDG contains an ether bond while PBTDG a thio-ether bond. We previously demonstrated that these minimal differences in the polymer backbone have a remarkable effect on polymer thermo-mechanical properties [11, 12]. In the present study, physical blends of PBS with PBDG and PBTDG and copolymers of PBS containing either BDG or BTDG units with long block length were considered for scaffold fabrication, since these polymeric systems possess physico-chemical and mechanical characteristics significantly different from each other and from the PBS homopolymer. The molecular characterization of the polymeric systems investigated in the present work is reported in Table 1.

Both the blends and the copolyesters are characterized by relatively high molecular weights. This result indicates that no relevant thermal degradation occurred during the reactive blending. Looking into more detail, a slight decrease of molecular weight can be observed for the block copolymers. This result has been previously described and can be explained on the basis of two opposite effects: chain scission and transesterification reactions [11, 12]. For short mixing times, i.e. those here employed, the chain scission reactions prevailed over transesterifications and therefore a decrease of molecular weight could be detected.

The chemical structure and composition of all copolyesters, determined by $^1$H-NMR were found to be consistent with the expected ones. $^{13}$C-NMR measurements were performed to study the structural changes due to the reactive blending. As far as the two
blends are concerned, the NMR spectrum is the simple addition of PBS and PBTDG ones, indicating that no appreciable chemical reactions took place during the mixing. On the contrary, in \( \text{P(BS}_{10}\text{BTDG}_{10}) \) and \( \text{P(BS}_{11}\text{BDG}_{11}) \) two new peaks developed in the region between 64.1 and 65.5 ppm (where the signals due to the carbon atoms of the 1,4-butane diol glycol subunit are located), due to the BS-BTDG and BS-BDG mixed sequences, respectively[11, 12]. Lastly, the average block length has been calculated from the equations previously reported [11, 12].

Electrospinning conditions parameters were optimized in order to obtain defect-free fibres. The scaffolds were highly porous with microscale interstitial pores made of randomly oriented fibres. In the case of \( \text{P(BS}_{11}\text{BDG}_{11}) \), two different polymeric solutions were prepared to obtain scaffolds with either micrometric or nanometric diameters (i.e. \( 890 \pm 410 \text{ nm and } 410 \pm 160 \text{ nm} \)) that were labelled \( \text{P(BS}_{11}\text{BDG}_{11})-\text{m} \) and \( \text{P(BS}_{11}\text{BDG}_{11})-\text{n} \), respectively (Figure 1 and Table S1).

Electrospun fibres were characterized by TGA and DSC. TGA did not reveal residual solvents in the fabricated fibres and confirmed that electrospinning process did not affect polymer thermal degradation behaviour. DSC was used to evaluate thermal transitions in the electrospun mats (Table 1). The calorimetric data of \( \text{P(BS}_{11}\text{BDG}_{11})-\text{m} \) and \( \text{P(BS}_{11}\text{BDG}_{11})-\text{n} \) were identical within the experimental errors. The phase behaviour of the polymers under investigation was similar. As far as the glass transition phenomenon is concerned, only one glass transition temperature \( (T_g) \) was always clearly evident suggesting the presence of a homogeneous amorphous state. All polymers were semicrystalline, being the corresponding calorimetric traces (Figure S1) characterized by a significant melting endotherm. The overall heat of fusion \( (\Delta H_m) \) decreased from blends to block copolymers because of the reduced length of crystallisable sequences. The copolymer containing ether-oxygen showed a slightly bigger \( \Delta H_m \) associated to the PBS crystal phase compared to the thio-ether containing copolymer. This implied that the PBS crystallization was less hampered in the copolymer containing BDG co-units. Interestingly, in the PBS/PBDG blend two distinct melting phenomena were evident, suggesting the simultaneous presence of the crystalline phases of the two homopolymers. This result has been previously confirmed by X-ray analyses [12]. For the blends, the melting temperature \( (T_m) \) was equal to that of PBS (and of PBDG homopolymer in the case of PBS/PBDG), while it decreased in the block copolymers because of the formation of a less perfect crystalline phase.
Table 1. Polymer and scaffold characterization.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>( M_n^a ) (g/mol)</th>
<th>PDI(^a)</th>
<th>( T_g^b ) (°C)</th>
<th>( T_m, \text{iso}^b ) (°C)</th>
<th>( \Delta H_m, \text{iso}^b ) (J/g)</th>
<th>WCA(^c) (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS</td>
<td>34600</td>
<td>2.5</td>
<td>-32</td>
<td>/</td>
<td>115</td>
<td>/</td>
</tr>
<tr>
<td>PBS/PBTDG</td>
<td>37000</td>
<td>2.6</td>
<td>-45</td>
<td>/</td>
<td>113</td>
<td>/</td>
</tr>
<tr>
<td>P(BS(<em>{10})BTDG(</em>{10}))</td>
<td>30200</td>
<td>2.5</td>
<td>-44</td>
<td>/</td>
<td>100</td>
<td>/</td>
</tr>
<tr>
<td>PBS/PBDG</td>
<td>23600</td>
<td>2.8</td>
<td>-29</td>
<td>63</td>
<td>114</td>
<td>28</td>
</tr>
<tr>
<td>P(BS(<em>{11})BDG(</em>{11}))</td>
<td>19700</td>
<td>2.3</td>
<td>-30</td>
<td>/</td>
<td>106</td>
<td>/</td>
</tr>
</tbody>
</table>

\(^a\) by GPC performed on polymers in form of powder; \(^b\) by DSC performed on polymeric scaffolds; \(^c\) by WCA performed on polymers in form of film

To investigate the effect of polymer chemistry on material hydrophilicity, water contact angle (WCA) measurements were performed on hot-pressed films and the results reported in Table 1. The significant decrease of WCA values in blends and copolymers was due to the introduction of highly electronegative atoms (O and S) along the PBS polymeric chain, which has the effect of increasing material surface wettability, as previously reported for other polymeric systems[13-17, 24]. The polymers containing BDG repeating units displayed a lower WCA than those containing BTDG units. Thus, the former were more hydrophilic as a consequence of the higher electronegativity of oxygen atoms with respect to sulphur ones.

In order to correlate bulk mechanical properties with polymer chemical structure, tensile stress-strain measurements were carried out on polymeric films rather than electrospun fibers. Indeed, it is somewhat difficult to make a direct comparison among stress-strain data of non-woven electrospun fibrous matrices since it is already established that mechanical data depend not only on the type of polymeric material and on single fiber features, but also on the arrangement of the fibers in the nonwoven matrices, such as fiber direction in the mat, fiber curvature, fiber interconnections, fiber fusion at contact points and mat porosity [25, 26]. Elastic modulus (E), stress at break (\( \sigma_b \)) and elongation at break (\( \varepsilon_b \)) of PBS, PBS/PBTDG, P(BS\(_{10}\)BTDG\(_{10}\)), PBS/PBDG, and P(BS\(_{11}\)BDG\(_{11}\)) are reported in Figure 2. Both the two blends and the copolymers were characterized by a lower elastic modulus than PBS and by a higher elongation at break (with the only exception of PBS/PBTDG). Moreover, E and \( \varepsilon_b \) were respectively lower and higher in the case of block copolymers compared to the corresponding blends. This behaviour could be due to the decrease in the amount of hard crystal phase when comparing the blend with the corresponding copolymer (Table 1). The presence of thioether bonds along the polymer backbone caused an increase of the elongation at
break and a decrease of the elastic modulus compared to the polymers containing ether linkages with the same molecular architecture. This result could be ascribed to the higher chain flexibility of BTDG units due to the presence along the polymeric chain of C-S bonds longer than the C-O ones and due to weaker interchain interactions because of the lower electronegativity of sulphur atoms compared to oxygen ones.

![Graph](image)

**Figure 2.** (A) Elastic modulus $E$; (B) stress at break $\sigma_b$ and (C) deformation to break $\varepsilon_b$ of the polymers under investigation.

2.3.2. *Hydrolytic degradation*

Hydrolytic degradation experiments were performed under physiological conditions on the electrospun mats in order to evaluate the effect of the following factors on the hydrolysis rate: (i) introduction of hydrophilic BDG and BTDG co-units in the PBS macromolecular chain; (ii) type of heteroatom (O or S) introduced; (iii) different molecular architecture.
Tailoring Chemical and Physical Properties of Fibrous Scaffolds from Block Copolyesters Containing Ether and Thio-ether Linkages for Skeletal Differentiation of Human Mesenchymal Stromal Cells

(blend or block copolymer). Hydrolytic degradation of these scaffolds can be interpreted on the basis of two different factors: crystallinity degree and surface hydrophilicity [27].

Figure 3. Hydrolytic degradation of polymeric scaffolds under investigation as a function of incubation time. (A) percentage weight loss; (B) percentage residual number molecular weight; (C) increase in BS mol% content; (D) normalized heat of fusion; (E) GPC traces of PBS/PBDG blend as a function of incubation time.

Comparative results were obtained for both P(BS<sub>11</sub>BGD<sub>11</sub>)-<i>m</i> and P(BS<sub>11</sub>BGD<sub>11</sub>)-<i>n</i> indicating that the fiber diameters considered in the present work did not affect degradation rate. Therefore, for this polymer, average results of the two mats were reported (Figure 3). After 162 days of incubation, PBS practically did not lose weight whereas polymer blends and block copolymers appreciably degraded. In particular, ether-oxygen containing polyesters showed a higher degradation rate with respect to sulphur containing ones. Moreover, Figure 3A shows that the weight loss of block copolymer is higher than the corresponding blend. The trend was particularly evident if we consider PBS/PBDG and P(BS<sub>11</sub>BGD<sub>11</sub>). On the contrary, only a small difference has been recorded in the case of PBS/PBTDG and P(BS<sub>10</sub>BGD<sub>10</sub>).

Overall, copolymers containing BDG sequences degraded faster than those containing BTDG ones because of the higher hydrophilicity of the former (Table 1). By comparing the blend with the corresponding copolymer, P(BS<sub>11</sub>BGD<sub>11</sub>) degraded to a higher extent than PBS/PBDG because of the lower crystallinity degree, while the two sulphur containing polymers underwent a similar degradation profile because of their comparable
crystallinity degree and hydrophilicity (Table 1). The percentage of residual number average molecular weight for PBS, P(BS<sub>11</sub>BDG<sub>11</sub>) and P(BS<sub>10</sub>BTDG<sub>10</sub>) (M<sub>n res%</sub>) is reported in Figure 3B as a function of incubation time. The three samples underwent a decrease of M<sub>n</sub> with time, although PBS did not show any weight loss in the time range investigated (Figure 3A). This is not surprising, taking into account that in the first stages of hydrolytic degradation a substantial decrease in molecular weight occurs even if weight losses are still negligible[28]. PBS decrease of M<sub>n</sub> was lower due to its high crystallinity degree, high crystal perfection and hydrophobicity, while in the case of copolymers the change of molecular weight was more evident and seemed to be affected by the type of polymer investigated: for P(BS<sub>11</sub>BDG<sub>11</sub>) the M<sub>n</sub> decrease was higher than that of P(BS<sub>10</sub>BTDG<sub>10</sub>). The calculation of M<sub>n</sub> was not possible for the two blends, because the GPC traces of the analysed polymers revealed a shoulder since the very beginning of the incubation period that turned into two partially overlapped peaks. With increasing of the incubation time, a second peak was detectable (Figure 3E). Therefore, to monitor the degradation process, the molecular weight of both peaks (M<sub>p,1</sub> and M<sub>p,2</sub>) was calculated (Table S2). For both blends the position of one peak slightly changed (M<sub>p,2</sub>), while M<sub>p,1</sub> remarkably decreased with the increase of incubation time. By considering the difference in repeating units hydrophilicity, we can reasonably assume that M<sub>p,1</sub> is related to either PBDG or PBT DG macromolecular chains that underwent a substantial hydrolytic degradation, while M<sub>p,2</sub> corresponds to PBS macromolecular chains, whose molecular weight decrease was significantly lower.

In order to gain a better understanding of the hydrolysis mechanism occurring in the polymers, <sup>1</sup>H-NMR measurements were performed on degraded retrieved samples. Figure 3C reports the content of BS units in mol%, at a certain time of degradation, normalized to the initial mol% BS (mol%BS<sub>0</sub>/mol%BS) as a function of degradation time. In all samples, an evident increase of BS content and a consequent decrease in the BDG and BTDG content was observed during degradation. The polymeric systems containing ether linkages (PBS/PBDG and P(BS<sub>11</sub>BDG<sub>11</sub>)) underwent a significant change of molar composition, more evident in the blend compared to the copolymer. On the other hand, PBS/PBT DG and P(BS<sub>10</sub>BTDG<sub>10</sub>), underwent a similar decrease of BTDG content. By comparing weight loss (Figure 3A) and chemical composition of the retrieved samples (Figure 3C), it is evident that samples undergoing higher weight loss concomitantly displayed a higher increase of BS units. Therefore, the weight loss was mainly due to the solubilisation of either BDG or BTDG chain fragments since ester cleavage preferentially
occurred on heteroatom-containing chain segments, which were easily solubilized in water due to their hydrophilic nature. Moreover, as expected, ether-oxygen-containing sequences demonstrated higher solubility in water with respect to sulphur-containing ones: the lower decrease in molecular weight and the contemporary higher weight loss of \( P(\text{BS}_{11}\text{BDG}_{11}) \) with respect to \( P(\text{BS}_{10}\text{BTDG}_{10}) \), indicated that longer BDG sequences were solubilized in water generating a more pronounced change in BS mol%.

A confirmation that the amorphous regions of a polymer degraded more quickly than the crystalline ones was obtained by analysing the degraded samples with DSC. All the calorimetric traces were found to be characterized by an endothermic peak associated with the fusion process of the crystalline portion of the material (similar to Figure S1). The corresponding heat of fusion, at a certain time of degradation, was normalized to the heat of fusion of the corresponding non-degraded sample \( \Delta H_t/\Delta H_0 \). In all the polymeric compositions, the normalized heat of fusion regularly increased with incubation time, with the exception of PBS, whose crystallinity degree was not significantly affected (Figure 3D). It is worth remembering that the increment of the crystalline/amorphous ratio can be also due to annealing, which occurs at a temperature comprised between \( T_g \) and \( T_m \) (i.e. 37 °C in our case). The increase of \( \Delta H_t/\Delta H_0 \) can, therefore, be considered as the sum of two contributions: (i) the higher degradation of the amorphous region with respect to crystalline one and (ii) annealing. In block copolymers, the annealing process plays a greater role than in the blends because of the lower crystallization rate of the former, as previously reported[12, 13]. As a matter of fact, the crystallization process becomes more difficult as the BS block became shorter due to the effect of the second block (either BDG or BTDG), which limited the transport of the BS blocks on the crystal surface and acted as defect during chain folding. In this respect, the more pronounced increase in the heat of fusion of \( P(\text{BS}_{10}\text{BTDG}_{10}) \) compared to PBS/PBTDG is not surprising, although they showed a comparable weight loss (Figure 3A). From a comparison between ether-oxygen-containing polymers and sulphur-containing ones, the higher increase in the heat of fusion of the former can be explained on the basis of their higher degradation rate.

2.3.3. Seeding hMSCs on scaffolds

To examine the performance of these new polyesters for skeletal tissue engineering, hMSCs were cultured on the samples for 28 days in BM, OM, CM, or MM. BM is a maintenance medium that does not trigger the osteogenic and chondrogenic differentiation of hMSCs, while CM is able to support the chondrogenic differentiation of
hMSCs [29, 30]. OM is a medium that supports the osteogenic differentiation of hMSCs and MM is a medium that also promotes the formation of a mineralized matrix[30, 31]. The bioactivity of hMSCs was assessed by monitoring the GAG production, cellular morphology, cellular metabolism, ALP activity, and relative osteogenic and chondrogenic gene expressions. Polystyrene tissue culture plates were used as a positive control (data not shown). To investigate for a possible influence of fiber diameter, scaffolds of the same chemistry but different fiber diameter were assessed between P(BS$_{11}$BDG$_{11}$)-n (430 ± 260 nm) and P(BS$_{11}$BDG$_{11}$)-m (890 ± 410 nm). To analyze the effect of the introduction of a co-monomer (either BDG or BTDG) in the PBS chain, scaffolds of similar fiber dimension but different chemical group were examined among PBS, P(BS$_{10}$BTDG$_{10}$) and P(BS$_{11}$BDG$_{11}$)-n. To investigate the effect of reactive blending, PBS/PBTDG and PBS/PBDG scaffolds were considered. Finally, to evaluate the effect of blending versus copolymerization, PBS/PBDG blend and P(BS$_{11}$BDG$_{11}$)-m, which had a comparable fiber dimension, were considered.

2.3.4. Cell morphology and metabolic activity

Cell attachment and morphology on scaffolds were examined by SEM (Figure S2, Figure S3 and Figure S4). After 28 days culture, it could be seen that the hMSCs migrated, attached and proliferated on all types of scaffolds in all investigated media. No significant difference in cell morphology was observed between P(BS$_{11}$BDG$_{11}$)-n and P(BS$_{11}$BDG$_{11}$)-m in all media, indicating that the fiber dimension considered in the present work did not affect cell morphology. In OM, hMSCs on PBS/PBTDG and P(BS$_{10}$BTDG$_{10}$) had a more spread morphology compared to PBS/PBDG and P(BS$_{11}$BDG$_{11}$)-n respectively, therefore revealing an effect of BTDG or BDG presence on cell morphology. However, in CM, hMSCs displayed morphology with more bi-polar or multipolar extensions on PBS/PBTDG scaffolds in comparison with PBS/PBDG. In addition, no difference in cell morphology was observed between PBS/PBDG blend and its corresponding block copolymer of P(BS$_{11}$BDG$_{11}$) in all media.

Cell metabolic activity on seeded scaffolds was evaluated at days 7 and 28 using a Presto Blue assay (Figure S5). Metabolic activity, which is indirectly related to cell viability, increased over the cell culture period in all media for all scaffolds. P(BS$_{11}$BDG$_{11}$)-n and P(BS$_{11}$BDG$_{11}$)-m had a similar metabolic activity cultured in both BM and CM through the culture time points. In OM, P(BS$_{11}$BDG$_{11}$)-n showed a significantly higher metabolic activity at day 28 compared with P(BS$_{11}$BDG$_{11}$)-m due to their difference in fiber
PBS constructs displayed a slightly higher metabolic activity compared to P(BS\textsubscript{10}BTDG\textsubscript{10}) in both BM and CM at day 28 due to the presence of co-units of BTDG. Similarly, the presence of BDG blocks along polymer chain did not enhance the cell metabolism in comparison with PBS under the same conditions. Interestingly, in the copolymers the introduction of BDG blocks generated a higher cell metabolism in CM compared to BTDG blocks, while in OM PBS/PBDG displayed higher cell metabolic activity compared to PBS/PBTDG.

2.3.5. ALP activity

To investigate early osteogenic differentiation of hMSCs cultured onto different scaffolds either in BM or in OM, we performed quantitative analysis of ALP at days 7 and 28 (Figure 4A and 4B). ALP is commonly known as an early marker of osteogenic differentiation, which is involved in the formation of calcium phosphate crystals [32]. The ALP activity increased during cell culture in BM and OM and showed no remarkable difference between scaffolds at day 7. After 28 days, P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-m} showed a similar ALP activity as P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-n} in OM. Shih et al. grew human hMSCs on electrospun type I collagen nanofibers[33]. They demonstrated that no significant difference of ALP activity of hMSCs after 12 days of culture in OM was observed between scaffolds with fiber diameter of 200-500 nm and fiber diameter of 500-1,000 nm. PBS, P(BS\textsubscript{10}BTDG\textsubscript{10}) and P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-n} scaffolds supported a similar ALP activity in both BM and OM, which indicated that the presence of block units of BTDG and BDG do not elicit a specific effect on ALP activity. We also observed that ALP activity in both BM and OM was comparable for both PBS/PBTDG and PBS/PBDG. This suggested that the introduction of a second component of the blend, being either PBTDG or PBDG, has a similar effect on ALP activity. It is worth mentioning that the ALP activity on P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-m} was higher than the corresponding blend in both investigated media.

The results of ALP activity were further confirmed by staining cells with ALP (Figure S6 and S7). Low levels of ALP staining were present on all types of scaffolds at day 7 in both media without significant difference. P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-n} showed similar density in ALP staining during the cell culture period in all media compared to P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-m}. At day 28, the intensity of ALP staining remarkably increased on PBS and P(BS\textsubscript{10}BTDG\textsubscript{10}) scaffolds in BM compared to P(BS\textsubscript{11}BDG\textsubscript{11})\textsubscript{-n}, suggesting a possible inductive effect on stem cell early osteogenic differentiation of these two chemical compositions. Furthermore, ALP staining increased in intensity on PBS, P(BS\textsubscript{10}TDG\textsubscript{10}), and
P(BS<sub>11</sub>BDG<sub>11</sub>)-n scaffolds in OM. Interestingly, PBS/PBDG and PBS/PBTDG scaffolds exhibited a constantly lower ALP activity at both time point in all media. In addition, P(BS<sub>11</sub>BDG<sub>11</sub>)-m showed a higher ALP staining in OM than the corresponding blend.

![Figure 4](image-url)

**Figure 4.** ALP activity of hMSCs cultured on scaffolds in BM (A) and OM (B) at days 7 and 28. The final results of ALP activity level was normalized by DNA amount. GAG production of hMSCs cultured on scaffolds in BM (C) and CM (D) at day 7 and day 28. *p<0.05, **p < 0.01, ***p < 0.001.

### 2.3.6 GAG assay

The quantification of GAG synthesis is shown in Figure 4C and 4D. No significant differences were found between P(BS<sub>11</sub>BDG<sub>11</sub>) and P(BS<sub>11</sub>BDG<sub>11</sub>)-n in BM. However, after 28 days, P(BS<sub>11</sub>BDG<sub>11</sub>)-m had a significant higher GAG deposition than P(BS<sub>11</sub>BDG<sub>11</sub>)-n in CM, due to their difference in fiber dimension. This results is consistent with the findings by Bean et al.[34], who demonstrated that seeding hMSCs onto microfibers at a higher cell density promotes GAG production better than nanofibers. In CM, at day 28, P(BS<sub>10</sub>BTDG<sub>10</sub>) scaffold presented a significant increase in GAG production compared to PBS and P(BS<sub>11</sub>BDG<sub>11</sub>)-n. In addition, PBS/PBTDG displayed a significantly higher GAG amount than PBS/PBDG in CM. Both these results may indicate that the systems containing BTDG units may have a better effect on production of GAG in CM with respect to those containing BDG units. This result could be due to the presence
of sulphur atoms along the polymer chain which are always observed in all of GAG[35].
P(BS$_{11}$BDG$_{11}$)-m highlighted a lower GAG production than the corresponding blend in all media, which is in contrast to ALP activity. A recent study showed that hMSCs cultured on PCL electrospun scaffolds cultured for 4 weeks in chondrogenic medium had a GAG/DNA concentration of about 15 $\mu$g/$\mu$g[36]. Compared to this study, our scaffolds showed a higher GAG production at the same time point, However, in doing such comparison with other scaffolds used in the literature, we may still be limited by the potentially different cell donors and related intrinsic variability between patients.

2.3.7. The mineralization of hMSCs

ARS has been used to examine the level of calcium deposition after 28 days (Figure 5). There were no apparently differences in ARS between scaffolds in BM. The intensity of ARS onto P(BS$_{11}$BDG$_{11}$)-n scaffolds was higher than both P(BS$_{10}$BTDG$_{10}$) and PBS scaffolds and no significant differences were found between P(BS$_{10}$BTDG$_{10}$) and PBS in MM. These results suggested that the introduction of BDG blocks supported better mineralization when compared to the introduction of BTDG blocks. No appreciable differences in ARS were observed between PBS/PBTDG and PBS/PBDG, as well as between PBS/PBDG and P(BS$_{11}$BDG$_{11}$)-m for all media. Xuejun et al. determined the cell viability, growth and differentiation of hMSCs in PLGA (PLA: PGA ratio of 85:15) nanofibers [37]. They demonstrated that hMSCs treated with osteogenic medium showed positive staining to alizarin red after 4 weeks. The intensity of alizarin red staining was lower than that of our scaffolds. However, as explained earlier for chondrogenic studies, different cell donors were used, which should be taken into consideration when comparing our results with the literature.
Figure 5. Mineralization of hMSCs on scaffolds was investigated after 28 days of culture in BM and MM. The production of calcium was stained with ARS (A). Scale bar = 1 mm. (B) Quantification of the calcium deposition per scaffold by ARS assay at day 28. *p<0.05, **p < 0.01, ***p < 0.001

2.3.8. Osteogenic gene expression

The investigation of osteoblast-related gene expression for hMSCs on scaffolds was performed at day 7 and day 28 (Figure 6). In both media investigated, P(BS11BDG11)-n and P(BS11BDG11)-m showed a similar osteogenic gene expression including ALP, Runt-related transcription factor 2 (RUNX2), Collagen type I alpha 1 (COL1A1) and Bone
sialoprotein (BSP), thus confirming that the difference in fiber dimension in the present work did not affect osteogenic differentiation. ALP expression for PBS, P(BS₁₀TDG₁₀) and P(BS₁₁BDG₁₁)-n scaffolds were almost the same in BM. However, ALP expression on PBS was higher than on P(BS₁₁BDG₁₁)-n scaffolds at day 28 in OM. RUNX2 strongly affects hMSCs osteogenic differentiation in the early stage, resulting in enhanced ALP, osteocalcin (OCN), osteopontin (OPN), and BSP expression [38]. The expression of RUNX2 on PBS was higher than that on P(BS₁₁BDG₁₁)-n scaffolds at both time points in BM. P(BS₁₀BTDG₁₀) displayed a significant higher expression of RUNX2 at day 7 in BM in comparison with P(BS₁₁BDG₁₁)-n. COL1A1 is associated with the formation of the ECM, being considered essential for the development of an osteoblastic phenotype[39]. PBS showed a significant higher COL1A1 expression than P(BS₁₁BDG₁₁)-n after 28 days in BM, although no significant differences were found in OM. BSP, a late marker of hMSCs osteogenic differentiation, is a highly sulphated and glycosylated phosphoprotein in bone matrix [40]. BSP expression on PBS was significantly higher than that on P(BS₁₁BDG₁₁)-n at day 28 in OM. All gene expression tested did not highlight significant differences between PBS/PBDG and P(BS₁₁BDG₁₁)-m. These results suggest that PBS scaffolds may elicit a higher early osteogenic differentiation than copolymers containing BDG or BTDG units.

![Figure 6. Quantitative RT-PCR gene expression analysis of osteoblast-related genes in BM and OM at days7 and 28. *p<0.05, **p < 0.01.](image-url)
2.3.9. Chondrogenic gene expression

To know the potential of chondrogenic differentiation of hMSCs on the scaffolds under investigation, we evaluated a panel of chondrocyte-related genes: collagen, type II alpha 1 (COL2A1), SOX9, aggrecan (ACAN) and activated leukocyte cell adhesion molecule (ALCAM) (Figure 7). No significant difference in expression of the investigated gene profiles was observed between P(BS\textsubscript{11}BDG\textsubscript{11})-n and P(BS\textsubscript{11}BDG\textsubscript{11})-m despite of their difference in fiber dimension. COL2A1 is a specific marker of hyaline cartilage ECM\cite{41}. P(BS\textsubscript{11}BDG\textsubscript{11}) and P(BS\textsubscript{10}BTDG\textsubscript{10}) scaffolds displayed a higher expression of COL2A1 than PBS scaffolds at day 28 in BM. It could be inferred that the introduction of BTDG and BDG blocks along the PBS macromolecular chain was favourable for COL2A1 expression. The expression of COL2A1 on PBS/PBTDG scaffold was consistently higher than that on PBS/PBDG scaffolds, indicating that the blend containing BTDG units better supported COL2A1 expression compared to BDG units. SOX9, an early marker for chondrogenesis, is the transcriptional activator of COL2A1, COL9A1 and ACAN\cite{42, 43}. The expression of SOX9 was similar for all scaffolds at both time points in BM, while the expression of this gene significantly increased onto PBS/PBTDG scaffold at day 28 in CM. ALCAM is expressed on human articular chondrocytes and function as a marker for stemness\cite{44}. At day 28, PBS/PBTDG scaffolds supported a significant higher expression of ALCAM in both BM and CM compared to PBS/PBDG scaffolds, thus suggesting that the blend containing BTDG units better supports ALCAM expression compared to BDG units. ACAN is one of the main proteoglycan in the articular cartilage matrix \cite{45}. In BM, ACAN on P(BS\textsubscript{10}TDG\textsubscript{10}) scaffold showed a significant up-regulation compared to PBS, due to the presence of BTDG units along the polymer chain. A significant higher expression of ACAN was observed on PBS/PBTDG scaffold compared to that on PBS/PBDG in both condition media at day 28. Together, gene expressions results indicated that BTDG units displayed better support of hMSCs chondrogenesis with respect to BDG units. This result could be ascribed to the presence of sulphur atoms in the BTDG units. It is also worth noticing that PBS/PBTDG blends showed better performances than the corresponding block copolymer.
Figure 7. Quantitative RT-PCR gene expression analysis of chondrogenesis-related genes in BM and CM at days 7 and 28. *p<0.05, **p < 0.01, ***p < 0.001.

2.4. Conclusions

In the present study, PBS-based scaffolds from block copolymers containing heteroatoms have been successfully prepared by electrospinning. Hydrolytic degradation studies under physiological conditions showed that the hydrolysis rate can be tuned by acting on the kind of heteroatom introduced and on the molecular architecture, as these factors deeply influence both the crystallinity and the surface wettability of the investigated materials. The presence of BTDG units in the copolymers demonstrated a better support of chondrogenic differentiation of hMSCs compared to the presence of BDG units. Additionally, PBS scaffolds induced enhanced early osteogenic differentiation of hMSCs and the additional presence of ether units (BDG) resulted in enhanced mineralization. It is also worth mentioning that the same chemical composition, but a different molecular architecture (i.e. blend or block copolymer) highlighted a very different effect on hMSCs cell function such as differentiation. Therefore, molecular architecture revealed as an important parameter in terms of modulating the chemistry of scaffold. Taken together, these results demonstrated that the new copolymer family may therefore find applications for the regeneration of skeletal tissues in regenerative medicine field. In the present work, we primarily focused on understanding cell-materials interactions in these new polymeric scaffolds, which is a fundamental required step in biomaterial science before continuing with more translational in vivo studies. In this respect, in vivo studies will be planned for future translational work.
References


[34] Bean AC, Tuan RS. Fiber diameter and seeding density influence chondrogenic differentiation of mesenchymal stem cells seeded on electrospun poly (ε-caprolactone) scaffolds. Biomedical Materials. 2015;10:015018.


Supporting Information

Table S1. Electrospinning operating conditions.

<table>
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<tr>
<th>Polymer</th>
<th>solvent mixture (v:v)</th>
<th>conc. (w/v%)</th>
<th>voltage (kV)</th>
<th>flow rate (ml/h)</th>
<th>D (cm)</th>
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<tr>
<td>P(BS₁₁BDG₁₁)</td>
<td>HFIP</td>
<td>25</td>
<td>23</td>
<td>0.5</td>
<td>15</td>
</tr>
</tbody>
</table>

<sup>a</sup> D = needle-to-collector distance; <sup>b</sup> "m" indicates micrometric fibers; <sup>c</sup> "n" indicates nanometric fibers.

Table S2. Peak molecular weight as a function of incubation time for PBS/PBDG and PBS/PBTDG.

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<th>PBS/PBTDG</th>
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Table S3. Sequences of primers used in real time RT-PCR.

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<th>Reverse primer 5’ to 3’</th>
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<td>GACTTGTCTTTTCAGCAAGGA</td>
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<tr>
<td>ALP</td>
<td>AACAAGCACTCCACACTTCATC</td>
<td>TTCAGCTGCTACTGCAATGTC</td>
</tr>
<tr>
<td>RUNX2</td>
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<tr>
<td>COL1A1</td>
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<td>CAGATCGTCATGCGACACAC</td>
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<tr>
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<tr>
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</tr>
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<td>ACAN</td>
<td>AGGCACCGTGATCCCTACC</td>
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<td>ALCAM</td>
<td>ACGATGAGGAGAGGAAGATAAGT</td>
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**Figure S1.** Calorimetric curves of the polymeric scaffolds under investigation.

**Figure S2.** SEM images of hMSCs cultured on scaffolds in BM at day 7 and day 28.
Tailoring Chemical and Physical Properties of Fibrous Scaffolds from Block Copolyesters Containing Ether and Thio-ether Linkages for Skeletal Differentiation of Human Mesenchymal Stromal Cells

**Figure S3.** SEM images of hMSCs cultured on scaffolds in OM at day 7 and day 28.

**Figure S4.** SEM images of hMSCs cultured on scaffolds in CM at day 7 and day 28.
Figure. S5. Metabolic activity of hMSCs cultured on scaffolds in BM(A), OM(B) and CM (C) at day 7 and day 28. *p<0.05, **p < 0.01 , ***p < 0.001.

Figure. S6. The staining of alkaline phosphatase enzyme activity for hMSCs on scaffolds in BM at day 7 and day 28. Scale bar =1mm.
Figure. S7. The staining of alkaline phosphatase enzyme activity for hMSCs on scaffolds in OM at day 7 and day 28. Scale bar =1mm
Flexible Yttrium-Stabilized Zirconia Nanofibers Offer Bioactive Cues for Osteogenic Differentiation of Human Mesenchymal Stromal Cells

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[+These authors contributed equally to this work.]
Abstract
Currently, the main drawback of ceramic scaffolds used in hard tissue regeneration is their low mechanical strength. Stabilized zirconia, especially the tetragonal 3 % yttrium-stabilized zirconia (YSZ) phase, has been considered as a bioinert ceramic material with high mechanical strength. In the present work, novel flexible nanofibrous YSZ scaffolds were prepared by electrospinning. The obtained scaffolds showed remarkable flexibility at the macroscopic scale, while retaining their stiffness at the microscopic scale. The surface nanoroughness of the scaffolds could be tailored by varying the heat treatment methods. Our results demonstrate the osteogenic differentiation and mineralization of seeded human mesenchymal stromal cells (hMSCs) were supported by the nanofibrous YSZ scaffolds, in contrast to the well-known bioinert behavior of bulk YSZ. These findings highlight that flexible ceramic scaffolds are an appealing alternative to the current brittle ceramics for bone regeneration applications.

Keywords:
Flexible ceramic scaffolds, electrospinning ceramic, surface roughness, human mesenchymal stromal cells, osteogenic differentiation
3.1. Introduction

Bioceramics have been widely used for several decades to repair bone damage[1-3], since they are promising materials due to their similarity to bone in terms of chemical properties[4-7]. Within the area of bioceramics, a clear distinction is made between bioinert and bioactive materials. Bioinert ceramics can be defined as those that do not bond with the surrounding tissues, nor induce a cell response. Examples of bioinert materials are alumina and yttria-stabilized zirconia (YSZ) [2, 3]. Conversely, bioactive ceramics are materials that can actively induce a specific cell response (e.g. cell differentiation) and, in some cases, are resorbable. These materials include hydroxyapatite (HA), β-calcium phosphates and bioactive glasses [1-3]. One of the major limitations of ceramic scaffolds for hard tissue regeneration is that they suffer from low mechanical strength, which may facilitate fragile failure when implanted [4, 6, 8]. Among the aforementioned bioceramic materials, zirconia has the highest mechanical strength, especially the 3% yttrium-stabilized tetragonal zirconia (YSZ) phase [9, 10].

The scaffold’s microstructure also plays a crucial role. Microporosity provides a large surface area for cell adhesion, thus improving the biological response [11]. However, the scaffolds’ mechanical properties are negatively affected [4, 7]. Porosity reduces the mechanical strength of materials [7]. Tailor-made design of the nanoarchitecture of porous biomaterials may help to overcome these issues [9, 12-14]. A recent study demonstrated a nanoarchitectured aluminosilicate ceramic with improved ductility and flexible recovery after compressive stresses [12]. Nanofiber mats of titania, silica and zirconia with enhanced flexibility have also been reported [13-20].

To fabricate scaffolds with fiber dimensions on the nano- to microscale mimicking the features of the natural extracellular matrix (ECM), electrospinning has been used broadly in the field of scaffold tissue engineering [21-23]. Cells do not only respond to chemical and physical stimuli from their neighboring cells, but also from the surrounding fluid and ECMs [24]. The characterization and control of surface properties, such as surface roughness, can therefore be used to optimize cell attachment, proliferation and differentiation.

In this article, we propose the fabrication of a flexible YSZ nanofiber material for bone tissue engineering. Recently, YSZ-based fibers have been fabricated by electrospinning for various applications, such as fuel cells [25] and filtration [18], but their application in the biological and biomedical field has not been explored to our knowledge. Herein, we introduce novel YSZ ceramic nanofibers in the field of regenerative medicine. The YSZ
fibers typically have lengths of the order of meters, instead of just a few millimetres as reported earlier [25], and a high gravimetric density. The sintering temperature that we employed to densify the ceramic nanofibers was much lower than in earlier studies [25], so that sintering between fibers was avoided, which contributed further to their flexibility and retention of mechanical properties upon cyclic deformation. Indeed, these YSZ ceramic nanofibers have remarkable flexibility compared to the traditional bioactive ceramics and their surface nanoroughnesses can be tailored for cellular response. Interestingly, in contrast to the well-known bioinert behavior of bulk YSZ, nanofiber meshes support in vitro osteogenic differentiation and mineralization of human mesenchymal stromal cells (hMSCs). This is the first study showing that flexible YSZ ceramic fibers with tailored surface nanoroughness can conduct osteogenic differentiation of hMSCs. The flexibility of the scaffold helps to overcome the problem of the fragile fracture that is characteristic for traditional bioceramic scaffolds. Therefore, it provides a complement to the currently available library of bioactive ceramic materials for bone tissue engineering.

3.2. Experimental procedures

3.2.1. Electrospinning

Zirconium n-propoxide (Alfa Aesar, Lancashire, U.K.) and yttrium acetate hexahydrate (Alfa Aesar, Lancashire, U.K.) were dissolved in n-propanol (Sigma Aldrich) in a 97 to 3 % molar ratio. Then, polyvinyl pyrrolidone (PVP) (M.W. 1.3 kg/mol, Sigma Aldrich, St. Louis, U.S.A.) was added to the solution and left stirring overnight to ensure complete dissolution of the PVP. The solution preparation was performed in a nitrogen atmosphere. Subsequently, the solution was placed into a syringe, which was later connected to Teflon tubing with a 0.8 mm needle and place in the needle holder of the electrospinning setup. The flow rate was set at 1 mL/h, the air gap was 20 cm, the voltage was 15 kV and the temperature and the relative humidity were set to 25 ºC and 30%, respectively.

3.2.2. Thermal annealing

The YSZ- PVP fibres were annealed to burn out the organic phase and to obtain a pure YSZ crystalline phase. The samples were annealed either in a conventional oven (CO) (Carbolite, Hope Valley, U.K.) or a microwave oven (MW) (Milestone MicroSynth, Shelton, U.S.A.) at different heating rates. The samples annealed in the CO were heated at 1 and 5 ºC/min to 850 ºC. The samples annealed in the MW were either heated at 5
Flexible Yttrium-Stabilized Zirconia Nanofibers Offer Bioactive Cues for Osteogenic Differentiation of Human Mesenchymal Stromal Cells

8°C/min to 850 °C, or by rapid thermal annealing (RTA). RTA consisted on placing the sample directly in a pre-heated MW at 850 °C, and after annealing the samples were cooled to room temperature without a temperature program.

3.2.3. Fiber morphology and diameter

The fiber morphology was studied by a Zeiss Merlin (Oberkochen, Germany) scanning electron microscope at high magnification. The fiber diameter was determined from low magnification SEM pictures. The imaging was done at 1.4 keV. Fifty measurements were performed per micrograph and per sample, respectively.

3.2.4. Phase identification and crystallite size

The crystallinity and phase purity of the samples were studied by X-ray diffraction in the 2θ range between 10 ° and 100 ° using a PANalytical X’Pert (Almelo, The Netherlands), with a step size of 0.015° and 8 s per step. The crystallite size was estimated using the Scherrer equation from the full width at half maximum of the (111) peak. The Scherrer equation is presented in equation 1.

\[ \tau = \frac{K \lambda}{\beta \cos \theta} \]  

Where \( \tau \) is the crystallite size, \( K \) is the dimensionless shape factor, \( \lambda \) is the X-ray wavelength, \( \beta \) the full width at half maximum of the peak and, \( \theta \) is the Bragg’s diffraction angle.

3.2.5. Macroscopic mechanical properties

The mechanical properties of the ceramic nanofiber mats (n=3) were measured by means of microindentation. A Piuma microindenter (Optics 11, Amsterdam, The Netherlands) equipped with a tip with a radius of 37.5 µm and a spring constant of 199 N/m was utilized. 25 points were scanned throughout 2500 µm2 for each sample.

3.2.6. Microscopic mechanical properties and surface morphology

The microscopic mechanical properties were measured by a 3-point bending test on a single nanofiber, as previously reported elsewhere [26, 27]. A polycarbonate substrate with 90 µm wide channels was used as a substrate to suspend the fibers. An atomic force microscope (Bruker Icon, Massachusetts, U.S.A.) equipped with a ScanAsyst-AIR (Bruker, Massachusetts, U.S.A.) tip was used to measure the force/displacement curves of the fiber. The tip had a spring constant of 0.4 N/m and a resonance frequency of 70
kHz. The apparent Young’s modulus was calculated with the Euler/Bernulli beam theory (Equation 2) using the average fiber diameter obtained from SEM.

\[ E^* = \left( \frac{E}{d} \right) \frac{l^3}{12\pi} \frac{1}{D^4} \] 

(2)

Here \( E^* \) is the apparent Young’s modulus, \( F/d \) is the slope of the force displacement curve, \( L \) is the gap distance and \( D \) is the fiber diameter. The fiber surfaces were also scanned using the same tip in tapping mode to obtain surface morphology data. The roughness values (Rq) were obtained by analyzing the AFM topography data using the Gwyddion software package (version 2.31).

3.2.7. Fatigue test

A three-point bending fatigue test was performed on the ceramic nanofiber mats. This version of the 3-point bending test was especially designed for the experiment, and consisted of (1) a plate with a well-defined gap, on which the sample was suspended on, and (2) a knife, which applied a force to deflect the sample. The gap was 2 mm wide and the knife was 0.5 mm thick. The width of the sample was kept constant at 10 mm. The plate and knife were set on a rheometer (Anton Paar Physica MCR 501, Graz, Austria), which measured the force displacement curves at a speed of 50 \( \mu \)m/s.

The sample’s deflection was calculated assuming a trigonometric system between the sample, knife, and gap. Thus, if the displacement of the knife is known, the deflection angle may be calculated using half of the gap distance. This is exemplified in Supplementary Figure 8.

3.2.8. Cell Culture

Pre-selected hMSCs (donor No. 8001L) were isolated from human bone marrow by the Institute of Regenerative Medicine (Texas A&M University, Temple, Texas) which provide standardized preparations of MSCs for research use under the auspices of an NIH/NCRR grant (P40 RR 17447-06) [28, 29]. Briefly, mononuclear cells were separated using density centrifugation and plated to obtain adherent hMSCs. The cells were harvested around a confluence of 60–80%, after which they were considered as passage zero (P0). The P0 cells were further expanded and harvested to obtain passage 1 (P1). To obtain sufficient cells for scaffold-based experiments, hMSCs (P1) were expanded in T-300 flasks at a density of \( 3\times10^5 \) cells/cm\(^2\) and cultured in BM, consisting of \( \alpha \)-MEM (Gibco), 10% fetal bovine serum (Lonza), 2 mM L-glutamine (Gibco), 0.2 mM ascorbic acid (Sigma), 100 U/mL penicillin and 100 mg/ mL streptomycin (Gibco). The cells were
harvested at approximately 80-90% confluence for further sub-cultures.

3.2.9. Cell seeding on scaffolds

The ceramic scaffolds were cleaned using oxygen plasma cleaning (Harrick Plasma PDC-002) for 10 min at high power and subsequently punched out into discs to fit inside a 24-well plate. Rubber O-rings (Eriks B.V., The Netherlands) were inserted into the well to hold the substrate on the bottom. The ceramic scaffolds were sterilized in 70% (v/v) ethanol for 15 min each time for 3 times. After sterilization, the scaffolds were rinsed with PBS for 5 min each time for 3 times and immersed in BM at 37 °C overnight before cell seeding. Ceramic scaffolds were seeded at a density of 2×10^4 cells/cm^2 and cultured in an incubator with 5% CO_2 humid atmosphere at 37 °C for up to 14 days in OM, and in both BM and MM for 28 days. BM is considered a normal growth medium, which cannot induce osteogenic differentiation nor form a mineralized matrix[30, 31]. OM consisted of BM supplemented with 10^{-8} M dexamethasone to induce osteogenic differentiation of hMSCs and the MM consisted of OM supplemented with β-glycerophosphate to support the osteogenic differentiation[30, 31], and promote the formation of a mineralized matrix. The culture medium was refreshed every 2 days.

3.2.10. Live and dead assay

To determine the cell viability on scaffolds, a LIVE/DEAD® assay (Life Technologies) was performed. After washing the scaffolds twice with PBS, they were incubated for 30 min with a mixture of PBS solution containing ethidium homodimer (red fluorescent) / calceinAM (green fluorescent) at room temperature following the manufacturer’s protocol. Scaffolds were washed three times with PBS before imaging with a fluorescence microscope (Nikon, Eclipse E600 and Nikon DS-Flic camera).

3.2.11. SEM analysis

After 3 days of culture, the medium was removed, and the scaffolds were washed two times with PBS and fixed in a 10% formalin solution for 2 h. After rinsing with PBS, the scaffolds were dehydrated with ethanol series (50%, 70%, 80%, 90%, 96%, and 100%). The scaffolds were further processed by critical point drying (Balzers CPD-030). The dried samples were sputtered with gold (Cressington) and imaged with a XL-30 ESEM-FEG SEM (FEI, Eindhoven, Netherlands).
3.2.12. Fluorescence microscopy

On day 3, samples for phalloidin staining were fixed with 4% (v/v) paraformaldehyde (Sigma-Aldrich) in PBS, left overnight at 4 °C and subsequently washed 3 times with PBS. Afterwards, the cell membrane was permeabilized with 0.5% TritonX-100 solution for 10 min at 4 °C and washed with PBS again. A 1% BSA in PBS solution was added for 2 h to block non-specific binding. After withdrawing the BSA solution, phalloidin (Alexa Fluor 488, Invitrogen; Excitation/Emission:495/518 nm) with a dilution (1:40) in 1% BSA/PBS was applied for 25 min at room temperature (RT) in the dark and rinsed 3 times with PBS. For nuclei staining, DAPI (1:100) in PBS was applied for 15 min at RT in the dark. The samples were observed under a fluorescence microscope (Eclipse E600, Nikon) equipped with proper filter. The average surface area for single cells was determined by the software CellProfiler 2.1.1(Broad Institute, Cambridge, USA). The surface area in each of at least 10 cells within an image for a total of six images per sample was quantified. All these images were from the same experiment.

3.2.13. Cell metabolic activity

The cell metabolic activity on day 7 and day 14 was assessed using a PrestoBlue™ Cell Viability Reagent (Life technologies), according to the manufacturer’s protocol. Briefly, a cultured medium was replaced with a medium containing 10% (v/v) PrestoBlue reagent and cells were incubated for 1 h at 37 °C. Fluorescence was measured at 590 nm on a Perkin Elmer Victor3 1420 multiple plate reader. Three separate experiments (n = 3) and two replicates each were performed.

3.2.14. ALP activity

The relative ALP activity was measured by chemoluminescence with a CDP-star® kit (Roche Life Science), according to the manufacturer’s instruction. Briefly, cell-scaffold constructs were washed 3 times with PBS and stored at -80 °C for at least 24 h. After thawing, the constructs were treated with 250 µL cell lysis buffer at a pH of 7.8 (composed of 0.1M KH₂PO₄, 0.1M K₂HPO₄ and 0.1% Triton X-100) for 1 h at RT. Cell lysates (50 µL) were transferred to Eppendorf tubes and stored at -20 °C for ALP assay. The remainder of the remaining cell lysates (200 µL) was used for DNA assay. For ALP analysis, 10 µL of cell lysates were transferred in duplicate to a well of a 96-well plate. Subsequently, 40 µL of CDP-star were added and plates were incubated for 15 min at room temperature. The plates were measured on a Multilabel reader (Victor3 1420,
Perkin Elmer, USA). Three separate experiments (n = 3) and two replicates each were performed. The results of ALP activity were presented after normalization to DNA quantity per scaffold.

3.2.15. Quantification of DNA amount

Cell lysates remainder, (200 µL) as mentioned above, were further digested for 16 h at 56°C with 200 µL Tris/EDTA buffer containing 1 mg/mL proteinase K (Sigma Aldrich), 185 µg/mL iodoacetamine (Sigma Aldrich) and 10 µg/mL Pepstatin A (Sigma Aldrich). The quantification of the total amount of DNA was performed using the CyQuant® DNA assay (Molecular Probes, Eugene, USA) according to the manufacturer’s manual. Briefly, a 40 µL aliquot of lysate was pipetted in duplicate into a black 96-well plate, followed by addition of 40 µL RNase (1000×) in NaCl-EDTA buffer composed of component B (20×) and incubated for 1 h at RT. Subsequently, 80 µL of Gr-dye solution was added and incubated for 15 min. The measurements were performed on a spectrophotometer reader (Victor3 1420, Perkin Elmer, USA) at an excitation wavelength of 480 nm and an emission wavelength of 520 nm. Three separate experiments (n = 3) and two replicates each were performed.

3.2.16. Gene expression analysis

Total RNA was isolated from samples at day 7 and 14 using a combination of TRizol with a NucleoSpinRNAII isolation kit (Bioke). Briefly, the scaffolds were washed three times with PBS and 1 mL of TRizol was added for each sample. Then, 0.2 mL of chloroform was added and the samples were centrifuged for 15 min at 12,000 RCF (relative centrifugal force). The aqueous phase containing RNA was transferred to a new Eppendorf® tube, mixed with an equal volume of 75% (v/v) ethanol and loaded onto an RNA binding column of the isolation kit. After a series of purification procedures, the RNA was re-suspended in RNASE-free water. RNA concentration and purity were measured by using an ND 1000 spectrophotometer (Nanodrop Technologies, USA). Finally, cDNA was synthesized using iScript (Bio-Rad, Veenendaal, The Netherlands) according to the manufacturer’s protocol. Quantitative PCR was performed using SYBR-green supermix (Bio-Rad). Four separate experiments (n = 4) were performed. Cycling parameters for quantitative PCR were as follows: initial denaturation for 10 min at 95 °C, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. The sequences of the analyzed primers are
shown in Table 1. Cycle threshold (CT) values were normalized to the housekeeping gene B2M, and the results of gene expression were calculated as $2^{-\Delta CT}$.

3.2.17. Alizarin red S staining and quantification

After 28 days of culture, samples were washed with PBS, fixed with 10% formalin for 1 h, and then subsequently with PBS and twice with distilled water. A freshly filtered 2% aqueous alizarin red S solution (pH 4.2) was applied to scaffolds for 2 min, after which the excess solution was washed off with distilled water. The quantification of orange-red coloration of alizarin red S was carried out as in previous studies[32, 33]. Briefly, constructs were transferred to a microcentrifuge tube before removing excess water. A 10% acetic acid solution was added and samples were shaken for 30 min at RT. After vortexing for 30 s, the solution was centrifuged for 15 min at 20,000 RCF and the supernatant phase was removed to a new 1.5 mL microcentrifuge tube. A 10% ammonium hydroxide solution was added drop-wise to neutralize the acid. Aliquots of the supernatant phase were measured in triplicate at 405 nm in a 96-well plate using opaque-walled, transparent-bottomed plates (Fisher Lifesciences). Three separate experiments (n = 3) and two replicates each were performed. Scaffolds without seeding cells were used as control.

### Table 1. Sequences of primers used in PCR.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer 5' to 3'</th>
<th>Reverse primer 5' to 3'</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMP-2</td>
<td>ACTACCAGAAGAGTGGGAA</td>
<td>GCATCTGGTTCTGGGAACCT</td>
</tr>
<tr>
<td>BSP</td>
<td>CCCCACCTTTTGGAACCA</td>
<td>TCCCCGTTCCTACCTACTGAT</td>
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<tr>
<td>OPN</td>
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</tr>
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<td>CGCCTGGGTCTCTCCTAC</td>
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<tr>
<td>ALP</td>
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</tr>
<tr>
<td>B2M</td>
<td>ACAAGTCACATGGTTCCA</td>
<td>GACTTGTCTTTAGCAGAAGA</td>
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</tbody>
</table>
3.2.18. Statistics

All data points are expressed as mean values ± standard deviation (S.D.). Statistical analysis was carried out using GraphPad Prism 5.01 (GraphPad software, San Diego, USA) for windows. The statistical differences were determined by using two-way analysis of variance (ANOVA), followed by a Tukey’s post-hoc test unless otherwise mentioned. A value of p < 0.05 was considered statistically significant.

3.3. Results and discussion

The ceramic nanofibers were fabricated by electrospinning with a mixture of a YSZ precursor and polyvinylpyrrolidone (PVP). Subsequently, the resulting fiber mats were annealed by means of conventional or microwave-assisted heating in static air at 850 °C for 2 h using different heating rates (i.e. to burn out the organic phases and for crystalline phase formation). The obtained nanofibers had a diameter of 530 ± 120 nm (Figure S1), and the fiber mats had a thickness of approximately 120 µm. A scanning electron microscopy (SEM) image of the material is shown in Figure 1A. The YSZ nanofiber mats exhibited extraordinary flexibility not often demonstrated for pure ceramic materials. It behaved as a paper-like material (Figure 1B and Video S1). The material could achieve total deflection (180° bending) without mechanical failure. Moreover, the deflection could be cycled at least 1000 times without breaking the material when a 3-point bending test was performed. To the best of our knowledge, the fabrication of ceramic nanofiber mats with such high flexibility and good fatigue behavior has not been reported until now. The flexibility of electrospun silica-based mats has been attributed to the amorphous nature of the material, which is therefore more robust than crystalline materials [14, 34]. However, that hypothesis does not explain the behavior of the crystalline YSZ nanofibers reported here (X-ray diffractograms showing crystalline nature are shown in Figure S2. Park et al hypothesized that the flexibility of crystalline TiO₂ was the result of the physical entanglement as single nanofibers are brittle [13].
Figure 1. The flexibility of YSZ nanofiber mats. (A) SEM images of a YSZ nanofiber mat, Scale bar is 20 µm. (B) Pictures of the YSZ mat showing its flexibility and fire resistance ceramic nature. (C) Mechanical properties of a nanofiber mat by microindentation (at macroscopic scale), a single nanofiber by 3 point bending using AFM (at microscopic scale), and bulk YSZ (taken from literature [27-29]). (D) Schematic overview of a 3 point bending test on a single nanofiber using AFM. (E) Schematic illustration of a 3 point bending fatigue test on the nanofiber mat. (F) Representative histogram of 200 cycles of the 3 point bending fatigue test on a YSZ nanofiber mat in which the force was monitored as a function of time. Error bars represent standard deviation of the mean. The statistical significance was determined using a one way ANOVA test, where ***p < 0.001.

However, we observed that also single nanofibers were able to withstand a high degree of deflection (Video S2). Probably due to the long length of the nanofibers, the mechanical energy could be dissipated under the elastic deflection. The coherence of fibers mats composed of long fibers is better than that of fiber mats composed of an assembly of fiber fragments. Moreover, the sintering temperature of 850 °C that we employed is much lower than in earlier reports on YSZ nanofibers where a temperature of 1500 °C was used [25]. As a result, our YSZ fibers densified but did not sinter together, which contributes to their freedom of movement and thus to their flexibility, and makes them less susceptible to fatigue-like phenomena. The flexibility of the mat was further quantified by microindentation tests. The Young’s modulus of the material was 1.11 ±
0.24 MPa, more than 5 orders of magnitude lower than that of bulk YSZ (Figure 1C), which is in the range of 178 - 222 GPa according to literature data [9, 35-37]. We also measured the mechanical properties of single nanofibers by a 3-point bending test using atomic force microscope (AFM) (Figure 1D). The measured apparent Young’s modulus of a single fiber was 30 ± 3 GPa, and a high deflection angle could be achieved. The apparent Young’s modulus of single fibers was several orders of magnitude higher than the macroscopic mat, but still a factor of 7 lower than bulk YSZ (Figure 1C).

![Figure 2. Plot of Young's Modulus vs density. The nanofiber mat and single nanofiber are compared with other traditional materials.](image)

This result is in accordance with other studies on electrospun ceramic nanofibers, which report similar decreases in Young’s modulus when measured via a three point bending test using AFM [26, 27]. The diameter and mechanical properties of the nanofibers at the different length scales did not vary significantly with the annealing treatment (Figure S3).
A special set-up was designed to perform a 3-point bending fatigue test on the mat (Figure 1E). The experiment consisted of 200 cycles of 45° deflection, while the applied force was monitored (Figure 1F). The sample did not break, and to our surprise we observed a slight increase of the force necessary to deflect the sample. Chen et al. reported slipping of individual nanofibers under tensile stress[18]. It may be possible that such a phenomenon occurs during deflection, gradually leading to a rearrangement of the fibers with an increasing number of cycles, ultimately resulting in a slightly stiffer material.

Moreover, the nanofibers mat offered a very low density, in the range of 0.06-0.09 g/cm³. Based on SEM observations, i.e. the absence of pores and grain boundaries, each individual fiber can be considered as nearly dense. The Young’s modulus of the nanofiber mats and individual fibers were plotted vs. the density as shown in Figure 2. For comparison, the values of other traditional materials were also plotted. It can be seen that the ceramic nanofibers mat behaved like a foam in terms of stiffness and density. This means that it is lighter and more flexible than other polymers, ceramics and porous ceramics. Nevertheless, individual nanofibers offer stiffness values close to ceramics and typical of porous ceramics or metals, 5 orders of magnitude stiffer than the mat.

The surface morphology and grain size of the fibers were modified by annealing the samples under different conditions. The green fiber mats were annealed in a CO at 1-5 °C/min heating rate, and in a MW at 5°C/min or by RTA (i.e. thermal shock). RTA is a method in which the MW is preheated to 850 ºC prior to placing the sample in the oven. The samples annealed in the CO showed rougher surface morphologies than the samples annealed in the MW even at the same heating rate, as shown by SEM (Figure 3A) and AFM (Figure 3B) analyses. This was further confirmed by the statistical analysis performed on the roughness data (Figure 3C). However, varying the heating rate did not result in significant differences among samples that were annealed by the same heating mechanism. The samples annealed in the CO at 1 °C/min and 5 °C/min had roughnesses of 0.8 ± 0.3 nm and 0.7 ± 0.2 nm, respectively. Both samples annealed in the MW had a roughness of 0.4 ± 0.1 nm. The heating mechanism had a significant influence on surface roughness. Samples annealed at 5 °C/min in a CO had a rougher surface than the samples annealed in a MW at the same heating rate (0.7 ± 0.2 nm vs 0.4 ± 0.1 nm). These results were consistent with the surface morphologies in SEM images (Figure 3A).
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Figure 3. The effect of annealing procedure on the fiber surface morphology. (A) SEM images of YSZ nanofibers, the scale bar is 200 nm. (B) AFM images of YSZ nanofibers, the scale bar is 100 nm. (C) Roughness data extracted from AFM scans. (D) Grain sizes of the fibers calculated from the FWHM of the (111) peak of the XRD patterns. Error bars represent standard deviation of the mean. The statistical significance was determined using a one way ANOVA test, where **p < 0.01 , ***p < 0.001.

However, it is noted that the roughness with AFM is measured in the out-of-plane direction, whereas the SEM images shows in-plane features of surface morphology. Different grain sizes were observed for each heating rate (Figure 3D) as calculated from the full width at half maximum (FWHM) in the XRD patterns (Figure S2). Samples annealed in the CO at 1 °C/min had the largest grain size, 24 ± 1 nm. Samples annealed in the CO and MW at 5 °C/min had intermediate grain sizes of 22 ± 1 nm and 18 ± 1 nm, respectively. The sample annealed in the MW using RTA had a much smaller grain size of 9.5 ± 0.1 nm. The samples annealed using lower heating rates showed larger grain sizes, irrespective of the heating mechanism. This can be attributed to the longer effective sintering times at high temperature under such conditions. The heating mechanism also affected the grain size. The samples annealed at 5 °C/min in a CO had a larger grain size than the samples annealed in a MW at the same heating rate. YSZ has previously been
reported to exhibit enhanced sintering behavior under microwave radiation heating\cite{38, 39}. Xie et al. reported smaller and more uniform grain sizes when YSZ was sintered in a MW, which is in accordance with our data (Figure 3D) \cite{39}. The YSZ nanofibers presented here have a different surface nanostructure than any previously reported YSZ topography. They are nearly 1-dimensional objects in a porous matrix, which affects the heat transport and crystallization rates. Nevertheless, the smoothening of the surface, when annealed using a MW, can be attributed to a step-bunching effect (i.e. aggregation of the ceramic steps) \cite{40}, sudden shrinkage and densification \cite{38}, and/or improved oxygen transport \cite{41}.

To elucidate the performance of our nanofibrillar YSZ in bone tissue engineering, hMSCs were grown on samples annealed at 5 °C/min using both CO and MW heating. We chose these samples as they have the same temperature profile during annealing, grain sizes in the same order of magnitude and similar fiber diameter distributions, yet different surface roughness. Recent studies revealed the importance of nanoscale and submicron-sized topographic cues to modulate fundamental cell behavior, including proliferation, migration, and differentiation \cite{42-44}. We hypothesized, therefore, that the nanofibrous structure and the different surface roughness of YSZ scaffolds may confer bioactive cues to an otherwise biologically inert ceramic material. Such topographical cues would ultimately result in varying cell morphology and, consequently, in different cell activities. The bioactivity of hMSCs on ceramic scaffolds was investigated by monitoring the cell viability, cellular morphology, cellular metabolism, alkaline phosphatase (ALP) activity, and gene expression in basic culture medium (BM) and osteogenic culture medium (OM) at days 7 and 14. Polystyrene tissue culture plates were used as a positive control (data not shown). The production of mineralized matrix was investigated after culturing cells-scaffold constructs in mineralization culture medium (MM) up to 28 days. Here, different culture media were chosen to investigate the influence of YSZ scaffolds on the osteogenesis potential of hMSCs in the presence or absence of osteoinducting soluble factors. BM is a maintenance medium that does not trigger the osteogenic differentiation of hMSCs \cite{30, 45}. On the other hand, OM induces osteogenic differentiation of hMSCs, whereas MM supports osteogenic differentiation and promotes the formation of mineralized matrix \cite{30, 31}.

To evaluate the viability of hMSCs on YSZ nanofibrillar scaffolds, a Live/Dead assay\textsuperscript{®} was performed after 3 days of culture. The fluorescence microscopy images of both live and dead cells on scaffold are shown in Figure S4. All investigated scaffolds showed a
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high rate of viable cells, which indicates no cytotoxic effects derived from the developed YSZ nanofibrillar scaffolds.

Figure 4. Characterization of cell morphology and actin cytoskeleton organization of hMSCs after 3 days of culture in BM. Representative SEM images of hMSCs on (A) microwave annealed (MW 5°C/min) nanofiber scaffolds, (B) conventional oven annealed (CO 5°C/min) nanofiber scaffolds, and (C) YSZ bulk disk scaffolds. Representative images of actin cytoskeleton organization of hMSCs on (D) microwave annealed (MW 5°C/min) scaffolds, (E) conventional oven annealed (CO 5°C/min) scaffolds, and (F) YSZ bulk disk scaffolds (actin filaments labeled with phalloidin are shown in green and nuclei labeled with DAPI are shown in blue). (G) Quantification of projected cell area on the investigated scaffolds. The statistical analysis was determined using two-tailed Student’s t-test and *p<0.05 was considered as significant difference. The scale bar in (A-C) and (D-F) are 10 µm and 50 µm respectively.

The cell morphology on YSZ scaffolds was investigated by SEM (Figure 4A-C). The cell cytoskeleton was further assessed by staining with phalloidin for actin filaments (Figure 4D-F), and the average surface area for single cell was quantified (Figure 4G). Cells on CO annealed scaffolds had a higher cell area compared to MW annealed
scaffolds, which could be correlated to the difference in surface roughness of the nanofibers. When comparing to that on bulk YSZ (having a roughness of $19 \pm 0.94$ nm), no significant difference in the average cell area was found possibly due to the combined effect of surface roughness and nanoscale fibrous structure.

Cell metabolic activity, which is indirectly related to cell viability, was determined using a PrestoBlue assay. The metabolic activity of hMSCs showed a similar profile in BM and OM (Figure S5). Bulk YSZ scaffolds had a significantly higher metabolic activity than all nanofibrous scaffolds, which could be due to the higher differentiation of hMSCs on nanofibrous scaffolds compared to bulk YSZ. Similar phenomena have been reported previously by Yuejun and his colleagues [46]. They found that hMSCs had higher metabolic activity on stiffer PDMS than the soft one while a higher differentiation activity of hMSCs were observed on soft PDMS compared to stiffer one. In addition, no significantly different metabolic activity was observed between CO and MW annealed scaffolds. This indicates that the surface roughness differences did not affect the cellular metabolism in the present work.

Polymerase chain reaction (PCR) was used to quantify the expression of a panel of osteogenic genes (Figure 5C and Figure S6). Gene expression of Runt-related transcription factor 2 (RUNX2) was significantly higher for CO and MW annealed scaffolds compared to bulk YSZ in both culture media. RUNX2 is considered as a focal point for integration of a variety of signals affecting the differentiation process of hMSCs into osteogenesis in the early stage, because it stimulates other downstream osteo-related genes such as osteopontin (OPN), osteocalcin (OCN), bone sialoprotein (BSP), and type I collagen [32]. Collagen type I alpha 1 (COL1A1) is fundamental for the development of the bone cell phenotype. It is actively expressed in the first proliferation period and then gradually down-regulated during subsequent osteoblast differentiation [47]. hMSCs cultured on annealed nanofibrous scaffolds showed higher expression of COL1A1 compared to bulk YSZ disks in both culture media. This is also associated with a significantly higher up-regulation of OCN and BSP on the nanofibrillar scaffolds in BM compared with bulk YSZ disks (Figure S6). Furthermore, the rougher meshes supported further enhanced OCN and BSP gene expression than the smoother ones, thus suggesting the possible role of surface topography on inducing hMSCs osteogenic differentiation, as also supported by other studies [48, 49]. OPN was only significantly up-regulated on nanofibrillar scaffolds at day 7 in OM, while no beneficial
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effect of surface topography or nanofibrillar cues were observed for OCN or BSP in OM both at day 7 and at day 14.

![Figure 5](image)

**Figure 5.** ALP activity (A and B) and gene expression analysis (C) from hMSCs cultured on conventional oven annealed (CO 5°C/min), microwave annealed (MW 5°C/min) and YSZ bulk disk scaffolds. ALP activity of hMSCs cultured in BM (A) and OM (B) at day 7 and 14 (the final ALP activity level was shown after normalizing to DNA amount). (C) Gene expression analysis for COL1A1, BMP2, and RUNX2 at day 7 and 14 in BM and OM (the final gene expression level was presented after normalizing to day 7 bulk YSZ). Error bars represent standard deviation of the mean. *p<0.05, **p < 0.01, ***p < 0.001.

Since one of the route for hMSCs differentiation towards the osteogenic lineage passes through endochondral development [50, 51], the down-regulation of OCN and BSP on nanofibrous scaffolds in comparison with their bulk disks could be also attributed to a higher hypertrophic differentiation of hMSCs, and therefore a less mature osteogenic
differentiation, on bulk disks due to their smooth surface properties in combination of medium stimuli. The smooth surface properties will affect the cell-to-material adhesion force and could have consequently influenced cell fate as well [52]. We also observed the expression of bone morphogenetic protein 2 (BMP2), which is known to be stimulating hMSCs along the osteogenic lineage [53]. BMP2 had a higher expression in all tested media on both nanofibrillar scaffolds compared to bulk YSZ except for day 7 in OM. Altogether, the ceramic nanofibrous scaffolds supported the osteogenic gene expression in both BM and OM, thus suggesting a possible role of the nanofibrillar cues inherently built in the YSZ meshes in contributing to hMSCs osteogenic differentiation. In addition, the varying surface roughness resulting from the different fiber annealing processes showed a net beneficial effect in case of the OCN and BSP genes, which occurred only for the rougher nanofibrous scaffolds obtained by employing conventional heating. It is also worth to mention that there were different trends in OM compared to BM, which might suggest that the differentiation of hMSCs in the present study was not only due to surface roughness but also other physical stimuli cues such as surface topography, porosity and physical cues associated to the nanoscale nature of the fibrillar meshes. Further studies should aim at clarifying the specific role of each of these material properties on cell differentiation, and the consequent osteoinductive nature of such nanofibrillar scaffolds, in addition to the osteoconductive properties here observed, both in vitro and in vivo.

At a protein level, the up-regulation of ALP activity is a common marker considered for the early stage of osteogenesis [32]. In BM, no significant differences were observed in ALP activity (Figure 5A). On the other hand, the ALP activity increased between day 7 and day 14 in OM and was significantly enhanced for CO annealed scaffolds compared to both MW annealed scaffolds and bulk YSZ (Figure 5B). These results indicate that the topographical properties of nanofibers, including larger surface area, higher porosity, and fibrous morphology, play a positive role in the osteogenic process of hMSCs. Similar results to our findings have been reported by Hae-Won Kim and colleagues who used bone marrow-derived stem cells to investigate the osteogenic potential of newly developed glass fibers [54]. Their results demonstrated that bioactive glass nanofibers showed a stronger ALP staining with respect to glass disks.
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Figure 6. Calcium mineralization on conventional oven annealed (CO 5°C/min), microwave annealed (MW 5°C/min), and YSZ bulk disk scaffolds. (A) The formation of mineralized extracellular matrix was stained with alizarin red S after 28 days of culture in BM and MM (the scale bar represents 1 mm). (B) Quantitative analysis of the mineral deposits per scaffold by alizarin red S staining at day 28. Error bars represent standard deviation of the mean. ***p < 0.001.

The ultimate deposition of a calcium-containing bone-like mineralized matrix is an index of extended osteogenic maturation of hMSCs. Mineralization was stained with alizarin red S after 28 days of culture (Figure 6 and Figure S7). When hMSCs were cultured on scaffolds in BM, little mineralized ECM was produced and no apparent differences were observed among the different scaffolds. In MM, a robust mineralized matrix was observed on bulk YSZ discs, while a strikingly significant increase in the amount of mineralized matrix was observed on the nanofibrous scaffolds. Quantification of the calcium deposits corroborated the staining results, showing significantly enhanced
calcium deposition on both fiber scaffolds compared to the bulk YSZ discs (Figure 6B). Here, the rougher CO annealed scaffolds resulted in a further enhancement of calcium content compared to the MW annealed scaffolds. This results is similar to the finding by Takeuchi and his co-workers, who demonstrated that osteoblast show a higher rate of matrix mineralization on rougher titanium surfaces [55]. A recent study showed that hMSCs cultured on calcium phosphate cements had a mineral concentration of about 1 mM per scaffold at day 21[56]. Compared to this study, our nanofibers annealed in a conventional oven had almost 3 folds higher mineral concentration. However, when comparing to other studies, care should be taken as different cell donors, time point and cell seeding density have been used.

3.4. Conclusions

In summary, new flexible nanofibrous YSZ scaffolds that display multiscale mechanical properties were successfully fabricated by electrospinning. Their surface roughness can be tailored by employing different heat treatment method. Our results demonstrate the osteogenic differentiation and mineralization of seeded human mesenchymal stromal cells was supported by the nanofibrous structure of YSZ scaffolds, in contrast to the well-known bioinert behavior of bulk YSZ. These flexible scaffolds can therefore present an appealing alternative to current brittle ceramics for bone regeneration applications.
References


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Supplementary Information

Figure S1. Fiber diameter distribution. Frequency distribution of fiber diameter as calculated from the SEM pictures.

Figure S2. XRD patterns of the nanofiber mats for different annealing conditions. The XRD patterns match the crystal structure of 3% mol yttria stabilized zirconia (ICDB 01-070-4426). CO = conventional oven; MW = microwave oven; MW RTA = rapid thermal annealing in microwave oven. It can be seen that the crystallinity of the samples is the same within experimental error.
Figure S3. The average fiber diameter and Young's moduli. (A) Fiber diameter calculated from SEM images as a function of annealing procedure; (B) Young's moduli measured by microindentation; (C) Young's moduli measured by a 3 point bending test of single fibers using an AFM. Polylactic acid (PLA) nanofibers were used as reference to verify the accuracy of the measurement. Error bars represent standard deviation of the mean. The statistical significance was determined using a one way ANOVA test, where *p<0.05. CO = conventional oven; MW = microwave oven; MW RTA = rapid thermal annealing in microwave oven.

Figure S4. Live & dead assay. hMSCs viability on ceramic scaffolds at day 3 in BM. Live and dead cells are represented by green and red fluorescence, respectively. The scale bar is 500 µm. CO = conventional oven; MW = microwave oven; BM = basic medium.
Flexible Yttrium-Stabilized Zirconia Nanofibers Offer Bioactive Cues for Osteogenic Differentiation of Human Mesenchymal Stromal Cells

Figure S5. DNA amounts (A-B) and metabolic activity (C-D) of hMSCs cultured on ceramic scaffolds in (A and C) BM and (B and D) OM at day 7 and 14. Error bars represent standard deviation of the mean. *p<0.05, ***p < 0.001. CO = conventional oven; MW = microwave oven; BM=basic medium; OM=osteogenic medium.

Figure S6. The expression of osteopontin (OPN), osteocalcin (OCN), and bone sialoprotein (BSP) in hMSCs cultured on scaffolds in BM and OM. The results were presented after normalizing to day 7 bulk YSZ discs in BM and OM, respectively. Error bars represent standard deviation of the mean. *p<0.05, **p < 0.01,
***p < 0.001. CO = conventional oven; MW = microwave oven; BM=basic medium; OM=osteogenic medium.

Figure S7. Calcium deposition test. (A) hMSCs cultured in a tissue culture plate as a control experiment, and stained for alizarin red S after 28 days culture in BM and MM; (B) The ceramic scaffolds without seeding hMSCs incubated in BM and MM for 28 days do not stain for alizarin red S. CO = conventional oven; MW = microwave oven; BM=basic medium; MM= mineralization medium.

Figure S8. Schematic representation of the 3 point bending fatigue test.
Tailoring Surface Nanoroughness of Electrospun Scaffolds for Skeletal Tissue Engineering

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Abstract:
Electrospun scaffolds provide a promising approach for tissue engineering as they mimic the physical properties of the extracellular matrix. Previous studies have demonstrated that electrospun scaffolds with porous features on the surface of single fibers, enhanced cellular attachment and proliferation. Yet, little is known about the effect of such topographical cues on cellular differentiation. Here, we aimed at investigating the influence of surface roughness of electrospun scaffolds on skeletal differentiation of human mesenchymal stromal cells (hMSCs). Scanning electron microscopy (SEM) and atomic force microscopy (AFM) analysis showed that the surface nanoroughness of fibers was successfully regulated via humidity control of the electrospinning environment. Gene expression analysis revealed that a higher surface roughness (Ra=71.0 nm) supported more induction of osteogenic genes such as osteopontin (OPN), bone morphogenetic protein 2 (BMP2), and runt-related transcription factor 2 (RUNX2), while a lower surface roughness (Ra=14.3 nm) demonstrated higher expression of other osteogenic genes including bone sialoprotein (BSP), collagen, type I, alpha 1(COL1A1) and osteocalcin (OCN). Interestingly, a lower surface roughness (Ra=14.3 nm) better supported chondrogenic gene expression of hMSCs at day 7 compared to higher surface roughness (Ra=71.0). Taken together, modulating surface roughness of 3D scaffolds appears to be a significant factor in scaffold design for the control of skeletal differentiation of hMSCs.

Key words:
Surface roughness, mesenchymal stromal cells, osteogenisis, chondrogenisis, 3D scaffold
4.1. Introduction

Human mesenchymal stromal cells (hMSCs) are considered as a promising candidate cell source for skeletal tissue engineering and regeneration due to their low immunogenic reaction in allogeneic hosts and multilineage differentiation potential including the chondrogenic, adipogenic and osteogenic lineages [1-4]. Moreover, hMSCs are an easy accessible cell source [5]. Intrinsic properties of the extracellular matrix (ECM), such as chemistry, and stiffness, have been demonstrated to regulate stem cell differentiation. It also has been proved that tailoring surface topography serves as one of the key roles regulating stem cell differentiation [6, 7]. Dalby and his co-workers cultured hMSCs on poly(methyl methacrylate) substrate with varying ordered nanopits [8] and found that semi-ordered nanopits (120 nm diameter, 100 nm deep, absolute or average 300 nm centre–centre spacing) induced the expression of bone specific proteins in hMSCs in contrast to completely ordered or random nanopits. In the work by Nuno et al., hMSCs were cultured on polycaprolactone substrates displaying a gradient in surface roughness [6]. Specific roughness (Ra ~ 2.1–3.1 μm; RSm ~ 71.1–48.1 μm) steered faster osteogenic commitment and stronger osteogenic expression compared to tissue culture polystyrene.

Recently, electrospun fibers have been intensively utilized as scaffolds in tissue engineering owing to their fibrous structure mimicking the physical dimensions of natural fibrous extracellular matrix [9-12]. Moreover, these fibers can be easily fabricated and modified [13]. The morphology of electrospun fibrous structure can be subdivided into the morphologies of single fiber and fibrous mesh [14]. The modification of the morphology of single fibers could be achieved via many approaches including varying collector temperature [15], solvent and relative humidity [9, 16], or by post-processing methods such as nanoimprint lithography [13], surface graft polymerization [17], physical coating or blending [18], wet chemical etching methods [19], and gas plasma [20]. Among these methods, varying the solvent and relative humidity has multiple superior advantages to other methods, since (i) it is a simple and direct process without any post-electrospinning treatment compared to nanoimprinting [13], wet chemical etching [19] and plasma [21]; (ii) it shows higher flexibility in controlling the degree of roughness than varying the collector temperature [15] and physical coating or blending [22]; (iii) it results in homogeneous treatment of each single fiber in contrast to nanoimprinting [13] and gas plasma [23] which can modify only localized surface areas of fibrous meshes. Therefore, this approach has been employed to produce electrospun scaffolds with nanoporouos features on their single fibers for a wide range of applications, such as drug delivery, filtration, and tissue
engineering [9, 15]. Previous studies have been demonstrated that the modification of the morphology of single fibers in electrospun scaffolds can directly affect cellular response [24-28]. We previously demonstrated that hMSCs cultured on nano-porous 10 µm fiber scaffolds had higher proliferation rates than hMSCs cultured on smooth 10 µm fiber scaffolds [24]. In addition, these hMSC cells were found to aggregate on smooth fiber scaffolds, whereas they showed a more spread morphology on the nano-porous fiber scaffolds. Shokrgozar et al. demonstrated that nanoroughness of PLGA electrospun scaffolds could enhance nerve cell growing rate up to 50% compared to smooth electrospun scaffolds [25]. Zhang et al. revealed that the introduction of elliptical nano-pores onto the surface of aligned electrospun PLLA microfibers allowed for enhanced cellular response of vascular smooth muscle cells, including increased cell adhesion, proliferation, alignment, and expression of vascular matrix proteins [26]. However, all the above mentioned studies mainly focused on the influence of surface roughness of electrospun fibers on cellular adhesion and proliferation. Little knowledge on the effect of electrospun fiber surface nanoroughness on stem cell differentiation is available.

Here, we aimed at investigating the influence of electrospun fiber surface nanoroughness on the skeletal differentiation of hMSCs. The surface roughness of electrospun scaffolds was tailored through varying the humidity during electrospinning. The topographical surface was determined by scanning electron microscopy (SEM) and atomic force microscopy (AFM). The biological relevance of each surface roughness on osteogenic and chondrogenic cell differentiation was assessed in terms of cell morphology, metabolic activity, alkaline phosphatase (ALP) activity or glycosaminoglycan (GAG) production, and specific gene expression profiles.

4.2. Materials and Methods

4.2.1. Fabrication of scaffolds

Poly (ethylene oxide terephthalate)/poly(butylene terephthalate) (PEOT/PBT, the weight ratio of PEOT/PBT=55/45, the molecular weight (g/mol) of the starting PEG segments used in the polymerization process is 300) was kindly provided by PolyVation B.V. (the Netherlands). A PEOT/PBT solution was prepared by dissolving the polymer in a mixture solvent of dichloromethane/1,1,1,3,3,3-Hexafluoro-2-propanol (v/v=97/3) with a final concentration of 20% (w/v). The electrospinning set-up was custom-made, consisted of an environmentally controlled electrospinning chamber connected to a syringe pump, and was able to move the spinneret in the chamber to obtain a homogeneous fiber distribution.
during the fabrication process. The polymer solution was fed at a rate of 8 mL/h using a syringe pump (KDS 100, KD Scientific). The applied voltage and tip-to-collector distance were 20 kV and 15 cm, respectively. The temperature in the electrospinning chamber was monitored during the process and kept constantly around 20 °C. The relative humidity in the spinning chamber was chosen to be around 20%, 50% and 70%, separately.

4.2.2. Fiber morphology and dimensions

The morphology of electrospun scaffolds was observed by SEM (Philips XL30 ESEM-FEG). Prior to SEM imaging, samples were gold sputtered with a Cressington Sputter Coater 108 Auto set at 30 mA for 40 seconds. To create cross section images, samples were first frozen in liquid nitrogen and then cut. The average fiber diameter and the distribution of fiber diameters was calculated by measuring at least 30 fibers in one SEM image and five images were used for each scaffold using Adobe Photoshop CS4.

4.2.3. Porosity and pore diameter of scaffolds

Apparent density and porosity of scaffolds were calculated using the following equation [12]:

\[ p = \left( 1 - \frac{\rho'}{\rho^\circ} \right) \times 100\% \]

Where \( p \) is the porosity of scaffold porosity, \( \rho' \) is the apparent density of scaffold and \( \rho^\circ \) the density of polymer materials used for fabricating scaffolds.

The pore diameter of scaffolds was calculated from SEM images using ImageJ according to previous study [29]. The diameters in each of at least 30 pores within an image for a total of four images per sample were quantified.

4.2.4. AFM analysis

Surface roughness analysis was performed through AFM using Tapping Mode (PicoScan Controller 2500, Molecular Imaging, USA) with a Super Sharp TESP cantilever: 42 N/m, 320 kHz, 2–5 nm ROC, No Coatings (Bruker AFM Probes) on a 1 µm² surface area (n=6). The average roughness (Ra), pore depth, and surface area differences were determined using the Scanning Probe Image Processor, SPIPTM, version 4.2.2.0 software. High quality images in 3D of the fiber surface were visualized at randomly different surface locations to verify the reproducibility of the observed characteristics.
4.2.5. Contact angle measurements

The wettability of scaffolds was determined by the sessile drop technique using an optical contact angle device (OCA15, Dataphysics, Germany). The sessile milli-Q water drop was deposited onto the sample surface with the syringe, and the drop contour was fitted by the Young-Laplace method. At least 3 different static contact angle measurements on each sample were performed.

4.2.6 Nile Red staining and analysis

The local environment of polymer surfaces was determined using a Nile Red adsorption technique. Fabricated scaffolds or spin-cast polymer films (controls) were exposed to a dilute solution (1μM) of Nile Red in milli-Q water for 1 minute. After this minute, the samples were briefly washed with fresh milli-Q water to remove excess Nile Red solution; samples were then sandwiched between a glass slide and a coverslip. Images were taken with a confocal laser excitation microscope DM6000 (Leica, Wetzlar, Germany) equipped with DFC365FX camera. For measurement of the local environment, Nile Red (λex = 514) emission was imaged over the range of 520 – 700 nm in 10 nm steps. Integrated intensity of the region of interest was then plotted against the average wavelength for each image, reconstituting the emission spectrum. All experiments were performed in duplicate, on two distinct samples (total of 4 measurements).

4.2.7. Protein absorption on scaffolds

Protein adsorption on electrospun scaffolds was investigated as previously described [21]. Briefly, samples were punched and placed in a 96 well plate. They were incubated in phosphate buffered saline (PBS) solution at 37 °C overnight. After removing PBS, 200 μL of bovine serum albumin (BSA) solution (2 mg/mL in PBS) was added to the wells and incubated at 37 °C overnight. The samples were rinsed with PBS to remove any non-adherent proteins. The amount of adsorbed protein was quantified using Bicinchoninic acid (BCA) protein assay kit (Thermo Scientific Pierce Protein Assay Kit). The absorbance was measured at 562 nm on a plate reader (Multiskan GO, Thermo Fisher, USA).

4.2.8. Cell culture and seeding

Human MSCs (donor No. 8001L) were isolated from bone marrow by Texas A&M Health Science Center [30]. Briefly, bone marrow was aspirated centrifuged to separate mononuclear cells that would be considered as passage zero (P0) cells after being
planted and harvested at 60–80% confluence. P0 cells were further expanded, harvested and frozen to obtain passage 1 (P1) cells. P1 cells were expanded in T-300 flasks at a density of $3 \times 10^5$ cells/cm$^2$ and cultured in basic medium (BM), comprising α-MEM (Gibco), 10% fetal bovine serum (Lonza), 2 mM L-glutamin (Gibco), 0.2 mM ascorbic acid (Sigma), 100 U/mL penicillin and 100 mg/ml streptomycin (Gibco). Cells were harvested at approximately 80% confluence for further scaffold-based study.

Electrospun sheets were punched into discs (1 cm in diameter) and placed in a 48-well plate. Rubber O-rings (Eriks B.V., The Netherlands) were put on the top of electrospun discs to prevent them from floating. For sterilization, the specimens were soaked in 70% (v/v) ethanol for 15 min three times. After that, they were rinsed with sterilized PBS 1 min for three times and immersed in BM overnight before cell seeding. After removing the medium from the scaffolds, hMSCs were seeded on each scaffold at a density of $1.5 \times 10^4$ cells/cm$^2$ in 80 µL BM. Cell-scaffold constructs were incubated for 4 hours to allow cell attachment, and topped up to 1 mL of BM, osteogenic medium (OM) (BM plus $10^{-6}$ M dexamethasone), or chondrogenic medium (CM). CM was composed of DMEM (Gibco), 1% ITS premix (Micronic BV), 50 µg/mL ascorbic acid (Sigma-Aldrich), 100 units/mL penicillin (Life technologies), 100 µg/mL streptomycin (Life technologies), 100 nM dexamethasone (Sigma-Aldrich), 40 µg/mL proline (Sigma-Aldrich), 100 µg/mL sodium pyruvate (Life technologies), 0.01 µg/mL TGF-β3 (R&D systems). The cell-scaffold constructs were cultured in an incubator with 5% CO$_2$ humid atmosphere at 37 °C up to 21 days. Culture medium was refreshed every two days.

4.2.9. Metabolic activity

PrestoBlue™ assay was used to assess cellular metabolism. Briefly, cell culture medium was replaced with a medium containing 10% (v/v) PrestoBlue™ reagent (Life technologies) and the plates were wrapped with aluminum foil to avoid light, followed by incubating at 37 °C for 1 h. Fluorescence was measured at 590 nm with a Perkin Elmer Victor3 1420 multilabel plate reader. Cell metabolic activity was analyzed on day 7 and day 21.

4.2.10. Quantification of DNA

The total amount of DNA per scaffold was quantified as described in a previous study [31]. Briefly, samples were washed twice with PBS and then processed to digest for 16h in a Tris/EDTA buffer composed of 1mg/mL proteinase K, 18.5 µg/mL iodoacetamine (Sigma-Aldrich) and 1 µg/mL Pepstatin A (Sigma-Aldrich). The lysate was divided into two parts.
One part was used for ALP or GAG assay and the others were used for DNA assay. For DNA quantification, the lysate was added to a buffer containing RNase (×1000), component B (part of the DNA quantification kit) and NaCl-EDTA buffer. Subsequently, quantification of total amount of DNA per scaffold was performed using the CyQuant® DNA assay (Life Technologies) according to manufacturer’s protocol. The signal was measured on a spectrophotometer (Victor3 1420, Perkin Elmer, USA) at an excitation wavelength of 480nm and emission wavelength of 520nm.

4.2.11. Alkaline phosphatase (ALP) activity assay

The ALP activity was assessed as in a previous study [21]. Briefly, Cell lysate mentioned in DNA assay paragraph were digested for 1h at room temperature with a buffer at a pH of 7.8 composed of 0.1M KH₂PO₄, 0.1M K₂HPO₄ and 0.1% Triton X-100. After that, a CDP-star® solution (Roche Life Science) was added and samples were incubated for 15 min at room temperature. The signal was analyzed using a Victor3 1420 plate reader (Perkin Elmer, USA). The value of ALP activity was normalized to DNA quantity per scaffold.

4.2.12. Glycosaminoglycan (GAG) assay

Cell lysates mentioned in DNA assay paragraph were pipetted in duplicate to 96-well plates. After addition of a dimethylmethylene blue dye (DMMB, Sigma-Aldrich) solution comprising 10 mM hydrochloric acid, 3.04g/L of glycine and 2.37g/L of NaCl, the plates were measured at an absorbance of 525 nm using a micro plate reader (Multiskan GO, Thermo Fisher, USA). The amount of GAG was calculated using a standard of chondroitin sulfate (Sigma-Aldrich).

4.2.13. RNA isolation and gene expression

Total RNA per scaffold was isolated following the protocol based on a previous study [32]. Briefly, samples were rinsed with PBS three times prior to adding 1mL TRIzol reagent (Invitrogen). Subsequently, 200 µL of chloroform was added to all samples, mixed vigorously, and centrifuged at 1200 g for 15 min at 4 °C. The aqueous phase containing RNA was transferred to filter columns provided by the NucleoSpinRNAII ISOLATON kit (Bioke) and the purification process was performed according to the manufacture’s protocol. The concentration and purity of total RNA were determined by using an ND 1000 Nanodrop (Life Technologies, USA). One µg of total RNA was used for first strand cDNA synthesis using iScript (Bio-Rad, Veenendaal, the Netherlands) following the
manufactures protocol. Quantitative polymerase chain reaction (qPCR) was carried out on a BioRad CFX96 real time PCR machine using SYBR-green supermix (Bio-Rad) and the primers as listed in Table 1. The Cycling conditions for running qPCR were as following: initial denaturation at 95 °C for 10 min, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. Relative expression of RNA was normalized to housekeeping gene B2M, and fold changes were calculated using the $2^{-\Delta\Delta CT}$ method, where $\Delta\Delta CT = (C_{T,\text{Target}} - C_{T,B2M})_{\text{time}} x -(C_{T,\text{Target}} - C_{T,B2M})_{\text{time}0}$.

4.2.4. Statistical analysis

Statistical analysis was determined using GraphPad Prism 5.01 (GraphPad software, San Diego, USA) for windows. Two-way analysis of variance (ANOVA), followed by a Tukey’s post-hoc test was performed to compare the gene expression, metabolic activity, GAG production and ALP activity. One-way ANOVA with Bonferroni’s multiple comparison tests was used to compare the surface roughness. A p value lower than 0.05 was considered statistically significant.

4.3. Results and Discussion

4.3.1. Characterization of electrospun scaffolds

The relative humidity showed a remarkable effect on the surface roughness of electrospun scaffolds. At a relative humidity of 20 %, the morphology of fibers was smooth (Figure 1A and 1D). Increasing the relative humidity to 50% resulted to apparent changes in surface morphology of fibers. Figure 1E shows that surface features or pores become evident. When the humidity increased from 50 % to 70 %, the pore depth increased as well as the pore size leaving little space between adjacent pores. The formation mechanism of pores on electrospun fibers could be elucidated using the combination of vapor-induced phase separation with thermal-induced phase separation [9, 16]. In a vapor-induced phase separation model, the non-solvent (water) from vapor phase penetrates into the polymer solution causing phase separation into a solvent-rich phase and solid-rich phase. On the other hand, in the thermal- induced phase separation model, the volatile solvent evaporates dramatically, the liquid jet cools down, leading to a phase separation into a polymer-rich phase and solvent-rich phase. In both models, the solvent-rich phase will cause formation of pores on the fiber surface, whereas the polymer-rich phase will eventually evolve into a solid matrix. The cross-section images (Figure 1G-I) indicated that all fibers had a solid interiors wrapped by an increasingly porous outer layer with increasing humidity.
Figure 1. SEM image of PEOT/PBT fibers (A-C) and their corresponding surfaces (D-F) and cross sections (G-I) fabricated by electrospinning under different relative humidity: 20% (A, D, G), 50% (B, E, H) and 70% (C, F, I). Scale bars are 10 µm (A-C) and 1 µm (D-I).

In addition, to investigate the effect of humidity on the topology of pores, AFM analysis was performed (Figure 2A). Three dimensional representations of surface topography for the electrospun fibers illustrate the differences in surface roughness between fibers by varying the humidity. These results are in agreement with the SEM data. The quantification results of surface roughness parameters including Ra, pore depth and surface area difference are presented in Figure 2B. A significant difference in surface roughness was observed between fibers fabricated at different humidity. Changing humidity from 20 % to 70 % corresponded to increasing Ra from $14.3 \pm 2.5$ nm to $71.0 \pm 11.0$ nm, surface area from $3.1 \pm 0.8$ nm to $63.3 \pm 6.2$ nm and pores depth from $27.2 \pm 4.7$ nm to $230.5 \pm 31.9$ nm. Altogether, increasing of relative humidity resulted in an increase of roughness, depth, and surface area of pores on the surface of electropun fiber. Similar findings were also reported by previous studies where a more porous morphology appeared on the surface of polystyrene fibers as electrospinning was performed in a higher-humidity environment [16, 33, 34].
Previous studies have demonstrated that fiber diameter, pore size and porosity of scaffolds influence cell behavior such as proliferation, attachment and differentiation [35-37]. To better understand the effect of nanosurface roughness on hMSCs differentiation, the surface roughness was varied while keeping other structural parameters, such as fiber diameter, pore size and porosity of the scaffolds constant. As shown in Figure 3A and B, no significant difference in porosity and pore size of the electrospun scaffolds was found when changing the relative humidity. In addition, the obtained electrospun scaffolds had average fiber diameters of 3.8 ± 0.8, 3.5 ± 1.0, and 4.0 ± 1.2 µm for spinning from relative humidity of 20%, 50% and 70%, respectively (Figure 3C-E). The relative humidity is known to have an impact on the fiber dimension which is essentially dependent on the polymer concentration as well as solvent used [34]. Given the fiber dimension scale range achieved in the present study, increasing relative humidity did not cause apparent difference in fiber dimension.
Chemical functional groups and/or blocks of the polymer chains may be reoriented during electrospinning process [38]. To determine the influence of relative humidity on the surface polarity of electrospun fiber, a Nile Red assay was employed (Figure 4). Nile Red is a hydrophobic and solvatochromic dye [39] that has previously been used to determine the hydrophilicity/phobicity of a wide range of polymers in the bulk [40] and the variability of polymer surface environments [41]. Shifts in the absorption or emission maximum of the dye adsorbed to a polymer indicate differences in hydrophilicity/phobicity (dielectric strength). The electrospun fibers were stained with Nile Red homogeneously (Figure S1). As shown in Figure 4, for all electrospun scaffolds (20%, 50%, and 70%), we found nearly identical emission intensity profiles (emission λ<sub>max</sub> = 600 nm); this is in contrast to control samples, where we observed shifts in the emission intensity based on polymer composition (see SI for details). The identical emission profile of Nile Red across all our samples suggests that the electrospun scaffolds have similar polarity on their surface. To confirm these measurements, control samples of spin-cast poly (methyl methacrylate) (PMMA) and polycaprolactone (PCL) were shown to have distinctly different Nile Red emission profiles. The PMMA had a lower emission profiles compared to PCL (Figure S2), which support the fact that PMMA was more hydrophobic than PCL [40].
Surface wettability of scaffolds is important for initial cell attachment and migration [42]. Wettability was assessed using contact angle measurements (Figure 5A and B). Generally, water sessile droplets penetrated into all scaffolds in the end, indicating hydrophilic properties of the mesh surface, probably due to the same chemical composition of scaffolds. In the present study, the material used for scaffold fabrication was PEOT/PBT which had a medium surface energy with an average contact angle of 48±1° [43]. Although water droplets would ultimately penetrate into all scaffolds, their spreading speed was different showing an inverse relationship with the surface roughness of scaffolds. Surface roughness, fiber diameter, and pore size are important factors affecting the contact angle of a surface, where the material itself is not varied [44]. As shown above, the scaffolds fabricated in different relative humidities displayed similar fiber dimension, pore size and surface chemistry, but varied surface roughness. Therefore, in the present study, the differences in surface energy were mainly attributed to the surface roughness. These findings are consistent with the results reported by Chuah et al., who demonstrated that increasing the surface roughness of polydimethylsiloxane (PDMS) substrates could enhance the hydrophobicity [45].

Figure 4. Nile Red emission profile of electrospun samples. Note that the samples have overlapping emission intensity profiles. $\lambda_{ex} = 514$. 
4.3.2. Protein adsorption

Protein adsorption on the different scaffolds was examined using BSA as a model protein, and the results are shown after normalizing to scaffolds fabricated under a relative humidity of 20%. As shown in Figure 6, no significantly difference in protein adsorption was observed between scaffolds from a relative humidity of 20% and 50%, which may be attributed to their close surface structure. On the other hand, scaffolds from a relative humidity of 70% showed significant higher protein adsorption than that of scaffolds from a relative humidity of 20%. This could be related to the increased surface roughness and surface porosity leading to a higher surface area. Such an increased protein absorption was also shown by Nandakumar et al.[21] who used gas plasma for surface modification of electrospun fibers. A higher protein adsorption on scaffolds is also known to influence the cellular response such as cell adhesion, proliferation and differentiation [46, 47].
4.3.3. Metabolic activity of cells

Metabolic activity of hMSCs seeded on scaffolds characterized by different surface roughness was investigated by prestoblue™ assay (Figure 7). The metabolic activity rate in BM at day 7 was similar to that of day 21, while a slightly lower metabolic activity was measured in OM and CM after 21 days. The decrease in metabolic activity rate in OM and CM could be ascribed to the differentiation of hMSCs. No statistical differences in metabolic activity were observed between the different surfaces roughnesses of the scaffolds in all media investigated in this study.

Figure 7. Metabolic activity of hMSCs cultured on scaffolds in BM (A), OM (B) and CM (C) at day 7 and day 21.
4.3.4. ALP activity and GAG production

ALP activity, a common marker for early osteogenesis, is known to be up-regulated in the early stages of osteogenic differentiation [48]. As shown in Figure 8A-B and Figure S3, at day 7, a similar ALP activity in all kind of scaffolds was found in both BM and OM. In contrast, fibers (the 70% sample) with a surface roughness of 71.0 ± 11.0 nm displayed a higher ALP activity compared to fibers (the 20% sample) with a roughness of 14.3 ± 2.8 nm in BM at day 21. The similar trend was also observed in OM. Therefore, it could be inferred that a higher surface roughness causes a higher ALP activity in the range studied in the present work. Our results are consistent with previous studies [49, 50], where osteoblasts were demonstrated to have a higher ALP activity on a rougher surface of titanium substrates. Furthermore, our previous study has also demonstrated that hMSCs express evidently higher ALP activity on ceramic nanofibers displaying a rougher surface (0.7 ± 0.2 nm) compared to a smoother one (0.4 ± 0.1 nm) under osteogenic culture condition [51].

GAG is a long-chain sugar molecule and plays an essential role in chondrogenic differentiation, since it is one of the main components of the proteoglycans forming articular cartilage extracellular matrix [52]. In a previous study by our group, additive manufactured three-dimensional (3D) scaffolds whose polymer fibers have tailored surface roughness (ranging from 0.07±0.01 µm to 1.95±0.55 µm) were fabricated in combination with a wet-spinning technique [53]. The results demonstrated that hMSCs on a highest roughness scaffolds (1.95±0.55 µm) showed a higher GAG production under chondrogenic culture condition. Despite the GAG production in all types of scaffolds enhanced over the experimental time course in all media (Figure 8C-D and Figure S4), no statistical difference in GAG production was found between the different surface roughnesses, indicating that the surface roughness nanometric range investigated in this study does not significantly influence GAG production by hMSCs during differentiation into the chondrogenic lineage.
Figure 8. ALP activity of hMSCs cultured on scaffolds in BM (A) and OM (B) at days 7 and 21. GAG production on scaffolds in BM (C) and CM (D) at day 7 and day 21. The final results of ALP activity and GAG production level was normalized by DNA amount. *p<0.05, ***p < 0.001.

4.3.5 Osteogenic gene expression

The influence of surface roughness on osteogenic differentiation was confirmed by real time RT-PCR (Figure 9). Osteopontin (OPN) is known as a major non-collagenous extracellular structural protein with respect to cell adhesion, migration, and biomineralization [54]. Bone morphogenetic protein-2 (BMP2) is known to induce osteogenesis of hMSCs [55]. At day 7 in BM, OPN and BMP2 expressions were significantly higher for rougher fibers (the 70% sample) compared to smoother fibers (the 20% sample). One the other hand, bone sialoprotein (BSP), collagen type I (COL1A1) and osteocalcin (OCN), which are extracellular matrix proteins of bone [56], showed higher expression on smoother fibers (the 20% sample) compared to rougher fibers (the 70% sample). To know the osteoconductive properties of different surface roughness, the same panel of osteogenic markers was also investigated in OM. In the investigated surface roughness range, RUNX2 expression was significantly higher on rougher fibers (the 70% sample) compared to smoother fibers (the 20 %-sample) at day 7. Conversely, BSP expression on smoother fibers (the 20 % sample) was significantly higher than that
on rougher fibers (the 70% sample) after day 21. Recent studies revealed that the hierarchical combination of micro- and nanoscale surface features promotes osteogenic differentiation [57, 58]. Khang. et al. reported that a stronger osteogenic differentiation of hMSCs was supported by a heterogeneous composition of nano and sub-micron hybrid surface features compared to only nano scale or sub nano scale features [57]. In another study, Gittens et al. demonstrated that a combination of micro-/submicro-scale surface roughness with nanoscale structures on Titanium surfaces enhanced osteoblast differentiation [58]. Therefore, it could be assumed that the hierarchical combination of different surface nano-roughness in electrospun fibers may be more beneficial on the osteogenic differentiation of hMSCs.

4.3.6 Chondrogenic gene expression

A panel of chondrocyte-related genes, including collagen type II alpha 1 (COL2A1), activated leukocyte cell adhesion molecule (ALCAM), SOX9, and aggrecan (ACAN) was investigated to assess hMSCs chondrogenesis (Figure 10). In BM, only SOX9 was differentially regulated. On the other hand, the gene expression profile was much different under chondrogenic differentiation culture conditions, most likely due to the dominant effect of chemical stimuli. In CM, upregulation of all tested chondrogenic markers was observed on the 20% sample compared to the 70% sample due to their different surface roughness. However, this trend was disappeared after 21 days. Recent work suggested that the tendency of hMSCs differentiation which is trigged by surface topography stimuli may be discontinued when the surface topography cues disappear [53, 59]. This might have also happened in this study when hMSCs were fully confluent on scaffolds and started being embedded within the endogenous ECM. SOX9 is known as an early chondrogenic transcription factor which control the expression of genes related to matrix secretion [60]. Upregulation of SOX9 could be activated in mouse stromal ST2 cells cultured in low oxygen tension (hypoxic environment) [61]. The expression of SOX9 was significantly higher on rougher fibers (the 70% sample) than on smoother fibers (the 20% sample) in both BM and CM after 21 days. This might be related to the higher spatial proximity of the hMSCs cultured in rougher scaffolds, which facilitate cells condensation resulting in hypoxic condition when cells are confluent on the scaffolds [53, 62].
Figure 9. Quantitative RT-PCR gene expression analysis of osteoblast-related genes for hMSCs cultured on scaffolds in both BM and OM at days 7 and 21. *p<0.05, **p < 0.01, ***p < 0.001.
Figure 10. Quantitative RT-PCR gene expression analysis of chondrocyte-related genes for hMSCs cultured on scaffolds in BM and CM at time points of day 7 and day 21. *p<0.05, **p < 0.01, ***p < 0.001.

4.4. Conclusion

In the present work, three types of electrospun fibers with surface nanotopography were directly fabricated by varying the relative humidity during electrospinning. These scaffolds had similar fiber dimensions, porosity and pore size, and chemical composition of the surface, yet a different surface roughness on the fibers. SEM and AFM results demonstrated that higher relative humidity resulted in higher surface roughness, pore depth and surface area in the nanoscale range. In order to investigate the impact of individual fiber surface nanotopography on cell differentiation, hMSCs were cultured on scaffolds for 21 days in osteogenic and chondrogenic differentiation media. We found that a higher surface roughness (Ra = 71.0 nm) showed enhanced expression of osteogenic genes such as OPN, BMP2, and RUNX2, while a lower surface roughness (Ra = 14.3 nm) demonstrated enhanced other osteogenic genes expression, including BSP, COL1A1 and OCN. Interestingly, a lower surface roughness (Ra = 14.3 nm) showed to better support the chondrogenic differentiation of hMSCs at day 7 compared to higher surface roughness (Ra = 71.0 nm). Tailoring surface nanoroughness of electrospun scaffolds could, therefore, be used as a promising approach to control the cellular response of hMSCs in tissue engineering and regenerative medicine.
Reference


Surpporting Information

**Figure S1.** Nile Red emission profile of PCL and PMMA casting film.

**Figure S2.** Representative of image of 20% humidity electrospun scaffold stained with Nile Red. $\lambda_{ex} = 514$. 
Figure S3. ALP activity (A and C) and DNA concentration (B and D) of MSCs cultured in scaffolds in basic medium (A-B) and osteogenic medium (C-D).

Figure S4. GAG activity (A and C) and DNA concentration (B and D) of MSCs cultured in scaffolds in basic medium (A-B) and chondrogenic medium (C-D).
Chapter 5

From Fiber Curls to Mesh Waves: a Platform for the Fabrication of Hierarchically Structured Electrospun Fibers Mimicking Natural Tissue Formation

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Abstract
Despite the tremendous progress in electrospinning in the past two decades, little progress has been made toward fabricating fibers having features mimicking the structure and architectural organization of extracellular matrix (ECM) fibrils and/or fibers in native tissue. Here, we present a simple, efficient, and versatile technique to produce curly or wavy patterns in electrospun fibers by using thermally shrinking materials. The pattern features could be easily manipulated by controlling various processing variables including the orientation of deposited fibers, the temporary content of sacrificial fibers, and the shape, geometry, chemistry and shrinking percentage of the thermally shrinking materials. In addition, our technique allows the retention of the imparted buckled pattern without the use of additional fixing agents or methods. Additionally, the technique allows fabrication of scaffolds with gradient and multiscale wavy patterns, thus mimicking the organization of multiple native tissues. Mechanical properties of the fabricated curly fibers could be varied through the shrinkage ratio of thermally shrinking materials, which enables the possibility to obtain a structure that could not only mimic the topographical patterns of connective tissues, but also mechanically resembles the properties of the same. We further investigated if the wavy patterns could regulate transforming growth factor-beta (TGF-β) production, a key signaling pathway involved in connective tissue development. Our results demonstrated that a TGF-β reporter cell line cultured on wavy scaffolds had higher TGF-β expression compared to non-wavy scaffolds. This platform may open up new opportunities in fabricating smart fibers for regenerative medicine.

Key words:
Electrospinning, shrink films, buckling, gradients, tissue regeneration
5.1. Introduction

Extracellular matrix (ECM) fibrils or fibers with a curly or wavy microstructure are observed in several human tissues, such as ligaments, tendons, heart muscle, valve leaflets, cornea, iris, and intervertebral disk [1-4]. Typically, the crimped or wavy structure of native fibrous collagen has a periodicity in the range of 10 to 200 µm [1]. Fibers with a crimped structure are capable of absorbing more deformation compared to their straight counterparts, and therefore show enhanced compliance at low strains while holding greater strength as the load increases [1, 5]. Thus, collagen fibers containing a crimped morphology serve to buffer mechanical loads and enable the tissue to show nonlinear stiffness under increasing tensile strains [5].

Scaffold-based regenerative medicine has emerged as a promising strategy to repair damaged or diseased tissues. Currently, many techniques for generating scaffolds have been exploited such as phase separation, molecular self-assembly, electrospinning and additive manufacturing. Among these techniques, electrospinning has been widely explored due to its ability to produce fibrous meshes comparable to native extracellular matrix (ECM) in a simple and versatile manner [6, 7]. In addition, a desired pattern and structure of deposited electrospun fibers can be readily achieved by varying the collection methods, for example, randomly oriented fiber collected by a stationary plate, aligned fibers fabricated by a rotating mandrel or electrodes and tubular constructs produced using a rotating mandrel [6, 8-10]. Recently, electrospun scaffolds have been fabricated to mimic the native ECM environment in various degrees, particularly resembling the curly features. Younan and colleagues demonstrated the use of a plasticizer-based treatment, especially using ethanol, to generate a crimp structure along each electrospun fiber [5]. The morphology of crimp pattern obtained from their work is similar to other previous studies. Amsden et al. found that aligned electrospun fibers would self-crimp after removal from the mandrel at a temperature higher than the glass transition temperature of the spun polymer [11, 12]. They also demonstrated that the degree of crimping could be varied by increasing the operating temperature or applying a plasticizer treatment on the polymer fibers. Using the same fabrication method, Brian and colleagues assessed the response of fibroblasts seeded on crimped-like structured scaffolds under dynamic mechanical loading [12]. Fibroblasts showed enhanced ECM synthesis and collagen accumulation in crimped structured scaffolds compared to uncrimped fibers [12]. Tonin and colleagues produced crimped polymer fibers utilizing a weak tangential air-flow fed
from the top of a cylinder placed between the spinneret and the target collector[13]. Although the above mentioned methods have shown to mimic the wavy patterns observed in native ECM in different degree, the major drawbacks of these methods include limited control of the crimp pattern, relative complicated processing steps, the use of plasticizers potentially being toxic for the cells, and difficulties in generating multiscale and gradient wavy features, which are common features in native tissues.

Here, we propose a simple, versatile and efficient method that uses electrospinning combined with thermally shrinking materials, which tend to return to their original unstretched shape when heated above their softening point ($T_s$). The thermally shrinking materials, such as corresponding films or tube-like semi-finished products, are typically either mono- or biaxially oriented according to their manufacturing protocol[14]. When the extending force is applied in one direction, monoaxial orientation occurs whereas when the stretching force is simultaneously applied in two directions, (equi)baxial orientation is obtained. This causes it to shrink back toward its initial dimensions. By applying proper cooling, the molecules will be frozen in this state until sufficient heat energy is applied to allow the chains to shrink back. Using this new method, we are capable of fabricating architectures resembling the ones of collagen fibrils in various types of connective tissues including tendons and ligaments, muscles, skin and connective tissues of the eye.

5.2. Materials and Methods

5.2.1. Depositing fibers on monoaxially oriented PLA shrink film

Block copolymers PEOT/PBT (weight ratio of PEOT/PBT is 55/45; starting poly(ethylene glycol) (PEG) segments have a molecular weight of 300Da) were purchased from PolyVation® (Groningen, The Netherlands). The polymer solution for electrospinning was prepared by dissolving PEOT/PBT in chloroform/1,1,1,3,3,3-hexaofluoro-2-propanol (HFIP) (v/v=4:1) with a final concentration of 20%(w/v), and stirred overnight at room temperature until complete dissolution. The solution was then loaded into a 5 mL syringe attached to a needle with a diameter of 0.8 mm and controlled by a syringe pump at a flow rate of 0.5 mL/h. A high voltage of 16 kV was applied and the distance between the tip of the needle and the collector was 15 cm. The temperature and relative humidity of the electrospinning chamber, which were kept the same for all electrospinning experiments, were controlled at 25 °C and 30%, respectively. Fibers were deposited on a oriented PLA film (4 cm x 4 cm) which was placed on a conducting collector plate (for a random deposition) or two electrodes (for an aligned deposition). For collecting aligned
fibers, changing the position of the PLA film during the electrospinning allows the deposition of fibers according to specific orientations: parallel, diagonal and perpendicular orientations relative to the direction of stretching and shrinkage were used in this experiment. The fiber density was controlled by controlling the deposition time. To know the influence of deposition orientation on curled patterns of fibers deposited in a low density, fibers were collected for 5 s on each PLA film. To investigate the effect of fiber density on curled patterns, fibers were collected up to 45 min. Fiber-film constructs were then kept in an oven pre-heated to 75 ± 1 °C for 1 min to allow shrinkage of the PLA film or film frame, and formation of a buckled fiber mesh.

5.2.2. Tailoring the buckled pattern by selective sacrificial fibers

PVA was dissolved in ethanol/water (v/v=1:4) at a final concentration of 8% (w/v) while the recipe for PEOT/PBT solution was the same as described above. To investigate low fiber densities, PEOT/PBT and PVA aligned fibers were deposited on a PLA film sequentially in a layer-by-layer way. The tip-to-collector distance and voltage applied, which were kept the same for the PEOT/PBT and the PVA solutions, were set to 15 cm and 16 kV, respectively. The feeding rates of the PEOT/PBT and the PVA solutions were 2 mL and 1 mL/h, respectively. To investigate the effect of sacrificial fibers on fiber meshes, PEOT/PBT and PVA fibers were electrospun simultaneously from two corresponding syringes and collected by a mandrel with a rotating speed of 300 rpm to which a PLA film was attached. The two syringes were set up at either side of the mandrel with a tip-to-collector distance of 13 cm. The feeding rates for PEOT/PBT and PVA were 5 mL/h and 1 mL/h, respectively. The same voltage of 16 kV was applied for both solutions.

Fiber-film constructs were then transferred to an oven pre-heated to 75 °C ± 1 °C for 1 min to allow shrinking of the PLA film and formation of a buckled fiber or fiber mesh. Finally, the fiber-film constructs were immersed in water for 30 min to remove PVA fibers.

5.2.3. Influence of shrinking percentage of thermo-shrinkable material on curled features

PEOT/PBT fibers were deposited on oriented PLA films which were placed on the top of a static collector. The fibers were collected for 30 min. To investigate the influence of shrinking ratios of the films on the curly patterns, the fiber-film constructs were shrunk at 65 °C for different time periods.
5.2.4. Influence of shape and geometry of thermo-shrinkable materials on curled features

To fabricate a fiber mesh with a multiscale crimped pattern, a frame punched out from a PLA film was put on the top of a pair of electrodes to collect the fibers. The spinning parameters for PEOT/PBT were the same as described above. The fibers were collected for 10 min.

To obtain a tubular curled scaffold, PEOT/PBT fibers were collected for 30 min using a mandrel wrapped with an oriented PLA film stripe and rotating with a speed of 150 rpm. The feeding rate of the polymer solution was 5 mL/h. The distance between the needle tip and the collector was set to 12 cm and the applied voltage was 16 kV. Fiber-film constructs were then transferred to an oven pre-heated to 75 °C ± 1 °C for 1 min to allow shrinking of the PLA film and formation of a buckled fiber or fiber mesh.

5.2.5. Buckled PVA, PAN and PS fibers using PLA film

PVA was dissolved in ethanol/water (1:4) at a concentration of 8 % (w/v) and stirred overnight before electrospinning. The applied voltage was 16 kV, the distance between needle tip and collector was 15 cm and the feeding rate of the polymer solution was 1 mL/h. Aligned fibers were collected using a pair of electrodes over which a PLA film was placed.

For electrospinning of PAN fibers, 10% polymer solution was prepared by dissolving PAN (MW=150,000, sigma) in dimethylformamide (DMF) and was stirred overnight at 60 °C to dissolve the polymer completely. The feeding rate of the polymer, the applied voltage, and the tip-to-collector distance was optimized at 1 mL/h, 16 kV and 15 cm, respectively. Aligned fibers were collected using a pair of electrodes with PLA film on top of it. The fibers were collected for between 10 s and 5 min.

PS (MW=350,000, from Sigma) was dissolved in DMF at room temperature under stirring overnight to obtain a 20 wt% homogeneous solution. The polymer solution was delivered at a rate of 0.5 mL/h. Electrospinning was carried out at 18 kV and 15 cm distance from tip to collector. Fibers were deposited on the PLA film which was placed on top of a pair of electrodes. Fiber-film constructs were then transferred to an oven pre-heated to 75 ± 1 °C for 1 min to allow shrinkage of the PLA film and formation of a buckled fiber mesh.
5.2.6. **Buckled PEOT/PBT and PAN fibers using shrinkable PS material**

The electrospinning process for PEOT/PBT fibers was the same as described above except for the collection methods. For collecting random fibers, a biaxially oriented thermo-shrinkable PS film with a shrinking ratio of around 50 % was put on top of a static plate collector. For collecting aligned fibers, the PS film was wrapped around a mandrel with a high speed of 5000 rpm. The thickness of fiber meshes was controlled by the deposition time.

The fabrication of PAN fibers was also the same as described above. PAN fibers were deposited on biaxially oriented thermo-shrinkable PS films placed on top of a pair of electrodes. The fibers were collected for 10 s. Fiber-film constructs were then transferred to an oven pre-heated to 120 °C for 15 min to allow shrinkage of the PS film and formation of buckled fiber materials.

5.2.7. **Characterization of buckled fibers**

To check the morphology of the scaffolds, the samples were sputter-coated with gold and then observed under a scanning electron microscopy (SEM) (Philips XL-30) at an accelerating voltage of 10 kV. For the low density mats, the wavelength and amplitude were measured using Image J; 40 fibers were measured for each condition (5 images with 8 fibers each). The wavelength of the curly features was considered as the distance between two consecutive peaks; the amplitude was measured as the distance between the maximum and minimum of the wave divided by two.

Uniaxial tensile tests were performed to evaluate the mechanical properties of the different curled structures. Six different percentages of shrinkage (10, 25, 30, 40 and 50%), obtained via varying the exposition time to a shrinkage temperature, were analysed while non-curly fibers were used as control. A total of 7 samples for each type were tested. The samples were given a rectangular shape using a surgical scalpel so that the length-to-width aspect ratio (4:1) provided uniform tensile stress in the region where the strain was measured. All the samples had a constant cross-sectional area (CSA) after shrinking, which was measured with a digital calliper (accuracy of 0.02 mm and resolution of 0.01 mm) and we assumed it had a rectangular shape. The samples were fixed in a standard clamps and axially aligned to the 500 N load cell of a Zwick Z020 material-testing machine. The scaffolds were preconditioned by a series of 10 cycles where they were exposed to 3% of strain at a strain rate of 0.1 %/s, in order to reduce hysteresis. The samples were then pulled until failure at a strain rate of 0.3 %/s. With CSA and strain
measurements, a stress-strain curve representing the tensile mechanical properties of the
different scaffolds was created. From the stress-strain curves, the following parameters
were obtained: Young’s Modulus (Pa), defined as the slope of the linear region of the
stress-strain curve, ultimate stress (Pa), ultimate strain (%) and strain energy density (Pa)
at failure. The failure modes were also noted.

5.2.8. hMSC culture and seeding

Pre-selected hMSCs (donor No. 8001L) were obtained from the Institute of Regenerative
Medicine at Texas A&M University. Briefly, a bone marrow aspirate was drawn and
mononuclear cells were separated using density centrifugation. The cells were plated to
obtain adherent hMSCs, which were harvested when cells reached 60-80% confluence.
These were considered passage zero (P0) cells. These P0 cells were expanded,
harvested and frozen at passage 1 (P1) for distribution. hMSCs were thawed and
expanded in T-flasks till reaching 80% confluence in basic medium composed of
Minimum Essential Medium (α-MEM, Gibco), 10% FBS (Lonza), 2 mM L-glutamine
(Gibco), 0.2 mM ascorbic acid (Sigma), 100 U/mL penicillin and 100 mg/mL streptomycin
(Gibco). Samples were punched to fit a 48-well plate. After sterilization with 70% ethanol
for 15 min for 2 times, the scaffolds were washed with phosphate-buffered saline (PBS) 3
times, afterwards incubated in basic medium for 3 hours at 37 °C before cell seeding.
Cells were seeded in scaffolds with a cell density of 5000 cells/cm². The cell culture was
carried on for 5 days with medium replacement every 2 days.

5.2.9. Cell penetration analysis

The samples were fixed with formalin for 30 minutes and washed with PBS. To get cross-
sections, samples were cut in half and embedded with Cryomatrix™ mounting media on
mounting blocks with the cross-section facing upwards and 8 µm cross-sections were
obtained using a cryotome (Shandon Cryotome, Thermo Scientific). To avoid polymer
auto-fluorescence, the cross-sections were stained with 1% (w/v) Sudan Black solution in
70% ethanol, incubated for 1 hour at room temperature, washed thoroughly with PBS for
5 min for 5 times. To visualize cell penetration into the scaffold, cross-sections were
stained with DAPI. Briefly, samples were incubated with 1% Triton x-100 for 10 min on ice,
washed with PBS and blocked with bovine serum albumin (10mg/mL) for 60 minutes to
block unspecific binding. After removal of this solution, DAPI (1:100 PBS) was added for
15 min at room temperature avoiding light. Finally, all samples were washed with PBS.
and kept at 4 °C in the dark. Images were then taken using a fluorescent microscope (Nikon Eclipse E600).

To measure the penetration depth associated with each cell in each cross-section, the pictures were analyzed using Image J. The approach taken was to define the borders of the section and then to create a grid for each image which divided the height of the cross-section in 5 bins (Figure S9), corresponding to regions from the top to the bottom of the scaffold. The first bin was considered the top of the scaffold while the last bin was identified as the bottom of the scaffold. To normalize the data, the number of cells in each bin was divided by the total amount of cells in that cross-section and the values presented in the x-axis were presented as the ratio of infiltration depth to the thickness of scaffold.

5.2.10 MLEC culture and seeding

MLEC, a kind gift from Dr. Daniel Rifkin from the Department of Cell Biology at the New York University School of Medicine, were stably transfected with an expression construct containing a truncated plasminogen activator inhibitor-1 (PAI-1) promoter fused to the firefly luciferase reporter gene. These cells are highly sensitive and specific for TGF-β production based on its ability to induce PAI-1 expression. MLEC were thawed and expanded in T-flasks till reaching 80% confluence in Dulbecco’s Modified Eagle’s Medium α (DMEM-α) supplemented with 10% FBS, 0.2 mM L-glutamine, and 100 units/mL penicillin and 100 mg/mL streptomycin. The samples were punched to fit a 96-well plate (Thermo Scientific). After sterilization with 70% ethanol for 15 min for 3 times, the scaffolds were incubated in the described medium (without TGF-β1) overnight. The next day, cells were seeded with a cell density of 50,000 cells/cm² on the scaffolds and on tissue culture plates used as a control. Cell culture was carried on for 5 days.

5.2.11. Luciferase and DNA assay

After removing the medium, samples were washed thoroughly with PBS, lysis buffer (5x) was added and diluted (5:1) in Milli-Q water. Samples were stored at -80 °C for at least 1h, then defrosted on a plate shaker at room temperature. Each sample was divided, one half for the Luciferase and the other half for the DNA assay and transferred into 2 different, white and black 96-well plates, respectively. For the Luciferase assay, Luciferase Assay Substrate (Promega) was added to the samples and the light produced was read on a plate reader (Victor 3™, PerkinElmer®) column by column.
The DNA assay was performed using the CyQUANT® cell proliferation assay kit (Molecular Probes, Eugene, USA) according to its manufacture manual. Briefly, samples were supplemented with lysis buffer containing RNAse (Sigma) and incubated for 1h on a plate shaker, meanwhile a standard curve was prepared using λDNA standard (provided from the CyQUANT kit). After the incubation period, CyQUANT® GR dye in lysis buffer was added to the samples and the fluorescence was measured on a plate reader (Victor 3 ™, PerkinElmer®) at an excitation wavelength of 480 nm and an emission wavelength of 520 nm.

5.2.12. Statistical Analysis

Statistical differences were determined using GraphPad Prism 5.001 software (San Diego, USA). Statistical differences between groups were determined using analysis of variance (ANOVA) followed by a Tukey’s post-hoc test unless otherwise mentioned. Statistical significance between the control group and the experimental groups are indicated by (*) which represents a p-value < 0.05, by (**) which represents a p-value < 0.01, and by (***) which represents a p-value < 0.001. Values are presented as mean ± standard deviation.

5.3. Results and discussion

5.3.1. Fabrication of curly fibers
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Figure 1. Schematic of a typical process for generating curled patterns in electrospun fibers using thermal shrinkage. (a) Depositing electrospun fibers over a thermo-shrinkable film. (b) Fiber-film construct before thermal shrinkage. (c) Wavy fibers form after thermal shrinkage. (d) and (e) Wavy fibers are detached from the thermally shrinkable film using a non-solvent. (f) Random fibers are collected through using a static ground plate. (g) Aligned fibers are obtained using a pair of electrodes.

Figure 1 shows the schematic of producing curly fiber. In a typical process, polymer fibers are deposited onto thermo-shrinkable materials (e.g. a corresponding planar film) via electrospinning (Figure 1a and b). By heating the construct above $T_s$, the fibers buckle as a consequence of the substrates shrinking while the fibers adhering to them. Cooling down the construct at room temperature leads to the thermal fixation of the curled pattern generated in the electrospun fibers, and this is one of the main advantages of our technique compared to previous studies, where an extra fixation agent or method was always required[1, 5]. In the present work, a block copolymer poly(ethylene oxide terephthalate)/poly(butylene terephthalate) (PEOT/PBT), which has been demonstrated as a biocompatible and -degradable polymer, was used as a material for electrospinning[15, 16]. In addition, a monoaxial poly(lactic acid) (PLA) film having a $T_s$ around 65 °C was utilized as the main thermo-shrinkable material. To determine its shrinking properties, three different temperatures (65, 75 and 85 °C) were tested during different periods of time (Figure S1). According to these results, standard values for the
film shrinkage were set at 75 °C for 1 minute which allowed 60% shrinkage of the PLA film and enough time to control the process manually. Importantly, the applied $T_s$ of thermo-shrinkable materials should be below the melting temperature ($T_m$) of the polymer used for fabricating the fibers. Figure S2 shows poly(caprolactone) (PCL) fibers which became partially melted after shrinking the PLA film onto which they have been deposited. No curls were formed because the $T_m$ of PCL (around 60°C) is lower than the $T_s$ of the PLA film.

5.3.2. From single fiber curls to multiscale mesh waves

Random fibers were collected using a static ground plate, while aligned fibers were generated using a pair of parallel electrodes. Changing the angle of the PLA film during electrospinning allows deposition of fibers relative to specific film orientations (Figure. 2a). In case of a monoaxially oriented film, a perpendicular configuration corresponds to deposit fibers at a 90° angle with respect to the shrinkage axis; a parallel configuration corresponds to deposit fibers along the shrinkage axis; a diagonal configuration corresponds to fiber deposition at a 45° angle; a random configuration corresponds to randomly oriented fibers. As shown in Figure 2b, after shrinking the PLA film, curled features were generated in fibers of all deposition direction except for the perpendicular one. The perpendicular configuration only allowed fiber compactness, because the space between each fiber decreased as the film shrunk, bringing the fibers closer together. The wavelengths and amplitudes of curls for parallel configuration showed a significant difference compared to that of diagonal configurations (Figure 2c). However, it is difficult to calculate the wavelengths and amplitudes for the random configuration, since the fibers were randomly deposited and crossed each other, leading to heterogeneous curls in fiber.
Figure 2. From single fiber curls to multiscale mesh waves. (a) A schematic illustration of varying fiber orientations by changing the position of the film over the electrodes. Red arrows indicate the shrinking direction of the oriented PLA film, while the green circular arrow means rotating the film to achieve different fiber orientations. (b) Comparison between fibers deposited in different directions relative to the direction of shrinkage before and after shrinkage. (c) Histogram of the wavelength and amplitude for parallel and diagonal configurations. (d) Topographical evolution as the time of deposition increased and with it the mesh thickness. Samples where fibers were deposited perpendicular to the direction of shrinkage showed waviness rather than only curls on individual fibers after 9 minutes of deposition, whereas the perpendicular samples show waviness after 5 minutes. (e) Depicting the thickness of samples with fibers deposited perpendicular or parallel to the shrinkage axis, after shrinking, according to time of deposition. Scale bars: 20 μm (b) and 500 μm (d). *p-value<0.1, **p-value<0.01 and ***p-value<0.001.
With the same process, it is possible to transfer the effect of generating curls on single fibers to the effect of generating multiscale waves on fiber meshes. The transition between the two effects relied on the density of deposited fibers. The density of deposited fibers in turn depends on the time of spinning; longer times correspond to a thicker fiber deposition, leading to the formation of fiber meshes evolved from low density fibers. For the given fiber chemistry and dimension in the present study, the multiscale waves on perpendicular samples started at 9 min deposition time with an average thickness of $0.3 \pm 0.1$ mm, whilst for the parallel samples it occurred at 5 min with the same average thickness of $0.3 \pm 0.1$ mm (Figure 2d and e). Although previous methods have proved capable of generating curled patterns on single fibers, they have no way of controlling the transition between generating curls on single fiber and generating multiscale waves on fiber meshes. With the present technique, the fabricated scaffolds can assume different structural configurations, which are promising as they may mimic the hierarchical organization of native tissues from the nano- to the macro-scale. The presented platform works also for other polymer fibers. As shown in Figure S3 and Figure S4, curly fibers of polyvinyl alcohol (PVA), polystyrene (PS) and polyacrylonitrile (PAN) could be generated using the oriented PLA films. Besides oriented PLA, the present technique could also be translated to other thermo-shrinkable materials. Theoretically, every thermoplastic material can be made into a stretched, oriented film, the frozen stress of which can be unlocked resulting in shrinking when arriving at the stretching temperature again, for instance, oriented PS films. Figure S5 and Figure S6 show that curly features were successfully generated in PEOT/PBT and PAN fibers, respectively, using an equibiaxially shrinkable PS film.

5.3.3. Maneuvering curled pattern using additional sacrificial electrospun fibers

The curled pattern of fibers could also be tailored by using sacrificial fibers, such as those made from water soluble polymers like PVA. Figure 3a shows the schematic of fabricating a mixture of PVA and PEOT/PBT (PVA-PEOT/PBT) fibers, where PVA fibers were firstly deposited on the PLA film, followed by PEOT/PBT fibers. After removal of the sacrificial PVA fibers, PEOT/PBT fibers held a distinguishably different curled pattern having both periodical larger arches and small curls within the same fiber, which did not appear in the absence of the sacrificial polymer. However, the removal of the sacrificial polymer was distinguishable whether it was the first or second layer. As shown in Figure S7a-c, the final PEOT/PBT-PVA fibers in which the PVA was deposited as a second layer have a
different curly morphology after removing PVA fibers. The comparison of curled features between final PVA-PEOT/PBT and PEOT/PBT-PVA fibers is summarized in Figure S7d. The wavelength and amplitude for the larger arches in PVA-PEOT/PBT fibers were $47.2 \pm 22.9$ and $6.2 \pm 1.8 \, \mu m$, respectively, which is similar to the values of crimped patterns in human ligament collagen (wavelength of 45-60 $\mu m$ and amplitude of 5-10 $\mu m$) [11]. On the other hand, the corresponding values for smaller curls in PVA-PEOT/PBT fibers were $5.2 \pm 1.6$ and $1.9 \pm 0.5 \, \mu m$, respectively. In the case of PEOT/PBT-PVA fibers, they had an average wavelength of $3.2 \pm 1.3 \, \mu m$ and average amplitude of $1.4 \pm 0.5 \, \mu m$, which is significantly different ($p< 0.01$) from the PVA-PEOT/PBT fibers. Previous methods used to curl electrospun fibers could produce similar patterns, to the ones achieved in the present work. [11, 17, 18]. For example, Tao Han et al. produced bucked electrospun fibers through manipulating the buckling instability of the electrospinning jet. The wavelength range corresponding to buckled pattern in their study was 1.2-30.6 $\mu m$ [17], which range was comparable to the ones produced in the present work (from $3.2 \pm 1.3 \, \mu m$ up to $47.2 \pm 22.9 \, \mu m$). Based on a complex method of calcination and reduction of copper, Guoqing Chang et al. produced 3D helical ribbons with fibers presenting a wavelength of $3.2 \, \mu m$ [18].

This strategy of using a sacrificial polymer could also be translated to fiber meshes for tuning their surface topography. It has been demonstrated that topographical cues are important to cellular behaviors, including cell morphology, proliferation, migration and differentiation [19, 20]. Figure 3b shows the comparison between meshes before and after immersion in water. A difference in surface topographical pattern between them was visible, inclusively a smaller fraction structure after immersion due to the removal of the PVA fibers. A similar phenomenon was also referred in previous work of Baker et al. where they combined electrospinning with sacrificial fibers to generate a scaffold having engineered delamination [21]. Taken together, through selective removal of fibers, it is possible to modulate the wavy or curled patterns on single fibers as well as fiber meshes.
Figure 3. Tayloring curled features by using additional sacrificial electrospun fibers or tuning thermo-shrinkable materials. (a) Tayloring curled patterns along fibers by sacrifying a second layer of fibers from poly(vinyl alchol) (PVA). Illustration of fiber layer deposition (top left); SEM image of curled PEOT/PBT-PVA fibers before (top right) and after (bottom right) sacrifying the PVA fibers; morphology of curled PEOT/PBT fibers without using a sacrificial layer of PVA fibers (bottom left). (b) Changing wavy patterns of fiber mesh by using scarificial PVA fibers. Schematic figure of fabricating a PEOT/PBT-PVA blending fiber mesh (top left); PEOT/PBT-PVA fiber mesh before (top right) and after (bottom left) shrinking; morphology of the wavy PEOT/PBT-PVA fiber mesh after sacrificing the PVA fibers (bottom right). (c) Schematic of fabricating fibers having multiscale wavy patterns. (d) Morphology of curled pattern in a square outlined area (left) and in encircled outlined area (right) in c. (e) SEM image of electrospun fiber having a gradient waviness. (f) Fabricating a buckled fiber mesh mimicking the structure of the trachea by using PLA shrink tubes. Morphology of PEOT/PBT fibers deposited on PLA tubes before (left) and after (right) shrinking. Scale bar: 20 µm (a), 100 µm (d), and 500 µm (b, e and f).

5.3.4. Influence of shrink properties and geometric shape of thermally shrinkable materials on curled features

As demonstrated in Figure S1, the shrinkage ratio of thermo-shrinkable materials is time dependent. Based on this concept, the curled features could be regulated via manipulating shrinkage ratios (Figure S8). When the shrinkage ratio increased from 10% to 40%, the amount of “wrinkles” in scaffolds increased, whereas the average wavelength
of the “wrinkles” decreased correspondingly. The structure and geometry of thermo-shrinkable materials could be designed for specific requirement. Figure 3c shows an electrospun mesh having multiscale buckled patterns generated using a planar oriented PLA film with an open central area. When the fiber mesh shrunk, the fibers possessed a lower ratio of shrinkage on the area where there was no film underneath compared to the area where fibers adhered to the film. Thus, multiscale buckled patterns could be generated on the fiber mesh. After shrinking, the fiber mesh had a coiled pattern on the sides, in contrast to a smoother wavy pattern in its central zone (Figure 3d). Native tissue interfaces, such as the corresponding regions of tendons, ligaments and bone show gradients in structure, composition and mechanics [22, 23]. In the specific case of the interface between tendon or ligaments with bone, a gradient in mechanical properties serves to mitigate stress concentrations [24]. Interestingly, with one corner of the oriented PLA film clamped and thus hindering from shrinking, a gradient wavy pattern could be generated (Figure 3e). A scaffold having such gradient waviness could be used in the field of interface tissue regeneration. Figure 3f shows electrospun fibers deposited on a tubular oriented PLA film. After shrinking, a corrugated hose-like construct mimicking the structure of a trachea segment of 104 ± 22.5 µm could be fabricated.

Overall, a desirable curly feature could be achieved via manipulating the shrink properties of thermally shrinking materials and their geometric shapes in particular. Currently, it is able to design the geometric shapes of thermally shrinking materials into a variety of stretch configurations, such as tubes and more complex geometries (e.g., through thermoforming of films or blow molding of tubes), than only mono- or (equi) biaxially oriented films.

5.3.5. Mechanical properties

The mechanical behavior of the 3D waves was evaluated performing uniaxial tensile tests. Scaffolds showed a limited hysteresis on the preconditioning phase. During the uniaxial tensile testing, stress-strain curves representing the mechanical properties of the structure were obtained. The curves show a stress-strain behavior characterized by a non-linear toe region between 0 and around 1% of strain, a linear region between around 1% and 12%, and a final region where the curve reached the ultimate stress. The obtained results are listed in Table S1 and Figure 4a. A statistically significant decrease of the Young’s modulus with the increase of the shrinking percentage was detected (Figure 4b). Similar trends were found for the ultimate stress, ultimate strain and strain.
energy density (Figure 4c-e). In particular, the 25% of shrinkage seemed to represent a threshold in the case of the ultimate stress and strain energy density.

In order to describe the mechanical behavior of the different curled networks, the experimental data obtained from the mechanical tensile tests were analyzed through a mathematical model. Hyperelastic models are commonly used to describe the behavior of skin, other soft-tissue and rubber-like materials because of their elastic behavior[25]. Here, the 3-parameter Mooney-Rivlin model was used to describe the material’s stress-strain relationship as a function of the empirically determined constants $C_{01}$, $C_{10}$ and $C_{20}$. Considering a uniaxial tensile test, the model can be expressed as

$$\sigma = 2 C_{10} \left( \lambda - \frac{1}{\lambda^2} \right) + 2 C_{01} \left( \lambda - \frac{1}{\lambda^2} \right) + 4 C_{20} \left( \lambda - \frac{1}{\lambda^2} \right) \left( \frac{\lambda^2}{\lambda^2} + \frac{2}{\lambda} - 3 \right)$$

where $\sigma$ is the applied stress, $C_{10}$, $C_{01}$ and $C_{20}$ are material parameter constants, and $\lambda$ is the stretch ratio, defined as $\lambda = 1 + \varepsilon$. The estimation of the model parameters is provided in the Supplementary Table S2 and Figure S11 in the Supplementary Note.

The use of the Mooney-Rivlin model in case of a uniaxial tension condition allowed to fit the experimental data obtaining three different adimensional constants. The statistical differences among the different parameters demonstrated a different mechanical behavior between the parallel networks and the ones with different percentage of shrinkage.
Figure 4. Mechanical properties of wave structures. (a) Figures of stress–strain behavior (left) and zoom in the encircled outlined area in left (right). (b) Young’s modulus. (c) Ultimate stress. (d) Ultimate strain. (e) Strain energy density.

5.3.6. Cell penetration

Cell penetration and growth throughout the scaffold are vitally important for tissue engineering applications[26]. Although electrospinning scaffolds have drawn much interest in the tissue engineering field, one major limitation is that their fibers are typically densely packed, therefore hampering cellular infiltration into the scaffold [26, 27]. Thus,
many approaches have been explored to overcome this problem, for instance sacrificial polymers electrospinning, salt leaching, and electrospinning in combination with electrospraying [26]. However, the current methods to improve cell infiltration reduce scaffold stability [21, 28]. Here, we hypothesized that wavy fibers with a larger void space into their porous structure will enhance cell penetration into electrospun scaffolds. Human mesenchymal stromal cells (hMSCs) were seeded at the same density on flat and wavy scaffolds in basic medium. SEM images revealed that hMSCs on flat scaffolds are likely to disperse on its surface while a big portion of cells on wavy fibers tend to entangle within its fibers (Figure 5a and b). To further investigate their infiltration and growth into flat and wavy scaffolds, nuclear staining was performed and cross-sections were analyzed. hMSCs on the flat scaffolds did not migrate further than superficial layers, whereas hMSCs on the wavy scaffolds migrated deeper and faster into the scaffold (Figure 5b). The distribution of cells in each depth section of flat and wavy scaffolds is summarized in Figure 5c and d. For comparison, the depth presented in the x-axis corresponds to an average ratio of infiltration depth to the thickness of scaffolds, since the wavy scaffolds had a higher thickness compared to the flat scaffolds, as a consequence of the buckling effect occurring during the underneath film shrinking (Figure S9b). By day 2, the largest amount of cells stayed at the top of the scaffolds and decreased in number towards the bottom of the scaffolds for both wavy and flat substrates. Despite presenting the same trend, flat scaffolds presented a higher concentration of cells in the upper region whereas wavy scaffolds showed a more homogeneous distribution. After 5 days of culture, more cells migrated into the deeper regions of both scaffolds. However, more cells were present in the deeper region of the wavy scaffolds compared to the flat ones. These results thus indicate that wavy scaffolds support a more homogenous distribution of cells throughout their thickness and a faster cell migration than flat scaffolds.
Figure 5. Cell penetration and Luciferase assay. (a) The morphology of hMSCs grown on flat (top) and wavy (bottom) scaffolds. (b) The scaffold cross-section showing cell nuclei labeled with DAPI. (c) and (d) The distribution of cells present in each depth section of the flat and wavy scaffold. For each scaffold region, the number of cells was counted and normalized to the total cell number present in the sample. The cellular infiltration depth is normalized to the thickness of scaffolds. (e) Quantification of DNA amount. (f) Comparison of Luciferase expression between flat and wavy scaffolds. (g) Luciferase expression with DNA normalization. Scale bars: 10 μm (a) and 200 μm (b).
5.3.7 TGF-β signaling

Given the resemblance of the obtained fibers with wavy patterns of connective tissues, we investigated if the wavy patterns could modulate the TGF-β pathway, a key signaling pathway involved in connective tissue development. A mink lung epithelial cell line harboring a TGF-β reporter construct (T-MLEC) was used to that end [29]. An increase in luciferase activity corresponds to an increase in TGF-β signaling. In the present experiments, fetal bovine serum (FBS) that does contain TGF-β was used as a positive control [30]. As shown in Figure 5f, cells seeded on a wavy scaffold emitted significant more light than cells seeded on a flat scaffold at day 5, suggesting a corresponding increase in TGF-β signaling.

5.3.8 Similarities between native tissues and the scaffolds fabricated

Figure S10 shows a few examples of curled pattern characteristics of connective tissues that could be recreated with the scaffolds proposed here. The organization of musculoskeletal tissues (Figure S10a), connective eye tissue like the iris (Figure S10c), and the artery wall (Figure S10e) composed of curled collagen fibers interconnected by a matrix can be mimicked by several of the curly fibers and wavy meshes that we can fabricate with this technique (Figure S10b, d and f). The trachea also displays a crimped pattern (Figure S10g) that could be mimicked by a wavy high-density scaffold that was electrospun for 45 min with the fibers lying down perpendicular to the shrinkage axis (Figure S10h).

5.4 Conclusions

In summary, we have demonstrated a simple but effective technique for generating crimped patterns on electrospun fibers by exploiting thermo-shrinkable materials. The wavy pattern features could be tuned by adjusting processing parameters, including the orientation of the deposited fibers, the percentage of shrinkage of the thermo-shrinking films and the use of sacrificial fibers. By controlling the spinning time, we were able to transfer the curls from a single fiber to multiscale waves in a fiber mesh. This approach opens up many opportunities in generating wavy patterns, for example gradient curls, and wavy patterns in a tubular geometry. In addition, the proposed technique can be readily extended to many other polymer fibers and thermos-shrinkable materials to generate functional scaffolds that could be beneficial for mimicking a variety of tissues organizations, such as iris, trachea, muscles, artery wall, and ciliary body. Furthermore,
this new method may also inspire innovation in other application fields in future, such as functional textiles.
Chapter 5

References

From Fiber Curls to Mesh Waves: a Platform for the Fabrication of Hierarchically Structured Electrospun Fibers Mimicking Natural Tissue Formation

Supporting Information

**Figure S1.** Shrinking properties of a monoaxially oriented PLA film at different temperatures over different times. (a) 65 °C, (b) 75 °C and (c) 85 °C.

**Figure S2.** Failed buckling PCL fibers with oriented PLA films. (a) PCL fibers before shrinking. (b) PCL fibers after shrinking. PCL fibers become molten and no crimped pattern could be observed. This was because the melting point of PCL is lower than the shrinking temperature of PLA. Scale bar: 20 µm (a and b).
Figure S3. Fabricating curled PVA fibers using oriented PLA films. (a) Single fiber with alignment diagonal to the shrinking direction of PLA film before (top) and after (bottom) shrinking. (b) Fibers with alignment parallel to shrinking direction of PLA films before (top) and after (bottom) shrinking. (c) Fiber mesh with alignment parallel to the shrinking direction of the PLA film before (top) and after (bottom) shrinking. Scale bar: 20 µm (a-c).

Figure S4. Fabrication of PAN and PS fibers using oriented PLA films. (a) PAN fibers deposited on PLA film before (top) and after (bottom) shrinking. (b) PS fibers deposited on PLA film before (top) and after (bottom) shrinking. Scale bar: 10 µm (a and b).
Figure S5. Fabricating curled PEOT/PBT fibers using oriented PS films. (a) Random PEOT/PBT fibers before (top) and after (bottom) shrinking. (b) Aligned PEOT/PBT fibers before (top) and after (bottom) shrinking. Scale bar: 20 µm (a-b).

Figure S6. Fabricating curled PAN fibers using oriented PS films. (a) PAN fibers before shrinking. (b) PAN fibers after shrinking. Scale bar: 10 µm (a and b).
Figure S7. (a) Illustration of fabricating PEOT/PBT-PVA fibers. PEOT/PBT fibers were firstly deposited on the surface of the PLA film, then a second layer of PVA fibers was deposited on top of it. (b) SEM image of curled PEOT/PBT-PVA fibers before sacrifying the PVA fibers. (c) The morphology of curled PEOT/PBT-PVA fibers after sacrifying the PVA fibers. (d) Figure depicting the ways to measure wavelengths and amplitudes for big loops and small loops. (e) Comparison of curly features between PEOT/PBT-PVA and PVA-PEOT/PBT fibers. PEOT/PBT-PVA corresponds to PEOT/PBT as first layer of deposition and PVA as second layer, and PVA-PEOT/PBT corresponds to PVA as first layer and PA as second layer. The distinction between big loops and small loops is explained in (d), where the same fiber has two different curls that were measured separately. ***p-value<0.001. Scale bar: 20 µm (b and c).
**Figure S8.** The influence of shrinkage ratio on wavy feature. (a) The morphology of wavy pattern at 10% shrinkage ratio. (b) The morphology of wavy pattern at 25% shrinkage ratio. (c) The morphology of wavy pattern at 40% shrinkage ratio. (d) Quantification of curly features generated from different shrinkage ratios. ***p-value <0.001. Scale bar: 1 mm (a-c).

**Figure S9.** (a) Grid created in ImageJ to measure cell infiltration along the cross-section. The numbers represent the bins for each scaffold cross-section, 1 corresponding to the top and 5 to the bottom; (b) The graph shows the doubling effect in thickness registered after shrinkage. **p-value <0.05.
Figure S10. Similarities between native tissues and the fibers obtained. (a) A SEM image of musculoskeletal tissue composed wavy fibers. (b) Fibers fabricated in the present study exhibit a similar wavy configuration to a. (c) A SEM image of the iris (scale bar: 1 mm; from www.sciencephoto.com). (d) The features of the scaffold in the present study show a similarity to that of the iris in c. (e) An image of the artery wall composed of curled elastin bundles separated by collagen (from www.sciencephoto.com). (f) A shrunk low density scaffold in the present study resembles the structure in e. (g) A SEM image of a tracheal wall displaying a crimp pattern (from www.sciencephoto.com), the scale bar is unknown. (h) High density shrunk scaffold in the present study resemble g. (i) A SEM image of a ciliary body. (j) The structure obtained in the present study having similarity to i.
**Table S1. Mechanical analysis.**

<table>
<thead>
<tr>
<th>Shrinkage</th>
<th>Young’s modulus (MPa)</th>
<th>Linearity range (%)</th>
<th>Ultimate stress (MPa)</th>
<th>Ultimate strain (%)</th>
<th>Strain energy density (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parallel</td>
<td>9.91 ± 2.24</td>
<td>1-7</td>
<td>2.75 ± 0.76</td>
<td>131.78 ± 29.42</td>
<td>3.04 ± 0.87</td>
</tr>
<tr>
<td>10%</td>
<td>5.12 ± 1.15</td>
<td>1.8-10</td>
<td>2.24 ± 0.33</td>
<td>245.96 ± 21.37</td>
<td>4.66 ± 0.77</td>
</tr>
<tr>
<td>25%</td>
<td>6.49 ± 1.64</td>
<td>1.8-10</td>
<td>2.69 ± 0.47</td>
<td>199.18 ± 37.93</td>
<td>5.28 ± 1.67</td>
</tr>
<tr>
<td>30%</td>
<td>4.61 ± 1.16</td>
<td>1-7</td>
<td>1.24 ± 0.20</td>
<td>112.64 ± 37.74</td>
<td>1.27 ± 0.50</td>
</tr>
<tr>
<td>40%</td>
<td>2.87 ± 0.46</td>
<td>1-9</td>
<td>1.37 ± 0.32</td>
<td>150 ± 32.56</td>
<td>1.76 ± 0.53</td>
</tr>
<tr>
<td>50%</td>
<td>2.40 ± 0.31</td>
<td>1-12</td>
<td>1.13 ± 0.15</td>
<td>124.9 ± 26.64</td>
<td>1.19 ± 0.25</td>
</tr>
</tbody>
</table>

**Supplementary Note: Mechanical modeling**

The results of the model fitting are summarized in Table S2.

**Table S2. Mooney-Rivlin model: parameters estimation.**

<table>
<thead>
<tr>
<th>Shrinkage</th>
<th>C01</th>
<th>C10</th>
<th>C20</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parallel</td>
<td>3.39 ± 1.33</td>
<td>-1.43 ± 0.89</td>
<td>0.130 ± 0.120</td>
<td>0.962 ± 0.011</td>
</tr>
<tr>
<td>10%</td>
<td>1.29 ± 0.24</td>
<td>-0.17 ± 0.08</td>
<td>0.007 ± 0.003</td>
<td>0.974 ± 0.007</td>
</tr>
<tr>
<td>25%</td>
<td>2.06 ± 0.54</td>
<td>-0.47 ± 0.25</td>
<td>0.022 ± 0.011</td>
<td>0.972 ± 0.003</td>
</tr>
<tr>
<td>30%</td>
<td>1.93 ± 0.79</td>
<td>-0.91 ± 0.57</td>
<td>0.081 ± 0.062</td>
<td>0.986 ± 0.003</td>
</tr>
<tr>
<td>40%</td>
<td>1.28 ± 0.41</td>
<td>-0.44 ± 0.27</td>
<td>0.032 ± 0.024</td>
<td>0.961 ± 0.011</td>
</tr>
<tr>
<td>50%</td>
<td>1.13 ± 0.50</td>
<td>-0.37 ± 0.34</td>
<td>0.032 ± 0.041</td>
<td>0.982 ± 0.018</td>
</tr>
</tbody>
</table>
A statistically significant decrease of the parameter $C_{01}$ with the increase of the shrink percentage was detected. A similar trend was found for the parameter $C_{10}$. No statistical differences were found among the different percentages of shrinkage for the parameter $C_{20}$.

Figure S11. Mooney-Rivlin model: comparisons of different conditions for each parameter.
Chapter 6

Direct Writing Electrospinning of Scaffolds with Multi-dimensional Fiber Architecture for Hierarchical Tissue Engineering

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Abstract
Nanofibrous structures have long been used as scaffolds for tissue engineering (TE) applications, due to their favorable characteristics such as high porosity, flexibility, high cell attachment and enhanced proliferation, and overall resemblance to native extracellular matrix (ECM). Such scaffolds can be easily produced at a low cost via electrospinning (ESP), but generally cannot be fabricated with a regular and/or complex geometry, characterized by macropores and uniform thickness. We present here a novel technique for direct-writing (DW) with ESP to produce complex three-dimensional (3D) multi-scale and ultrathin (~1 µm) fibrous scaffolds with regular pore sizes and controllable fiber bundle morphology. Using this technique, a scaffold mimicking the zonal organization of articular cartilage was fabricated as a proof of concept, demonstrating the ability to better mimic native tissue organization. The DW scaffolds directed tissue organization and fibril matrix orientation in a zone-dependent way. Comparative expression of chondrogenic markers revealed a substantial upregulation of Sox9 and aggrecan (ACAN) on these structures compared to conventional electrospun meshes. Our novel method provides a simple way to produce customized 3D ultrathin fibrous scaffolds, with great potential for TE applications, in particular those for which anisotropy is of importance.

Keywords:
Direct-writing, electrospinning, tissue engineering, articular cartilage, human mesenchymal stromal cells
6.1. Introduction
Articular cartilage is present in joints to provide a smooth, lubricated surface for articulation and resistance to high dynamic compressive loads[1]. Histologically, it is characterized by sparse chondrocytes distributed within a dense extracellular matrix (ECM), mainly composed of water, proteoglycans, glycoproteins, and collagen in a zonal configuration. These layers are distinguished as superficial, middle, deep, and calcified zone, which vary in molecular composition, cell density, and collagen orientation[2]. Articular cartilage is devoid of mesenchymal progenitors and blood vessels, and as a result, the potential for self-repair is limited. Therefore, when damage occurs, there is a chance for progressive degeneration that may lead to osteoarthritis[3].

Tissue engineering (TE) of cartilage constructs is a promising approach to replace the currently suboptimal clinical treatments, aiming to develop a biomimetic replacement with appropriate mechanical behaviour. Additive manufacturing (AM) techniques have greatly contributed to this goal. These technologies have provided valuable tools for the custom fabrication of 3D scaffolds with reproducible patterns, tunable porosity, and tailored physicochemical properties[4]. However, limitations inherent to these processes still persist. Notably, the required setups are typically expensive and the range of usable materials is dependent on the fabrication technique, for instance extrudable thermoplastic polymers for fused deposition modelling. These techniques also introduce other concerns such as thermal degradation of the polymer during the fabrication process[4]. Moreover, current AM techniques possess low spatial resolution and thus the features of the produced scaffolds are typically several orders of magnitude larger than ECM fibers, particularly collagen fibrils. This results in a poor cell-seeding efficiency and proliferation, as well as non-uniform scaffold coverage.

Nanofibrous scaffolds are interesting alternatives for cartilage TE, with a highly porous structure reminiscent of the ECM milieu that has been shown to promote cell attachment, and stimulate cell function while facilitating the efficient exchange of nutrients, oxygen, and metabolites[5]. Nanofibers are also very flexible due to their high aspect ratio, allowing cells to remodel their surroundings with produced matrix[6]. To fabricate nanofibrous mats, solution electrospinning (ESP) is one of the most popular methods, since it is easy to operate, relatively inexpensive, and versatile in terms of materials selection[7]. ESP is a complex hydrodynamic process governed by many variables and characterized by an unpredictable electrified jet[8]. The chaotic nature of this jet results from the build-up of surface charges generated by an applied electrostatic force, which
initially leads to droplet elongation, Taylor cone formation, and jet emanation, but soon overcomes the liquid surface tension, causing bending instabilities[8, 9]. As charge density increases, repulsion forces alter the stable jet into a whipping jet. At this point, fiber deposition occurs randomly and thus accurate pattern designing is impeded. Therefore, most ESP products are dense meshes that lack a regular pore-size network and have an architecture defined by the collector shape (e.g., flat or grooved plate, rotating drums, spaced electrodes, etc.) [9, 10].

To achieve control over an ESP process would enable the combined benefits of the topography provided by electrospun mats with the reproducibility and designing potential of AM techniques. Therefore, much interest has been taken in the pursuit of direct writing (DW) with ESP, with some efforts already reported. One strategy is based on the application of an electrostatic lens system to focus the electrical field into a narrow region. Through the use of a ring electrode positioned between the spinneret and collector, Neubert et al. and Bellan et al. showed the targeted deposition of fibers into a single spot [11, 12]. By either moving the collector or using steering electrodes, simple lines could be patterned onto the collector, but no complex patterning was achieved. A similar but improved strategy was followed by Jongwan Lee et al., who employed a side-wall electrode, a thin glass plate collector and a sharp-pin electrode to focus the electrified jet and draw regular 3D nanofibrous patterns[13], which could later be stacked to build a larger scaffold[14]. Although successful, the aforementioned approach relied on a complex setup that is difficult to adopt in other labs. An earlier and simpler approach that dispenses additional setup modifications is near-field electrospinning (NFESP), in which the unstable region of the jet is bypassed, by closely approximating the spinneret to the collector and using very low voltages[15-17]. However, drawbacks to this technique are evident, as it relies on a solution-dipped probe for jet feeding, which greatly limits the possible scaffold size.

In another approach, melt ESP replaces the polymer solution with a molten polymer to generate fibers[18, 19]. Application of melt ESP for DW of poly (ε-caprolactone) (PCL) fibrous structures for hydrogel reinforcement has revealed great potential in engineered cartilage constructs[20]. This technique has been also used to fabricate cell-invasive 3D scaffolds constructed from photo-cross-linkable poly(L-lactide-co-ε-caprolactone-co-acryloyl carbonate) microfibers[21, 22]. Because of the use of high melting temperatures, the technique may cause polymer degradation and is restricted to the use of
thermoplastic polymers. In addition, it excludes the addition of bioactive molecules, such as collagen, which undergo denaturation at high temperatures[19, 23]. The minimum feature size is also a concern in melt ESP, since the Barus effect induces a jet enlargement[24] and obstructs formation of nanofibers, although recent efforts have shown the production of sub-micron fibers in special conditions[25]. This process also produces a single fiber at a time, while tissue development benefits from the presence of multiple interconnected fibers that act as anchoring points for cells to assemble focal adhesions[26].

Here, we show the development of a novel DW technique that modifies conventional solution ESP to produce accurately patterned ultrathin 3D scaffolds. We determined the optimal conditions for DW ESP, including solution, ambient conditions, and processing parameters, to focus the jet to a single point and collect the fibers in a dry state. Patterns could be designed via predefining XY translation of the collector, and the resulting fibers stacked to create 3D constructs. After characterizing this technological process, we fabricated scaffolds that mimic the zonal configuration of articular cartilage. These scaffolds were seeded with human mesenchymal stromal cells (hMSCs) and cultured in chondrogenic differentiation medium over the course of 21 days. The DW ESP scaffolds successfully supported hMSC proliferation and chondrogenic differentiation, as evaluated by matrix production and orientation, tissue development, and gene expression. These data argue that DW ESP can fabricate ultrathin scaffolds with multidimensional fiber architecture and favorable biological characteristics, which has great potential for wide TE applications.

6.2. Results and Discussion

Traditional ESP generally deposits fibers with random morphology due to the bending instability of the electrospinning jet (Figure 1A). However, controlled fiber deposition, and thus patterning, is possible by DW with ESP, and requires the jet to follow a stable path from the moment it emanates from the spinneret tip, until it reaches the collector (Figure 1B). To fabricate suitable scaffolds to mimic articular cartilage, we deployed DW with ESP. First, we optimized the ESP process for DW, including polymer solution, flow rate, applied voltage, working distance and ambient conditions.

6.2.1. Solution conditions
For these studies, we used a polymer solution of Poly(ethylene oxide terephthalate)/poly(butylene terephthalate) (PEOT/PBT) for the scaffolds, because PEOT/PBT has been demonstrated as a biocompatible and degradable polymer with promising application for skeletal regeneration[27, 28]. Elongation and stability of the ESP jet depends on optimal solution parameters. Solutions with a very high viscosity usually exhibit longer stress relaxation times, which can prevent fracturing of the jet during ESP[9]. We tested solution mixtures with polymer concentrations between 26–30% (w/v) and CHCl₃:HFIP solvent ratios between 70:30 and 80:20 (v/v). We found that the optimal range of polymer concentration that permits DW was 27–29% (w/v), which yielded fairly viscous solutions. The optimal range of solvent ratio that permits DW was between 74:26 and 78:22 (v/v), which ensured good polymer dissolution and yielded a solution with high evaporation rate. A solution with high evaporation rate is particularly important for the DW technique, since it enables fibers to dry quickly within shorter working distances (Wd) used in these studies. Within the optimal range of polymer concentration and solvent ratio, the solution with 28% (w/v) polymer in 75:25 CHCl₃/HFIP (v/v) revealed the best outcome for a stable and continuous DW process (Table S1).

6.2.2. Solution flow rate

The solution flow rate through the spinneret is an important factor, as it influences the amount of transferred material and jet velocity[9]. We tested flow rates from 0.05 mL/h to 2 mL/h. We found that flow rates between 0.08 and 0.3 mL/h, a range that enables a low but steady flow of the polymer solution through the spinneret, were necessary for successful DW ESP. The influence of flow rates (0.05, 0.1, and 0.5 mL/h) on the jet and fiber deposition is shown in Figure S1A–C to exemplify the conditions tested. At a low flow rate of 0.05 mL/h, the amount of solution that was supplied through the spinneret was insufficient to maintain a regular jet for the applied voltage (8 kV) and scan speed, thus leading to the deposition of an erratic line (Figure S1A). As the flow rate increased to 0.1 mL/h, a thin and steady jet was produced, resulting in the formation of straight line patterns (Figure S1B). When the polymer solution supply was 0.5 mL/h, the jet could no longer be focused and consequently, the result was the collection of dispersed fiber mesh (Figure S1C). Thus, we concluded that the optimal flow rate was ~0.1 mL/h.

6.2.3. Applied voltage and working distances
In addition to the solution conditions and flow rate, the interplay between the applied voltage and the distance from the collector affects the solution jet. The applied voltage needs to reach a certain threshold to initiate the ESP process[8, 29]. As the voltage increases, the solution ejection also increases and higher electrostatic repulsive forces build up on the fluid jet[8, 9]. The optimal applied voltage is also dependent on the Wd, as both parameters are related in the determination of the electric field strength[30]. A typical electrified jet will follow a short stable path before acquiring a whipping profile[31], thus selecting an appropriate Wd is of utmost importance. We tested applied voltages between 2–16 kV, with corresponding Wd values between 2–12 cm. The threshold voltages needed to initiate the ESP process increased as the Wd increased: the minimum voltages were 4 kV at 2 cm, 4.5 kV at 3 cm and 5.5 kV at 4 cm (Table S2). It is worth mentioning that following jet initiation, the spinning voltage could be reduced to a value lower than its jet initiation voltage. For example, at 2 cm, the lowest Wd tested, the minimum voltage for jet initiation was 4 kV during DW ESP. However, following jet initiation, the spinning voltage could be reduced to a minimum value of 2.5 kV. At 7 kV and a Wd of 3 cm, we observed an unfocused jet, which broke down into multiple thin jets as a consequence of jet bending instability (Figure 1C). In this case, the amount of ejected solution provided insufficient viscoelastic force to balance surface charge density[32]. When we deceased the voltage to 5 kV, a stable jet was achieved during its flight time (Figure 1D), depositing and vertically building the fibers in a single point in the static collector. However, this process could not continue indeterminably due to an increase in charge accumulation, resulting in an increase of repulsive forces at that point (Figure S2). We then increased the Wd to 8 cm and tested different applied voltages. At 8 kV, the jet trajectory was stable and straight lines could be fabricated (Figure S1D). At 12 kV, the amount of ejected fluid increased, resulting in too much volume at the collector to produce single steady lines at the given scan speed (Figure S1E). At 16 kV, the repulsive charges accumulated and led to the onset of bending instabilities, resulting in the complete loss of the focusing effect (Figure S1F). When the Wd was further increased to 10 cm and 12 cm, either the jet was collected in its whipping phase or the electric field strength was too low to focus it, resulting in unpredictable fiber deposition (Figure S1H–I). From these data, we concluded that Wd ≤ 8 cm and voltages ≤ 8 kV were necessary to collect the jet in a stable and focusing manner.

From voltage/Wd parameters that resulted in successful focused spinning, we examined the morphology of the deposited fibers (Table S2) and found three different
regimes (Figure 2D–F). The transition between these regimes depended both on Wd and electrical field strength (V), which could be simplified as the ratio V/Wd. Typically, for low V/Wd, the fibers were randomly deposited in a bundle and dispersed from each other (Figure 2D). As the V/Wd ratio increased, fiber density increased and the distance between fibers decreased, resulting in increased fiber cohesiveness (Figure 2E). If the V/Wd ratio was increased by voltage, the fiber bundle displayed a more three-dimensional core, with the fibers aligned in the direction of the jet movement, as well as a thicker fiber dimension (Figure 2A and Figure S3). If the V/Wd ratio was increased via reducing the Wd, similar morphological changes was achieved together with an increase of fiber diameter (Figure 2B and Figure S4). The transition between a random deposition (lower limit) and an anisotropic alignment (upper limit), was defined by the V/Wd ratio, thus suggesting that these morphological changes are determined by the amount of solution ejected. To obtain a single fiber configuration (Figure 2F), a V/Wd > 2 was required. For example, at a Wd of 2 cm, an applied voltage of 4.5 kV resulted in single fiber deposition. However, this was only possible at Wd ≤ 2 cm, since the threshold that defines the balancing of repulsive charges by the solution cohesive forces was also low.

6.2.4. Scan speed

The velocity at which the fibers were collected was also a parameter of great interest. If the voltage and Wd were maintained constant, the physical behaviour of the jet would be in principle the same. However, the velocity at which the jet or collector moves would influence the amount of fibers reaching a single point on the collector. Consequently, the amount of charges in a single spot can also vary and can determine the consistency of the fiber deposition. We maintained the voltage at 5 kV, Wd at 5 cm, but varied the scan speed between 4 and 10 mm/s to determine its effect on the deposited fibers (Figure S5). At a collection speed of 4 mm/s, the deposited layer was slightly incoherent with respect to its central axis and some fiber dispersion occurs. When the scan speed was increased to 7 mm/s, the accumulation of repulsive charges at a point decreased and the spun layer showed a more consistent pattern, but still presented some fiber dispersion from its core (Figure S5B). At a collection speed of 10 mm/s, the deposited layer showed little deviation from its central axis having comparatively less fiber dispersion (Figure S5C). In summary, as the collector scan speed increased, the line focusing was improved whereas line patterns assumed a proportionally shorter width (Figure 2C). From these data, we conclude that the optimal scan speed was ~10 mm/s.
6.2.5. Ambient parameters

Finally, the ambient parameters for ESP, such as temperature and relative humidity, were also important to tune. Because an inverse relationship exists between temperature and viscosity (crucial for jet elongation as mentioned above)[33] we reasoned that lower temperatures would facilitate jet focusing and relative higher temperatures may cause faster solvent evaporation, and consequent needle blocking by the polymer solution. Therefore, we tested temperatures between 18.5–30 °C and found the optimal temperature is ~20 °C.

Relative humidity within the fairly large range of 25 to 55% allows for jet eruption and focusing, but we needed to optimize this parameter for long fabrication times. The production of large patterns requires the hemispherical drop at the spinneret tip to be maintained in a fluid state, which is facilitated by humidity levels. Furthermore, solutions
containing volatile solvents may dry quickly in dry environments. High humidity also benefits fiber stacking by reducing the charge accumulation. It should be noted that excessive humidity results in melting the fiber morphology due to insufficient solvent evaporation. From the humidity tested, we found that the optimal relative humidity is 40 ± 2%.

![Figure 2](image)

**Figure 2.** (A–C) Influence of the ESP processing parameters on fiber diameter (A and B) and line width (C). Applied voltage (A), working distance (B), and collector speed (C) were varied. (D–F) SEM micrographs of different possible configurations of the fiber bundle within the DW range: (D) Bundle with dispersed fibers (V=5kV and Wd=3cm); (E) Bundle with a core of aligned fibers (V=6kV and Wd=3cm); (F) Single fiber (V=4.5kV and Wd=2cm). Scale bars: D and E (100 µm) and F (50 µm).

### 6.2.6. Optimal ESP parameters

The range of processing parameters that permitted DW ESP of PEOT/PBT is summarized in Table S3. We found that the optimal range of polymer concentration was from 27 to 29% (w/v), which yielded fairly viscous solutions. Wd of ≤ 8 cm were necessary to collect the jet in its stable region, with corresponding optimal voltages of ≤ 8 kV. Flow rates between 0.08 and 0.3 mL/h were necessary and the scan speeds had to be kept above 5 mm/s. The optimal temperature was kept ~20 °C and relative humidity was ranged from 40 ± 2%.

### 6.2.7. Pattern fabrication

Having established the optimal parameters of DW ESP for PEOT/PBT, we proceeded to design scaffolds. The design of a scaffold, such as its architecture and pore network,
determines its success in a targeted TE application. While the architecture may guide and influence cartilaginous tissue formation[34], porosity is essential not only for nutrient diffusion and waste removal[5, 26], but also allows and impacts cartilage ECM formation[35, 36]. Conventional ESP processes lack on-demand patterning ability and typically produce compact meshes which hinder cell infiltration and ECM formation throughout their volume[18].

**Figure 3.** Pattern fabrication with DW ESP. (A and B) SEM micrographs of a 0/90° lay down pattern (A) and detail of a strut intersection (B). (C) Photo of a tympanic membrane design. (D–E) SEM micrographs showing 0/90° lay down pattern with 800 µm pore space with 80 layers (110 ± 9.0 µm thick) (D), 100 layers (140.5 ± 8.9 µm) (E), and 200 layers (190.8 ± 20.8 µm) (F). Scale bars: A and D (1 mm); B (100 µm), C (1 cm) and E and F (100 µm).

DW ESP permits the tailored patterning of 3D ultrathin scaffolds with multiscale and controllable porosity, with a setup that dispenses any complex add-ins. Applying the optimized DW ESP parameters we fabricated regular structures with a 0/90° lay down pattern (Figure 3A-B). The accuracy of this process also allowed the design and fabrication of more complex structures, such one that mimics the anatomy of a tympanic membrane (Figure 3C). By tuning the distance between struts and the number of deposited layers, the pore size and scaffold thickness could be controlled. True 3D structures could be built, where 0/90° scaffolds with an approximate fiber space of 800 µm were fabricated with varying thickness based on the number of deposited layers: 110 ± 9.0 µm with 80 layers (Figure 3D), 140.5 ± 8.9 µm with 100 layers (Figure 3E), and 190.8 ± 20.8 µm with 200 layers(Figure 3F). The fabrication of scaffolds in the centimetre thickness range was hindered by the loss of the macroporous structure. As fibers were
stacked, both the height and the width of the line pattern increased. This is attributed to a higher concentration of fibers on a single spot and consequently the accumulation of repulsive charges, leading to dispersion during the deposition process[37].

To circumvent this issue, some strategies can be implemented. A stepwise approach of the layer-by-layer deposition could provide enough time for the dissipation of the residual charges, hence reducing the repulsive effect and yielding a more focused deposition. However, if the ESP process is interrupted between the fabrication of layers, multiple jet ejections are required. This is a limiting factor, since minor deviations can occur from the focusing spot, causing a mismatch between the deposited layers. However, if the ESP process is continued, the ejection time will be greatly enhanced and further jet instabilities may arise. Another strategy could be the hybridization of this system with a mechanism for charge compensation used in other applications, such as scanning electron microscopy[38, 39]. Ji et al. reported a noticeable charge decrease on Al₂O₃ samples by employing an oxygen atmosphere through a local oxygen pressure device[39]. Finally, another possible solution is to invert the electrode polarity between layer depositions. In such a setup, the residual charges on the deposited fibers would be the opposite of the charges from the upcoming fibers, thus resulting in an attractive effect. Xu et al. followed this strategy and reported an increase in nanofiber deposition accuracy in an NFESP device[37].

6.2.8. Cell seeding and proliferation on DW ESP scaffolds

Electrospun meshes generally facilitate cell attachment and distribution due to their ECM resemblance[6]. To determine whether our DW ESP scaffolds supported cell seeding and proliferation, we fabricated scaffolds with fiber spacing of 860 µm and an average line width of 150 µm, then seeded them with hMSCs (Figure S6C). For comparison, we fabricated scaffolds with similar dimensions by 3D printing (Figure S6B). Seeding efficiency was evaluated by DNA quantification 1 day post-seeding (Figure S6A) and revealed that DW ESP scaffolds promoted 7 times more cell retention than those fabricated by AM techniques (17.66% v. 2.43%). To assess cell distribution and proliferation, the scaffolds were stained with methylene blue after 1 and 4 days of culture (Figure S7). At day 1, cells seeded on the 3D-printed scaffolds were mostly retained at the fiber intersections (Figure S7A), while those seeded on the DW ESP scaffolds were more abundant and well distributed (Figure S7C). At day 4, both scaffolds supported cell proliferation, but a larger population was present on the DW ESP structure (Figure S7B.
These data argue that the DW ESP scaffolds promoted better cell seeding and retention compared to the 3D-printed scaffolds.

6.2.9. Fabrication of an articular cartilage mimetic scaffold

Having established that DW ESP could be used to fabricate scaffolds that support cell seeding and proliferation, we then tested whether this novel technique could be used to create a scaffold that mimics articular cartilage with its distinct multi-zone morphology (Figure 4A); such a scaffold may offer a superior template for *in vitro* cartilage formation and ECM organization[40]. We fabricated a scaffold that recapitulates three of the four main zones of cartilage (Figure 4B-E). The superficial zone, denoted by tangentially oriented collagen fibrils, was represented by two parallel electrospun layers with an average distance of 238 ± 79.3 µm. To reproduce the oblique collagen structure of the middle zone, we created crossed diagonal layers. Finally, the deep zone was represented by parallel layers, radially oriented to the top structure, with an average distance of 523 ± 156.2 µm. The mimetic scaffolds had pore dimensions ranged from 238.0 ± 79.3 to 523.6 ± 156.2 µm of which the values were within the ideal dimensions (250 to 500 µm) reported for chondrocyte proliferation and ECM secretion on gelatin scaffolds[41]. Detailed views of the fiber bundle and individual ultrathin fibers with an average diameter of 1.39 ± 0.6 µm (Figure S8) are shown in Figure 4D and 4E, respectively.

Next, we further characterized the mimetic scaffold’s physical and mechanical properties (Table S4). Overall, the measured porosity was 91.2 ± 1.0% (Table S4), which is fairly large compared to other structures for the same application[34]. The mechanical properties of the scaffolds were also characterized using a nanoidenter. The measured Young’s modulus in compression was 2.65 ± 0.65 MPa, which proved to be higher than human native cartilage as measured by *in situ* biphasic creep indentation (0.45 MPa to 0.80 MPa) or bovine cartilage measured by nanoidentation[42, 43].

In order to engineer a larger tissue construct, four individual scaffolds (each approximately 208.5 ±19.4 µm thick) were stacked and bonded to yield an assembled structure of 1.39 ± 0.6 mm in height. The introduction of sacrificial layers of PEO between the spun mats promoted lateral pore formation, and the bonding process did not affect the inner core of the scaffold (data not shown).
Figure 4. Design of an articular cartilage mimetic scaffold. (A) Illustration of the zonal configuration in native cartilage[44]; (B) CAD design of the devised scaffold; (C) stereomicroscope image of the DW ESP structure, depicting the different zones; (D and E) SEM micrographs of the fiber bundle corresponding to the bottom columns of the scaffold (D) and detailed view of individual fibers at 100× magnification (E). Scale bars: C (1 mm), D (200 µm), and E (2 µm).

6.2.10. Cell culture and chondrogenic differentiation

To investigate the potential of designed scaffolds for articular cartilage regeneration, we cultured hMSCs on stacked DW ESP scaffolds (unless otherwise stated) in chondrogenic and basic medium (negative control medium). As a control, cell-seeded random fiber (RF) scaffolds were also prepared and analyzed. The cell proliferation on the DW and RF seeded scaffolds was evaluated at three time points during the first week of culture, using a Presto Blue assay (Figure S9). Proliferation was registered in all the conditions through an increase in fluorescent signal. In both media, DW scaffolds showed 4-fold higher intensity compared to RF scaffolds, until day 7 when cell proliferation increased steeply on RF sheets. This observation can be explained by the higher number of fibers in the vicinity of cells, due to the lack of large pores on the RF structure, which leads to easier cell spreading, and thus a faster proliferation rate. For chondrogenic differentiation, cells on DW scaffolds showed a steady increase in fluorescence over the 7 days. In contrast, the signal from cells incubated on RF scaffolds in differentiation medium showed little change over the same period. These data indicate that cells proliferated on both RF and DW scaffolds,
To determine the topographical influence of DW mimetic scaffolds on tissue development and matrix secretion, scanning electron microscopy (SEM) observations were performed after 21 days of culture in chondrogenic medium. Whereas the RF and DW scaffolds both demonstrated full coverage of cells, only the DW scaffolds showed cells and matrix throughout its volume (Figure S10).

Figure 5. SEM images showing the topographical influence of the scaffold’s architecture on tissue development (B, D, F) and matrix orientation (C, E, G) after 21 days of culture in chondrogenic medium. (A) Empty scaffold fabricated by DW ESP. Low-magnification images (B, D, F) and high-magnification images (C, E, G) of the superficial zone (B–C), middle zone (D–E), and deep zone (F–G). Scale bars: B, D and F (100 µm), C, E and G (10 µm).

When cultured on the RF mesh, hMSCs dispersed randomly and as a consequence their secreted matrix was isotropically distributed (data not shown). In comparison, the matrix organization was influenced by the zonal topography of the DW scaffolds (Figure 5). The horizontal layers forming the superficial zone (Figure 5C) and the vertical layers of the deep zone (Figure 5G) promoted tissue development along their respective axes, which resulted in the anisotropic formation of fibrils. Coherence analysis of the SEM micrographs showed a slightly higher coefficient of directionality for the deep zone (0.49 ± 0.08) compared to the superficial zone (0.41 ± 0.09) (Figure S11).
The oblique layers that represent the middle zone, induced tissue formation in a more random manner with secreted fibers displaying a crossed pattern (Figure 5E). The coefficient of directionality for this middle zone was 0.33 ± 0.11 (Figure S11). The presence of holes (Figure 5D) suggested the formation of tissue in different and opposite directions. In summary, the DW scaffold imparted a tissue and matrix organization in a zone-dependent way, in contrast to the isotropy seen in the RF sheets.

The morphology and feature size of the fibrils observed in the SEM micrographs suggested the presence of collagen. Among collagen types, type II collagen is the basis for articular cartilage. To investigate the formation of collagen type II, the surface of the scaffolds was immunostained against collagen type II after 21 days of culture in chondrogenic medium, but no signal was detected. hMSC pellets after 21 days in chondrogenic medium showed collagen type II staining (Figure S12C–F), suggesting that pellet conditions (such as greater cell–cell contact, maintenance of cell rounded shape, softer substrate, and reduced oxygen tension), resulted in improved differentiation of hMSCs and consequent production of collagen type II[45-47]. We are investigating whether modifications to the cell culture conditions that provide an environment more similar to the pellet conditions such as using a hydrogel for scaffold coating or a cell carrier like hyaluronic acid (Figure S13) will improve chondrogenic differentiation and collagen I and II production. When cultured on flat substrates, differentiated chondrocytes might dedifferentiate into fibroblast-like cells and produce collagen type I instead[48]. Therefore, we further immunostained the scaffolds for collagen type I (Figure 6). We observed distinct collagen type I staining that resembled the matrix patterning. The RF scaffold revealed random, non-preferential direction collagen type I staining (Figure 6A). For the DW scaffold, collagen type I anisotropy was observed for the top (Figure 6B) and bottom layers (Figure 6D), and an arc-like configuration seen for the middle layer (Figure 6C). These results indicate that the DW scaffold enabled collagen matrix organization in a zone-dependent way in contrast to the isotropy seen in the RF mesh.
Direct Writing Electrospinning of Scaffolds with Multi-dimensional Fiber Architecture for Hierarchical Tissue Engineering

Figure 6. Immunostaining of collagen type I on hMSCs seeded scaffolds after 21 days of culture in chondrogenic medium. (A) RF scaffold; (B–D) Sections of the DW scaffold, as labelled. Scale bar in D (100 µm) applies to all panels. The dashed lines in panels B and D delineate the collagen fiber orientation.

To determine whether the observed patterns in SEM were cartilage-specific matrix, we visualized sulphated glycosaminoglycan (GAG) by staining the scaffolds with Alcian Blue after 21 days of culture. Cultured cell pellets were also stained as a positive control (Figure S12A–B). As shown in Figure S14, both RF and DW scaffolds present higher amount of GAG staining in chondrogenic medium compared to basic medium (control medium). In chondrogenic medium, GAG staining was abundant and distributed all over the RF scaffold (Figure 7A). For the DW scaffolds, we observed GAG staining in a zonal distribution and distinct cell morphology (Figure 7B–D). For the top (Figure 7B) and bottom (Figure 7D) layers of the scaffolds, we observed anisotropic GAG staining and cell elongation along the same direction. This was particularly evident in the top layer, with the presence of some flattened cells on the most superficial zones, which resembled the native state. GAG staining was more randomly dispersed in multiple directions for the
middle section (Figure 7C), and the cells were more rounded compared with the other scaffold zones.

Figure 7. Staining with Alcian Blue for sulphated GAGs and Nuclear Fast Red for cell nuclei in histological sections of hMSCs after 21 days of culture in chondrogenic medium on RF (A) and DW (B–D) scaffolds. (A) 10× view of the RF scaffold; (B–D) 20× view of the top (B), middle (C), and bottom (D) sections of DW ESP scaffolds. The dashed lines represent the direction on the layer (parallel to its axis) and the arrows point to cells. Scale bar in D also applies to panels B and C.

Quantification of GAG production showed similar levels for both scaffolds (Figure 8A). However, cell adhesion and proliferation were significantly greater in DW scaffolds compared to RF sheets (Figure 8B). As a consequence, the RF sheets exhibited significantly higher GAG/DNA ratios (Figure 8C), or more matrix production normalized to cell content, compared to DW scaffolds. Although seeded at the same density (per volume), RF sheets may achieve cell confluency at earlier time compared to DW scaffolds due to their smaller available growth area. Therefore, cells grown on RF
scaffolds were subjected to higher cell–cell interaction throughout the culture period. Reports in literature show that higher cell densities, as a consequence of higher cell–cell interaction, enhance matrix production in cultures with hMSCs[49] and articular chondrocytes[50], which could explain our observed differences. Quantification of GAG and DNA levels after cells achieve full confluency on the scaffolds may help clarify this difference.

Finally, the cell-seeded scaffolds were examined for the presence of several chondrogenic markers, including Collagen type II alpha 1 (Col2a), Sox9 and aggrecan (ACAN). Col2a and ACAN are major proteins found in cartilage matrix[2, 51]. Sox9 is a transcription factor that plays a key role in chondrogenesis[52, 53]. The cell-seeded scaffolds cultured in basic medium were used as control. Levels of mRNA for all three markers were higher for DW scaffolds in chondrogenic medium compared to basic medium (Figure 8D–F). In chondrogenic medium, expression of Sox9 and ACAN was significantly higher in DW scaffolds compared to RF scaffolds after both 14 and 21 days of culture (Figure 8E–F). However, the higher expression of Sox9 and ACAN did not always result in a higher expression of Col2a. Col2a expression was higher for the RF scaffolds compared to the DW scaffolds at day 21 in chondrogenic medium (Figure 8D). This uncoupling of Col2a and Sox9 expression was also reported by Aigner et al. who demonstrated that Sox9 was not the central regulatory mediator of Col2a in human articular chondrocytes[54]. They demonstrated that high gene expression of Sox9 did not lead to an increase of Col2a expression in normal articular chondrocytes and the amount of Col2a expression was higher in spite of reducing Sox9 expression in osteoarthritic chondrocytes. The higher expression of Sox9 in DW scaffolds could be responsible for the higher expression of ACAN in DW scaffolds. Previous studies demonstrated that Sox9 supported the aggrecan promoter activity in clonal chondrocytic cells[53]. These data argue that DW scaffolds supported chondrogenic differentiation of hMSCs.
Figure 8. Cartilaginous matrix production (A–C) and gene expression profiles (D–F) of hMSCs cultured for 14 and 21 days in basic and chondrogenic medium. (A) Total GAG amount; (B) total DNA amount, and (C) GAG normalized to the DNA content. mRNA levels of Col2A (D), Sox9 (E), and ACAN (F) are represented as fold differences relative to the expression obtained in RF samples cultured in basic medium at day 14. * p < 0.05.

6.3. Conclusion

We have shown here the development of a technique for direct writing with electrospinning, with capability to control the fiber bundle morphology and fabricate regular 3D ultrathin structures. Applying this technique, we produced a scaffold that mimics the zonal organization of articular cartilage and compared its performance for cartilage tissue formation to a regular electrospun sheet. In contrast to nonspecific deposition by the typical electrospun mesh, DW scaffolds were able to direct tissue organization and fibril matrix orientation. Expressions of chondrogenic markers, including Sox9 and ACAN, were also significantly enhanced in this structure. Our work provides a new technique for the patterning of nano/microfibrous structures with potentially wide application for the engineering of many tissues, and particularly those presenting regional differences. Further optimization might enable the fabrication of articular cartilage suitable for therapeutic applications.
6.4. Materials and Methods

6.4.1. Polymer solution preparation

A PEOT/PBT block copolymer (PolyVation B.V.; the weight ratio of PEOT/PBT=55/45, the molecular weight (g/mol) of the starting PEG segments used in the polymerization process is 300) was dissolved in a mixture of chloroform (CHCl$_3$, Sigma-Aldrich) and 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP, Biosolve-Chemicals) to prepare the polymer solutions. Solutions of 26 to 30% of polymer (w/v) in 70:30 to 80:20 CHCl$_3$/HFIP (v/v) were formulated.

6.4.2. Electrospinning setup

The ESP system configuration consisted of a two parallel plate setup, composed by a top unit (21.5x30 cm) of a 2 cm thick copper plate encased in Teflon and a 0.5 cm thick copper collector (10x10 cm) mounted on an XY translational stage. The stage is a modified 3D printer (k8200, Velleman), specifically customized for ESP experiments and controllable using Repetier-Host software (v0.84, Repetier).

6.4.3. Electrospinning tests and jet imaging

Initial electrospinning experiments to assess the drawing region were carried with polymer solutions of 26 to 30% PEOT/PBT in 70:30 to 80:20 CHCl$_3$/HFIP (v/v). The applied voltage was varied from 2.0 to 16 kV, the working distance from 2 to 16 cm, the flow-rate from 0.05 mL/h to 2 mL/h, the temperature from 18.5 °C to 30 °C and the relative humidity from 25% to 55%. The scan-speed was varied from 1 to 13 mm/s and spinneret diameter were fixed to 0.5 mm. After jet initiation, the collector was set to a linear motion to collect the fibers in a single direction. A gross visual examination of the outcome and scanning electron microscope (SEM) analysis were done to analyse the contribution of each parameter to the jet stability and to define the parameters range for drawing capability.

ESP tests within the drawing region were carried with a 28% PEOT/PBT in 75:25 CHCl$_3$/HFIP(v/v). The applied voltage varied from 4 to 8 kV, the working distance from 2 to 7 cm and the scan-speed from 1 to 13 mm/s. The flow-rate was set to 0.18 ml/h through a spinneret with 0.8 mm diameter. A temperature of 20°C and a relative humidity of 30% were used as ambient parameters. SEM micrographs were then taken to investigate the fiber bundle morphology within the drawing region.
Images of the jet flight were captured using a digital camera (Panasonic Lumix DMC G3) and the resulting photographs were further treated with ImageJ.

6.4.4. Direct writing and scaffold fabrication

Direct writing experiments and scaffold fabrication were carried with a polymer solution of 28% PEOT/PBT in 75:25 CHCl3/HFIP (v/v) with the experimental parameters listed on Table S5. For the fabrication of the assembled scaffolds, the electrospun mats were stacked via a manual process illustrated in Figure S15. Briefly, the scaffolds were detached from the aluminium foil by placing them in deionized (DI) water for 1 to 2 min and gently tapping the structure with a tweezers to promote detachment. After drying with filter paper, the scaffolds were placed on top of a polydimethylsiloxane (PDMS) block covered with a Teflon sheet, and held in place with thin metal wires. Overall, four scaffolds were stacked together to achieve the final constructs. To promote lateral pore formation, a thin poly (ethylene oxide) (PEO, Mw=900,000 g/mol, Sigma-Aldrich) sheet (300 µm thick) fabricated through solvent casting of 3% PEO in 80:20 H2O/ethanol (v/v), was placed between each scaffold. The stacked layers were then bound together by joining the vertexes of each scaffold and melting them at 180°C (using a micro soldering iron). The assembled constructs were then placed overnight in a container filled with DI water under agitation to dissolve the sacrificial PEO layers. Following this process, the scaffolds were retrieved and let to dry.

The fabrication of random fiber (RF) sheets was carried using the same polymer solution and setup, with the process parameters listed on Table S5, which were selected to fabricate a mesh with similar fiber dimensions as the DW scaffold. The collector was set to a raster scan path with a step size on the Y axis of 1 mm to ensure thickness uniformity on the deposited mesh. Scaffolds with a diameter of 12 mm were then punched out for further cell culture study.

6.4.5. Scaffold characterization

The scaffold geometry and architecture was characterized through measurements on SEM micrographs. The height of the final assembled construct was measured using a digital caliper. The porosity of the electrospun scaffold was calculated using the theoretical approach described in equation 1:
Porosity (%) = \left(1 - \frac{m_{\text{scaffold}}}{\rho_{\text{material}}V_{\text{scaffold}}}\right) \times 100\% \quad \text{Equation 1}

Where $m_{\text{scaffold}}$ is the measured weight of the scaffold, $\rho_{\text{material}}$ is the density of the material (1.2 g/mL)[55] and $V_{\text{scaffold}}$ is the apparent volume of the scaffold (assuming a solid block of equal dimensions). To determine the Young’s modulus under compression of the electrospun scaffolds, a mechanical test was performed using a nanoidentation device (Piuma, Optics 11). The compression test on the scaffold was carried with a tip of radius ($r$) of 22.5 $\mu$m and spring constant ($k$) of 78.3 N/m. 40 indentations were performed with a step size of 50 $\mu$m on the X axis and 5 $\mu$m on the Y axis.

6.4.6. Cell culture

Pre-selected hMSCs (male age 22) were retrieved from the Institute of Regenerative Medicine at Texas A&M University [56, 57]. Briefly, a bone marrow aspirate was drawn from the patient after informed written consent, and mononuclear cells were separated using density centrifugation. hMSCs at passage 2 were expanded in basic medium (BM) composed of $\alpha$-MEM (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (Lonza), 2 mM L-glutamin (Thermo Fisher Scientific), 0.2 mM ascorbic acid (Sigma-Aldrich) and 100 units/mL penicillin plus 100 mg/mL streptomycin (Thermo Fisher Scientific) until reaching 80-85% confluence. The medium was refreshed every 2 days.

6.4.7. Pellet culture

To check the donor potential for chondrogenesis, a pellet culture study was performed over 21 days as a positive control. 2x10^5 cells suspended in 200 $\mu$L of chondrogenic medium composed of DMEM (Thermo Fisher Scientific) supplemented with 1% ITS premix (Micronic BV), 50 $\mu$g/mL ascorbic acid (Sigma-Aldrich), 40 $\mu$g/mL of proline (Sigma-Aldrich), 100 $\mu$g/mL of sodium pyruvate (Life Technologies), 100 units/mL penicillin plus 100 mg/mL streptomycin (Thermo Fisher Scientific), 0.01 $\mu$g/mL of TGF-β3 (R&D systems) and 100 nM dexamethasone (Sigma-Aldrich) were seeded in round bottomed non-treated tissue culture 96-well plate. For pellet formation, the cells were centrifuged at 2000 rpm for 3 min at 4 °C. As a control the cells were seeded in BM using the same protocol. The medium was refreshed every 2 days, taking out 100 $\mu$L and adding 100 $\mu$L. The pellets were fixed in 4% paraformaldehyde for 2 h at 4 °C, and then washed three times with phosphate buffered saline solution (PBS) and maintained wet.

6.4.8. Seeding efficiency and proliferation on 3D-printed v. DW ESP scaffolds
3D printed scaffolds made of PEOT/PBT polymer were fabricated via additive manufacturing (Bioscaffolder, SysENG, Germany) [35]. hMSCs were seeded on DW ESP and 3D printed scaffolds at a density of 5x10^4 cells/cm^2, and cultured in BM during 4 days, with medium changes every other day. Cell attachment was evaluated by DNA quantification using CyQUANT® Cell Proliferation Assay Kit (Thermo Fisher Scientific). Briefly, the scaffolds were washed three times with PBS, transferred to a 1.5 mL Eppendorf tubes and placed at -80 °C overnight. After this, three freeze/thaw cycles (from -80 °C to room temperature) were applied and 250 µL of proteinase K digestion buffer (0.001% (w/v) proteinase K, 10% of 1.85 mg/mL iodoacetamide (Sigma-Aldrich) solution and 10% of 100 µg/mL of pepstatin A (Sigma-Aldrich) solution in Tris/EDTA buffer) was added to every tube, followed by incubation at 56 °C for 16 h under agitation. The lysate was collected and used to measure the DNA amount in the scaffolds using the kit protocol. Cell distribution and proliferation were assessed with methylene blue staining. Briefly, samples were fixed in 10% formalin solution for 30 min, followed by a rinse with demi water for 1 min. After that, samples were put in 1% methylene blue solution for 60s, and washed with demi water again for three times. Images were acquired were a stereomicroscope (SMZ25, Nikon).

6.4.9. Cell seeding and culture on RF sheets and DW ESP scaffolds

Prior to cell seeding, samples were placed in a non-treated tissue culture 24-well plate (O-rings were used to prevent scaffolds floating), sterilized in 70% ethanol for 1hr and washed three times with PBS. After removing the PBS, the scaffolds were pre-wetted in basic medium overnight for next step cell seeding. Cells were seeded on each scaffold at a similar density (12000 cells/mg of material) unless otherwise stated. After cell attachment, a final volume of 1 mL CM was applied for each well. Cell culture on scaffolds in BM was used for negative control. Cell culture on scaffolds was kept for 21 days, and the media refreshed every 2-3 days.

6.4.10. Scanning electron microscopy (SEM) Analysis

The cell-seeded scaffolds (n=4) were washed three times with PBS, fixed during 2 h in 10% neutral buffered formalin, washed again three times with PBS, dehydrated in a series of ethanol dilutions (1x 50, 60, 70, 80, 90 and 96% and 2x 100%) for 30 min and then critical point dried with liquid carbon dioxide (Balzers CPD 030 and Leica EM CPD 300). Following this, samples were gold sputtered (Cressington Sputter Coater 108 auto).
for 40s at 30 mA and then imaged using a SEM (Philips XL-30 ESEM, Philips; and Versa 3D, FEI) at a V=10 kV. For analysis of the fiber diameter and morphology, and characterization of the electrospun scaffolds, the samples were gold sputtered and imaged using the same parameters.

6.4.11. Metabolic Activity

The cell metabolic activity was measured at day 1, day 4 and day 7 using the PrestoBlue® Cell Viability Reagent (Life Technologies). Briefly, a 10% (v/v) presto blue solution in culture medium was added to each sample (n=4) and incubated at 37 °C, 5% CO₂ during 2 h, protected from light. After that time, 150 µL aliquots were transferred to a black 96-well plate and the fluorescence measured in a Perkin Elmer Victor3 plate reader (Perkin Elmer) using the following parameters for Ex/Em: 560/590 nm. Blanks of medium only were made to correct for background fluorescence.

6.4.12. Real-time quantitative polymerase chain reaction (qPCR)

Gene expression of the chondrogenic markers including Col2a, Sox9, and ACAN was evaluated after 14 and 21 days of culture. Total RNA was isolated from the samples using a TRIzol based extraction method followed by the Isolate II RNA Mini Kit (Bioline). Briefly, 1 mL of TRIzol reagent (Thermo Fisher Scientific) was added to each sample, followed by centrifugation at 12000 rcf for 2 min at 4 °C, addition of 0.2 mL of a CHCl₃: isoamyl alcohol mixture and again centrifugation at 12000 rcf for 15 min at 4°C. The aqueous phase was then collected and the rest of the procedure was conducted using the Isolate II RNA Mini kit protocol. RNA quantification and purity assessment were done in a NanoDrop 2000 (Thermo Fisher Scientific). cDNA was synthesized using the SensiFAST™ cDNA Synthesis Kit (Bioline) according to the manufacturer’s protocol. qPCR was performed in the CFX Connect™ Real-Time PCR Detection System (Bio-Rad, UK) using the SensiMix™ SYBR® & Fluorescein Kit (Bioline) and the primers listed in Table S6. 25 µL of sample volume composed of 3 µL of cDNA, 12.5 µL of kit reagent, 7.5 µL of distilled water and 1 µL of forward and reverse primer were used in the amplification process. For this, cDNA was first denatured for 10 min at 95 °C, followed by 45 cycles, consisting of 15 s at 95 °C, 15 s at 60 °C, 15s at 72 °C. Ct values were determined and normalized to the housekeeping gene B2M and ΔCt ((average of Ct-control) – (Ct-value)). The relative mRNA of the markers of interest is represented as fold changes relative to the
expression obtained in the random fiber samples cultured in basic medium and calculated by $2^{-\Delta \Delta Ct}$.

6.4.13. Immunohistochemistry

Samples (n=3) were retrieved after 21 days of culture, washed three times with PBS and fixed for 30 min in 4% paraformaldehyde, at 4 °C. Samples were permeabilized in 0.5% Triton X-100 for 5 min at room temperature, followed by incubation with blocking buffer (0.05% Tween 20, 5% goat serum, and 1% BSA in PBS) overnight. Afterwards, a mouse monoclonal anti-collagen type I (1:100; Thermo Fisher scientific) or a mouse monoclonal anti-collagen type II (1:50; Novus Biologicals) were added and incubated overnight again at 4 °C. The following day, samples were washed three times with wash buffer (blocking buffer without goat serum) and incubated with goat anti-mouse Alexa Fluor 647 secondary antibody (Invitrogen, USA) (1:300) in wash buffer for 1.5 h at RT. Cell nuclei were stained with DAPI (0.1 µg/mL in PBS) for 5 min, followed by washing in PBS (three times). Samples without primary antibodies were used as negative control. Fluorescent images were acquired using an inverted fluorescent microscope (Nikon Eclipse Ti-S).

6.4.14. Glycosaminoglycan (GAG) assay

GAG and DNA measurements were performed after 14 and 21 days of culture. Samples (n=4) were washed with PBS, transferred to a 1.5 mL Eppendorf tube and stored dry overnight at -80 °C. Three freeze/thaw cycles (from -80 °C to room temperature) were applied to all the samples and 250 µL of proteinase K digestion buffer (the formulation as describe above) were added to each tube, followed by incubation at 56 °C for 16 h, under agitation. The lysate was collected and used to measure the GAG amount and respective DNA content. The GAG amount was determined spectrophotometrically after reaction of 25 µL of lysate mixed with 5 µL of NaCl solution (2.3 M) with 150 µL of dimethylmethylene blue dye (DMMB, Sigma-Aldrich). The intensity of colour change was quantified immediately by measuring the absorbance at 525 nm in a microplate spectrophotometer (CLARIOstar, BMG Labtech, Germany). The GAG content was then calculated using a standard of chondroitin sulphate (Sigma-Aldrich). The DNA amount was determined following the protocol described above.

6.4.15. Histology

Pellets and cell-seeded scaffolds (n=3) were retrieved after 21 days of culture, washed with PBS and fixed overnight in 10% neutral buffered formalin, at 4°C. Subsequently,
samples were dehydrated through a graded ethanol series (1x 50, 60, 70, 80, 90 and 96% and 2x 100%), cleared with butanol (2 h), infiltrated with liquid paraffin (overnight), embedded in paraffin and stored at -20 °C. Sections with 10 µm in thickness were cut with a microtome (HM355S, Thermo Fisher Scientific), mounted in poly-L-lysine pre-coated slides and stained with 0.5% (w/v) Alcian Blue solution in 1M HCl (pH =1.0, Sigma-Aldrich) for sulphated GAG and with Nuclear Fast Red for cell visualization. Microphotographs were then acquired using an inverted fluorescent microscope (Nikon Eclipse Ti-S) equipped with a colour camera (DS-Ri2).

6.4.16. Image analysis

SEM images were analysed for fiber alignment quantification using the ImageJ software. The plugin OrientationJ was employed for this purpose and the images were analysed over the same ROI. Measurements (n=30) were performed per condition and the results are expressed in terms of directional coherency coefficient, in a scale from zero to one. A coherency coefficient close to one, represented as a slender ellipse, indicates a strongly coherent orientation of the local fibers in the direction of the ellipse long axis. A coherency coefficient close to zero, represented geometrically as a circle, denotes no preferential orientation of the fibers[49].

6.4.17. Statistics

Statistical significance was assessed with an unpaired t test on the fiber diameter study. A p-value<0.05 was considered significant. For the biological study, statistical significance was assessed using non-parametric tests. For comparison between two groups, the Mann-Whitney U test was used and for comparison between three groups, the Kruskal-Wallis test was applied. A p-value<0.05 was considered significant. All the values are expressed as the average ± standard deviation.

Acknowledgements

H. Chen thanks China Council Scholarship programme for financial support (Grant # 2011614016)
References


Direct Writing Electrospinning of Scaffolds with Multi-dimensional Fiber Architecture for Hierarchical Tissue Engineering


[26] N. Abdul Rahman, Electrospun Conducting Polymer Nanofibers for Biomedical Applications, ResearchSpace@ Auckland, 2012.


Supplementary Information

1. Materials and Methods: Polymer Solution Characterization

1.1. Viscosity

The viscosity was measured using a viscometer (Brookfield DV-E, Brookfield Engineering). Solution (8mL) was loaded into the device’s container and a cylindrical spindle was used and set to 30, 50, and 100 rpm. Three different measurements were taken for each rotational speed and the final viscosity value was averaged from these results.

1.2. Surface Tension

The evaluation of the polymers solution’s surface tension was carried out using an optical contact angle measuring system (OCA 20, Dataphysics) at 25°C. Since the organic solvents in the solution have high evaporation rates, the preferred method for surface tension measuring was an axisymmetric drop shape analysis (ADSA) of an air bubble/polymer solution interface. ADSA is a powerful tool for fast, accurate and reliable measurements of fluid-liquid interfacial tensions. ADSA is based on the numerical fit between the shape of an experimental bubble (or drop) and the mathematical model given by the classical Laplace equation of capillarity[58]. In this procedure, a closed container was filled with the polymer solution and a capillary (with the tip bent upwards) was dipped in it. An air bubble was precisely dosed through the capillary and the evolution of the drop shape and contact angle was automatically analysed and used to extrapolate the surface tension. The surface tension of deionized (DI) water at 25°C (γ=71.99 ± 0.005) was used as an accuracy control.

1.3. Conductivity

Conductivity measurements were obtained using an impedance spectroscopy-based method. Impedance spectroscopy provides the relation between the real (Zr) and imaginary (Zi) impedance of a sample, as a function of the applied frequency, and the results can then be graphically expressed in the Nyquist plot, as Zr v. Zi. The impedance of a resistance is independent of the frequency, and thus it has no imaginary part [25]. Hence, it is possible to determine the resistance of a solution from a Nyquist plot by fitting
the resulted curve and calculating the impedance when the imaginary part is zero. Once determining the resistance (R), the electrical resistivity of a solution (ρ) can be determined using the relation \( R = \rho \frac{L}{A} \), where L is the distance between the electrodes and A is the area of the electrode. To determine these parameters precisely, solutions of known conductivity (σ) can be used to calibrate the system, since \( \sigma = \frac{1}{\rho} \). Potassium chloride (KCl) solutions of 1M, 0.1M, 0.01M, 0.001M and 0.0001M were then prepared and the resistance measured. The results were fitted into a curve and the calibration factor of the system (L/A) could then be determined. Finally, to obtain the polymer solution conductivity, the resistance was first measured through the Nyquist plot fit and the conductivity calculated using the calibration factor. The impedance spectroscopy measurements were carried using the Autolab software (Metrohm Autolab B.V.) and using a setup built in-house.

2. Figures

Figure S1. Influence of the processing parameters on the deposited line consistency. This gross examination was used to identify the direct writing region. (A–C) Effect of different flow rates at a voltage (V) is 8 kV and working distance (Wd) of 8 cm; (D–F) Effect of the voltages at Wd=8 cm and flow-rate is 0.08 mL/h; (G–I) Effect of different working distance at V=8 kV and the flow-rate is 0.08mL/h Scale bars: A–I (500 µm).
Figure S2. Point pattern created when the solution is let to spin at a single point for a prolonged period. (A) Digital camera image; (B) SEM micrograph. Scale bars: A (5 mm) and B (1 mm).

Figure S3. Bundle morphology resulting from different voltages: (A) 4 kV, (B) 5 kV and (C) 6 kV. The working distance was maintained at 5 cm, the flow-rate at 0.18 mL/h and the scan-speed at 10 mm/s. Scale bars: A–C (100 µm).

Figure S4. SEM images showing the effect of working distance on the deposited fibers at (A) Wd = 3 cm, (B) Wd = 4 cm and (C) Wd = 5 cm. The applied voltage was set at 5kV, the flow-rate at 0.18 mL/h and the scan speed at 10 mm/s. Scale bars: A–C (100 µm).
Figure S5. Effect on the line pattern at different scan speeds (A) 4 mm/s, (B) 7 mm/s and (C) 10 mm/s. The voltage was set at 5 kV, the working distance at 5 cm and the flow-rate at 0.18 mL/h. Scale bars: A–C (100 µm).

Figure S6. (A) Seeding efficiency of human mesenchymal stromal cells (hMSCs) on (B) 3D printed scaffolds and (C) direct-written (DW) electrospun (ESP) scaffolds. Efficiency measured as attached cells relative to total cells seeded on scaffolds and expressed as mean ± SD. Scale bars: B–C (500 µm).
Figure S7. Methylene Blue staining on (A and B) 3D-printed and (C and D) DW ESP scaffolds after (A and C) 1 day and (B and D) 4 days of culture in basic medium. Both scaffolds have the same pore space of 860 µm and an average line width of 150 µm, although the DW scaffolds appear larger due to the fiber protrusions. Scale bars represent 500 µm.

Figure S8. Frequency distribution of individual fiber diameters from the articular cartilage mimetic scaffold.
Figure S9. Presto blue measurements of the DW and RF seeded scaffolds taken at 3 times points during the first week of culture.

Figure S10. SEM micrographs of the cell-seeded scaffolds RF (A) and DW (B, C) after 21 days in culture on chondrogenic medium. (C) cross-section of the DW scaffold shows cells and matrix grow inside the scaffolds. Scale bars: A and B (1 mm), and C (500 µm).

Figure S11. Directional coherency of the fibril matrix in the different sections of the DW ESP scaffold. 0 – no coherency; 1 – full coherency (full alignment). *p < 0.05; ***p < 0.001.
Figure S12. Chondrogenic differentiation of the hMSCs cultured in a pellet form in differentiation (left column) or control (right column) medium for 21 days. (A and B) Alcian Blue staining; (C and D) Collagen type II immunostaining; (E and F) DAPI staining. Scale bars: A–F (100 µm); scale bar in F applies to C–E panels.
Figure S13. (A) Schematic of the process of scaffold embedding with a hyaluronic acid (HA) hydrogel and (B) photograph of such construct. Scale bars: 1 cm.

Figure S14. Alcian Blue and Nuclear Fast Red staining of histological sections of RF(A and B) and DW ESP (C and D) scaffolds after 21 days culture in basic (A and C) and chondrogenic medium(B and D). Scale bar in D represents 50 µm and applies to all panels.
Figure S15. Scaffold mounting process: (A) The scaffold is placed on top of a PDMS block and fixed with thin metal bars; (B) a thin PEO sheet is then placed directly on top of the scaffold; (C) the process is repeated until 4 mats are assembled. The scaffold vertexes are then joined together and the sacrificial layers are dissolved in DI water.

3. Tables

Table S1. Summary of some physical properties from a solution of 28% PEO55PBT45 dissolved in 75:25 Chloroform:HFIP.

<table>
<thead>
<tr>
<th>Physical property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity (cP)</td>
<td>326.2 ± 2.9</td>
</tr>
<tr>
<td>Conductivity (µS/cm)</td>
<td>4.7 ± 0.2</td>
</tr>
<tr>
<td>Surface Tension (mN/m)</td>
<td>28.0 ± 0.7</td>
</tr>
</tbody>
</table>
### Table S2. Summary of the obtained fiber morphologies with varying working distances and voltages. The values in bold on the voltage column represent the minimum voltage for jet initiation. Once the jet starts to spin, the applied voltage can be reduced.

<table>
<thead>
<tr>
<th>Working Distance (cm)</th>
<th>Voltage (kV)</th>
<th>Focused spinning?</th>
<th>Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td>2</td>
<td>3.5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>+</td>
<td>RF/AC</td>
</tr>
<tr>
<td></td>
<td>4.5</td>
<td>+</td>
<td>SF</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>3.5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>4.5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>5.5</td>
<td>+</td>
<td>RF/AC</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>+</td>
<td>AC</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>3.5</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>4.5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>+</td>
<td>RF</td>
</tr>
<tr>
<td></td>
<td>5.5</td>
<td>+</td>
<td>AC</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

- **N/A, no spinning, no fiber**
- **RF, Bundle of random fibers**
- **+, yes**
- **AC, Bundle of fibers with aligned core**
- **–, no**
- **SF, Single fiber**

### Table S3. List of processing parameters that permitted DW ESP of PEOT/PBT.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Polymer Solution</strong></td>
<td></td>
</tr>
<tr>
<td>▪ Concentration (w/v, %)</td>
<td>27–29</td>
</tr>
<tr>
<td>▪ CHCl₃:HFIP ratio (v/v)</td>
<td>74:26–78:22</td>
</tr>
<tr>
<td><strong>Process</strong></td>
<td></td>
</tr>
<tr>
<td>▪ Working distance (cm)</td>
<td>≤ 8</td>
</tr>
<tr>
<td>▪ Voltage (kV)</td>
<td>2.5 ≤ V ≤ 8</td>
</tr>
<tr>
<td>▪ Flow-rate (mL/h)</td>
<td>0.08 ≤ Q ≤ 0.3</td>
</tr>
<tr>
<td>▪ Scan-speed (mm/s)</td>
<td>≥ 5</td>
</tr>
<tr>
<td><strong>Ambient</strong></td>
<td></td>
</tr>
<tr>
<td>▪ Temperature (°C)</td>
<td>18.5–21</td>
</tr>
<tr>
<td>▪ Relative Humidity (%)</td>
<td>28–44</td>
</tr>
</tbody>
</table>
**Table S4.** Characterization of the articular cartilage mimetic scaffold

<table>
<thead>
<tr>
<th>Features</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individual scaffold thickness (µm)</td>
<td>208.5 ± 19.4</td>
</tr>
<tr>
<td>Mounted scaffold thickness (mm)</td>
<td>1.39 ± 0.6</td>
</tr>
<tr>
<td>Fiber diameter (µm)</td>
<td>1.39 ± 0.6</td>
</tr>
<tr>
<td>Pore Space Superficial zone (µm)</td>
<td>238.0 ± 79.3</td>
</tr>
<tr>
<td>Pore Space Deep zone (µm)</td>
<td>523.6 ± 156.2</td>
</tr>
<tr>
<td>Porosity (%)</td>
<td>91.2 ± 1.0</td>
</tr>
<tr>
<td>Young’s Modulus in compression (MPa)</td>
<td>2.65 ± 0.65</td>
</tr>
</tbody>
</table>

**Table S5.** List of ESP parameters used for scaffold fabrication.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>DW scaffold</th>
<th>RF sheet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voltage (kV)</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Working distance (cm)</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Flow-rate (mL/h)</td>
<td>0.12</td>
<td>0.4</td>
</tr>
<tr>
<td>Scan-speed (mm/s)</td>
<td>35–50</td>
<td>5</td>
</tr>
<tr>
<td>Spinneret diameter (mm)</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>18–21</td>
<td>20</td>
</tr>
<tr>
<td>Relative Humidity (%)</td>
<td>38–43</td>
<td>30–32</td>
</tr>
</tbody>
</table>

**Table S6.** List of qPCR primers sequence.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward</th>
<th>Reverse</th>
</tr>
</thead>
<tbody>
<tr>
<td>B2M</td>
<td>5’-ACAAAGTCACATGGTTCACA</td>
<td>5’-GACTTGTCTTTCCAGCAAGGA</td>
</tr>
<tr>
<td>Col2a</td>
<td>5’-CGTCCAGATGACCTTCTACG</td>
<td>5’-TGAGCAAGGGCCCTCTTGGAG</td>
</tr>
<tr>
<td>Sox9</td>
<td>5’-TGGGCAAGCTTGGAGACTTC</td>
<td>5’-ATCCGGGTGGTCTTTGTG</td>
</tr>
<tr>
<td>ACAN</td>
<td>5’-AGGCAGCGTGATCCCAACC</td>
<td>5’-GGCCTCTCCAGTCTCATTTC</td>
</tr>
<tr>
<td>ALCAM</td>
<td>5’-ACGATGAGGCAGACGAGATAAGT</td>
<td>5’-CAGCAAGGAGGAGCCAACAAAC</td>
</tr>
</tbody>
</table>
Bibliography

General Discussion

Despite it was created in the middle of 1990s, the field of regenerative medicine is expanding fast [1]. Whereas regenerative medicine first aimed at developing biological constructs that restore damaged or diseased tissues and organs [1-3], it is more and more progressing towards a field where the engineered biological constructs could potentially enhance tissues and organs. To this end, stem cells are often seeded onto an artificial scaffold, which is designed to facilitate cell reorganization and facilitate cell differentiation to form a functional tissue [4]. Traditional regenerative medicine scaffolds are simply biocompatible and biodegradable, which do not fairly resemble the cell microenvironment. The extracellular matrix (ECM) in the human body is dynamic, hierarchical and tissue specific [5]. As a consequence, there is a need to develop bioactive scaffolds that could instruct basic cellular functions like adhesion, proliferation, and differentiation [4]. Such bioactive scaffolds can be constructed via designing and controlling the chemical, physical and mechanical properties of scaffolds. Physical and mechanical cues including surface topography, elasticity and structural features can regulate cell functions such as the above mentioned [6]. The chemistry of engineered scaffolds, which determine their surface chemistry, mechanical properties and biodegradable rate, are dependent on the class of materials [2]. Thus, an engineered scaffold with well-designed material chemistry, structural and mechanical properties is essential to steer the regenerative potential of stem cells [6, 7]. Among current fabrication methods, electrospinning remain a unique and effective technique to fabricate continuous nano or micro fibers. Due to its capacity to produce fibrous scaffolds resembling the features of ECM, electrospinning has been intensively investigated in regenerative medicine [8].

The aim of this thesis is to make a contribution in creating bioactive scaffolds for regenerative medicine and regenerative medicine through electrospinning-based techniques. To achieve this goal, here we mainly focus on designing new materials, tuning surface topography and developing new manufacturing techniques. Chapter 2 and chapter 3 show functional electrospun fibers were fabricated from new materials that are
a new class of poly(butylene succinate)-based (PBS-based) polyesters and yttrium-stabilized zirconia (YSZ) ceramics, respectively. In chapter 4, electrospun fibers encoded with different surface topographies were achieved by simply varying humidity during electrospinning. In chapter 5, we have proposed a new approach to fabricate crimped electrospun fibers, which mimics the structural properties of collagenous fibers in different kinds of connective tissues, by using the principles of mechanical buckling through thermal shrinkable materials. Finally, in chapter 6, we further developed a new solution electrospinning technique which allows depositing fibers in predetermined patterns, in a controlled manner. This new direct writing electrospinning technology allows bridging the gap between additive manufacturing technologies and melt electrospinning and enabling to fabricate scaffolds with a more closely degree of biomimicry compared to native tissues. In chapter 7, a general discussion of the results presented in this thesis is hereafter elaborated and divided in the following sections.

7.1. Fabrication of nanofibrous scaffold from PBS-based polyesters

An essential requirement for ideal scaffolds in regenerative medicine is that they should have suitable mechanical properties and degradation rate to match tissue regeneration dynamics. Although commercial polyesters such as poly(ɛ-caprolactone) (PCL), poly(lactic acid) (PLA), poly(glycolic acid) (PGA) and copolymers (PLGA) have been broadly explored in regenerative medicine, they show limited flexibility in tuning mechanical properties and degradation rate. In addition, they lack of functions that can render chemical cues to steer cell differentiation. These disadvantages limited their application in regenerative medicine. Therefore, there was considerable demand for new polyesters. Commercial polyester named poly(butylene succinate) (PBS) could serve as a possible alternative to the above mentioned aliphatic polyesters due to its good thermal stability, proven biodegradability and biocompatibility, and low cost of raw materials [9, 10].

Chemical and physical modification of PBS may improve its performances with respect to the intended application. In chapter 2, functional electrospun scaffolds were fabricated from a class of multiblock PBS-based copolyesters containing either butylene thiodiglycolate (BTDG) or butylene diglycolate (BDG) sequences. These new polyesters displayed controllable wettability, hydrolysis and mechanical properties rate depending on the molecular architecture and on the kind of etheroatom introduced along the polymer backbone. The polymers containing BDG were more hydrophilic than those containing BTDG units attributed to the higher negative electric charge of oxygen atoms with respect
to sulphur ones. The hydrolytic degradation rate of these scaffolds can be interrupted by crystallinity degree and surface hydrophilicity [11]. Copolymers containing BDG sequences degraded to a higher extent than those containing BTDG ones, due to the higher hydrophilicity of the former. The copolymer P(BS\textsubscript{11}BDG\textsubscript{11}) degraded faster than its corresponding blend of PBS/PBDG, because of the lower crystallinity degree. P(BS\textsubscript{10}BTDG\textsubscript{10}) and its corresponding blend of PBS/PBTDG, they underwent a similar degradation profile because of their comparable crystallinity degree and hydrophilicity. Previous studies have demonstrated that the chemical composition has a strong relationship with mechanical properties, degradation rate and hydrophilicity of the resulting scaffolds. For example, fibrin gels were reinforced with collagen and glycosaminoglycans to improve their mechanical strength [12]. The degradation rate of PLA can be increased by the incorporation of glycolide components in the polymer backbone chains through copolymerization [13]. The hydrophobicity of PLGA scaffolds can be physically modified by blending with the hydrophilic poly(ethylene glycol) [14]. The chemical composition of scaffolds is also known to be one of the key mediators of manipulating cell behavior such as cell attachment, proliferation, and differentiation [15, 16]. In the work by Chang et al., type II collagen and chondroitin sulfate were effectively grafted on PCL porous scaffolds to optimize the biocompatibility of the system [17]. They demonstrate that these modified scaffolds promote type II collagen and GAGs production from the seeded chondrocytes. Amino-functionalized coating surface are most favorable for osteogenic and adipogenic differentiation of human mesenchymal stem cells as compared to OH-functionalized, COOH functionalized, and CH\textsubscript{3}-functionalized coating [15]. Here, we demonstrated that pure PBS polymers enhanced early osteogenic differentiation of hMSCs and the additional presence of ether units (BDG) resulted in enhanced mineralization. Vice versa, the presence of BTDG units in polymer chains enhanced the differentiation of human mesenchymal stromal cells (hMSCs) into chondrocytes while this chemistry inhibits their osteogenic differentiation.

Overall, these results highlight that scaffold chemistry is critical to its function and important to influence cell activity. Thus, chemical properties of scaffolds should be taken into stronger consideration when designing scaffolds for regenerative medicine application.
7.2. Fabrication of bioactive nanofibrous scaffolds via tuning structure and texture

Besides polymers, ceramics are also an important material candidate for regenerative medicine. Although ceramics including hydroxyapatite (HA) and tricalcium phosphate (TCP) have been widely used as scaffolds for skeletal tissue regeneration, the principle drawback of these materials is their brittleness. Among the ceramics so far studied, zirconia is widely used for orthopedic and dental implantation due to its excellent biological and mechanical properties [18]. However, zirconia is a bio-inert material, which restricts its application in regenerative medicine [19]. To overcome the aforementioned problems, in chapter 3, novel paper-like flexible nanofibrous YSZ scaffolds were prepared via electrospinning a mixture of YSZ precursors and polyvinylpyrrolidone (PVP), followed by annealing the green fiber in either a conventional oven or microwave. Electrospun silica-based mats were flexible than crystalline materials owing to the amorphous nature of the material [20]. However, these new YSZ nanofiber mats, which were crystallized, showed remarkable flexibility compared to traditional ceramics. The mats could withstand a total deflection of 180° bending and cycled at least 1000 times without mechanical failure. This phenomenon may be attributed to the long length of the fabricated nanofibers leading to physical entanglement. Nevertheless, individual nanofibers offer stiffness values close to bulk zirconia, 5 orders of magnitude stiffer than the mat. Live and dead assay showed high cell viability on the developed nanofibrillar scaffolds. Gene expression and ALP activity of seeded hMSCs was promoted by the nanofibrous structure of zirconia scaffolds compared to the bulk ceramic. Interestingly, significant higher amount of calcium content was observed on the scaffolds, in contrast to the well-known bioinert behavior of bulk zirconia. These results suggested that electrospun fiber matrices provide a more suitable environment for osteogenic differentiation of hMSCs. This difference could be ascribed to their structural features, since their scaffold chemistry and culture condition were the same. The morphology and fiber dimension of electrospun fibers are comparable with those observed in the ECM of human. Thus, electrospun nanofibrous scaffolds provide improved environments for cell responds including cell attachment, proliferation, and differentiation [21]. Hae-Won Kim et al. demonstrated bone marrow derived stem cells had enhanced osteogenic potential on glass nanofibers than glass disks due to their nanofibrous structure [22].

Apart from structural features elaborated above, other physical cues can also be used as potent regulators of cell behavior such as surface topography and stiffness of scaffolds, mechanical force (like shear stress imposed by culture media) and external force (like
light signal) [7, 23]. Recently studies have shown that the design of surface topography of materials can be used for controlling stem cells differentiation [24, 25]. Increasing the surface roughness and surface porosity of materials causes an improvement in cellular attachment and differentiation [26]. In Chapter 3, we demonstrated that the modulation in surface roughness of zirconia ceramic fibers could be achieved by applying different sintering mechanisms. Nanofibers annealed in a conventional oven showed a surface roughness of 0.7 ± 0.2 nm while nanofibers annealed in a microwave oven had a roughness of 0.4 ± 0.1 nm. A step-bunching effect (i.e., aggregation of the ceramic steps), sudden shrinkage and densification, and/or improved oxygen transport can be used to explain the formation of smooth surface when annealed in MW [29-31]. However, modulating the heating rate did not show obvious influence on surface morphology among samples that were annealed by the same heating mechanism. It was worth to notice that the grain sides of samples depended on heating rates of annealing and heating mechanism. Using a lower heating rate would cause larger grain sizes due to the longer effective sintering times. Conversely, the samples annealed in a MW showed a smaller grain size than the one annealed in a CO using the same heating rate. A similar result was reported by Xie et al. who demonstrated smaller and more uniform grain sizes for YSZ when annealed in a MW [32]. It has been demonstrated that surface topographical cues influence cellular activity such as adhesion [27, 33], migration [34], proliferation [27], and differentiation [35, 36]. The results of mineralization assay revealed that the rougher scaffolds annealed in CO promoted higher calcium content compared to the MW annealed smooth scaffolds (0.4 ± 0.1 nm). This finding was consistent with the previous work where a rougher titanium surface promoted matrix mineralization of osteoblasts compare to smooth titanium surface.

As a logical consequence of the findings of chapter 3, the influence of surface roughness of electrospun fibers on cell activity was further investigated on polymeric scaffolds (chapter 4). By varying the relative humidity during electrospinning, it was possible to make fibers with different surface roughnesses, yet keeping constant other scaffolds properties including fiber dimension, scaffold pores and porosity, and scaffold chemistry. Another advantage of this approach lies in a homogeneous modification, which is applied to every single fiber of the scaffolds rather than happening on the surface area of scaffolds. In chapter 4, electrospun fibers having varied surface roughness were fabricated from copolymer PEOT/PBT under different humidity condition. The surface roughness (Ra) of obtained fibers increased from 14.3 nm to 71.0 nm when the humidity increased from
20% to 70% during electrospinning. These results are in line with the work reported by Casper et al. where increasing humidity caused an increase of pores on the surface of polystyrene fibers [37]. Adsorption is a surface phenomenon. The amount of adsorbed protein on a surface is dependent on the surface roughness and chemistry [38]. The fibers fabricated from 70% relative humidity had a higher protein adsorption compared to the one obtained from 20% relative humidity due to the higher surface roughness of the former one. In addition, surface energy is also influenced by surface topography. Fibers having a surface roughness of 71.0 nm showed a different surface wettability compared to the one with a surface roughness of 14.3 nm. Similar results were reported by Chuah et al. who demonstrated that increasing the surface roughness caused variation in surface tension.

hMSCs were cultured for 21 days on scaffolds with different surface roughnesses. We found that the influence of surface roughnesses on hMSCs osteogenesis was medium dependent. The control medium (basic medium) showed much more pronounced effect compared to the differentiation medium (osteogenic medium). In basic medium, some specific osteogenic genes (OPN and BMP2) displayed higher expression on a higher surface roughness (Ra=71.0 nm), whereas other genes, including BSP, COL1A1 and OCN, showed higher expression on a lower surface roughness (Ra=14.3 nm). In osteogenic medium, the result was not conclusive. In addition, a higher surface roughness (Ra=71.0 nm) had higher ALP activity than a lower surface roughness (Ra=14.3 nm). A recent study demonstrated that a surface composed of nano and sub-micron mixed features promote better osteogenic differentiation of hMSCs compared to the surface only containing singular features (nano or sub-micron scale) [39]. This can be used to explain the finding that both lower and higher roughnesses here investigated supported osteogenic differentiation of hMSCs. In the case of hMSCs chondrogenesis, a lower surface roughness (Ra = 14.3 nm) showed to better support the chondrogenic differentiation of hMSCs at day 7 compared to higher surface roughness (Ra = 71.02 nm) in differentiation medium. At a protein level, no significant difference in GAG content was found among the roughnesses tested in the present study.

### 7.3. Developing new manufacturing techniques

One of the principles in regenerative medicine is that scaffolds should resemble the ECM of the tissues to be replaced. There are numerous types of ECM in human tissues and their functions are tissue-specific. For example, collagen fibers, one of the main components of ECM, present crimped structure in several human tissues, such as tendon, ligament,
muscle, and dermis. In these tissues, the wavy collagen structure gives its tissue structural and mechanical properties. For instance, the wavy collagen fibers in tendon, which are highly resistant to large stretching, is associated with the high tensile strength of tendons. Despite electrospinning has been extensively used in regenerative medicine, little progress has been made for fabricating crimped fibers mimicking the fibril pattern of natural tissues. In chapter 5, we developed a new technique for generating wavy patterns in electrospun fibers. This new technique comprise three steps: first, a construct was achieved by depositing electrospun fibers on a thermally shrinkable material; second, by heating the construct above the glass temperature of the shrinkable material results in shrinkage, thereby buckling the adhered fibers; third, by cooling down the construct at room temperature, the curl pattern is retained. In general, the developed technique is a simple and effective approach to fabricate crimped fibers and showed following advantages compared to previous studies [40-42]: i) the deposited fibers adhere to thermally shrinking film without extra bonding process; ii) the buckled fiber patterns are retained by cooling down at room temperature without extra fixating agents or methods; iii) a wide range of crimped patterns can be achieved by tuning the processing conditions; iv) the capability to create multiscale crimped patterns spanning from fibers to fiber meshes by controlling the thickness of fiber layers, the depositing time of fibers and the shape or geometry of the thermo-shrinkable materials.

Cellular migration is one of the major concerns for electrospun scaffolds in terms of their application in regenerative medicine. hMSCs present a more even distribution and cellular infiltration on wavy scaffolds compared to non-wavy scaffolds. This could be attributed to the wavy patterns which increase of free volume inside the scaffold. The wavy scaffolds have a pattern mimicking the natural configuration of connective tissues, such as tendons and ligaments. It was demonstrated that such topographical pattern supported the expression of TGF-β which is a key signal pathway related to connective tissue development. Some further investigations are still necessary to evaluate if the same trend could be observed on more specific cell populations than the mink lung epithelial cell line used on the scaffolds in this study. The present technique in chapter 5 is simple and universal, which can also be translated to other polymers and thermo-shrinkable materials. We have shown successful translation to polymers including PVA, PAN and PS through using PLA shrinkable films. In addition, PEOT/PBT and PAN fibers having wavy patterns were also obtained using PS biaxial shrinkable films. Thus, the technique shown in chapter 5 could serve as a technical platform to fabricate functional scaffolds that could be
beneficial for mimicking a variety of tissues organization, such as tendons, ligaments, trachea, muscles, skin, and the connective tissues of the eyes. Electrospinning, which was first studied in the 1930s, remains one of the most effective approaches for generating nanofibers [43]. One of the main issues for traditional electrospinning lies in the difficulty to control the morphology and geometry of the electrospun meshes, resulting from the rapidly whipping electrospinning jet [44, 45]. Despite the fact that attempts to control the fiber deposition in an accurate manner using melting electrospinning writing (MEW) have been reported, fibers produced by melt electrospinning usually have larger dimensions compared to conventional solution electrospinning. Another notable drawback of melt electrospinning is the use of elevated temperature, which may cause the degradation of thermo-sensitive polymers. Furthermore, many natural polymers including collagen cannot work with melt electrospinning [46]. In addition to MEW technique, near-field electrospinning (NFESP) have been developed as a process capable of depositing nanofibers in a controllable and continuous manner. The NFESP is characterized by using a shorter spinneret-to-collector distance together with very low voltages. However, this technique has difficulty in fabricating large scale scaffolds as it relies on a solution-dipped probe for jet feeding [47]. To solve the above mentioned problems, one possible solution is to develop a new technique which allows solution electrospinning to deposit fibers in a controlled manner. Recently, solution electrospinning have been reported to fabricate patterned nanofibrous mats [45, 48]. However, in these cases, the used apparatus was conventional electrospinning in combination of extra components including a side-wall, scanning functions and sharp-pin electrode, which were required to improve the focusing [45, 48]. In contrast to the setup they used, the solution electrospinning apparatus utilized in chapter 6 is much simpler. It was only composed of a conventional electrospinning system able to move in an x-y-z programmable manner. Interestingly, three different regimes could be obtained: the fiber deposition into a bundle of random fibers, a bundle with aligned fibers at its core, and single fiber deposition. The transition between these regimes was dependent on the applied working distance and the resulting electrical field strength. The transition between regimes of fiber deposition into a bundle of random fibers and a single unit has been demonstrated in previous studies [45, 49]. However, to the best of our knowledge, the deposition of fibers into a bundle with aligned at core through a direct writing electrospinning process have not been reported yet.

The ECM in the human body is complex and tissue specific with an often hierarchically organized structure [4, 46]. For example, articular cartilage is a highly organized tissue with
different functional zones [50-52]. Articular cartilage has limited capacity to self-repair after being damaged [53-55]. For this reason, the development of solutions that aid in the regeneration process of damaged articular cartilage remains an important issue in the regenerative medicine field. In this respect, we attempted to design a new fictional scaffold for cartilage repair. With the new technique, we fabricated lattice patterned mats that mimic the zonal organization of collagen fibers in articular cartilage. Four lattice patterned mats were further stack together to form a final functional construct. Compared to the typical electrospun mesh, the direct writing electrospinning (DW-ESP) scaffold was able to direct tissue and ECM fibril organization in a zone dependent manner. Additionally, some chondrogenic gene markers, including Sox9 and aggrecan (ACAN) were significantly up-regulated in this structure. The new technique provides a new approach for patterning structures mimicking the configuration of virtually any human tissues. However, some major issues relating to this technique still need to be addressed in the future: for example, to expand the amount of biomaterials that can be used with DW-ESP and to overcome the problem of charge repulsion to obtain thicker constructs in a more reliable manner.

7.4. Future Perspective

Nanofibrous scaffolds should be designed more and more as bioactive systems rather than just passive cell carriers. There are some important factors, such as mechanical properties, degradation rate, and geometrical structure and texture that need to be addressed in the “design stage” for tailoring nanofibrous scaffolds. All of these factors play an important role in governing cellular responses and host integration. Future research on bioactive artificial nanofibrous scaffolds may focus on the following aspects: The first strategy is to dig deeper in the basic mechanisms of tissue formation in biological systems. In the human body, there are multiple types of tissues of which structures are complex and tissue specific. The mechanisms underlying the formation, repair and maintenance of human tissue are not well understood. Without this fundamental knowledge that is crucial to custom-design functional scaffolds, regenerative medicine would practice in blindness. Secondly, developing and improving new techniques for manufacturing nanofibrous scaffolds is always necessary. Currently, phase separation, electrospinning, and self-assembling are the most popular approaches for fabrication of nanofibrous scaffolds. However, all these methods do not fully meet the increasingly demand for designing smart and bioactive scaffolds having tailored functions. In future studies, we need to fully explore the current techniques of fabricating
nanofibrous scaffolds. For example, we should understand how to improve direct writing electrospinning, which would allow depositing fibers forming a thick cm-scaled 3D dimension construct. Another approach is to integrate electrospinning with other techniques, which open up much possibility to manufacture artificial functional scaffolds. Electrospinning combined with other surface modification methods, such as plasma, lithography, ultrasonic spray coating and chemical grafting, could be used to generate bioactive topographic cues for the desired scaffold’s function. For instance, through using electrospinning and nanoimprinting techniques, nanofibrous scaffolds could be decorated with nanotopographic patterns, such as ridges and grooves, to better match the nanostructure of ECM [56]. Nanofibrous scaffolds having a hierarchical structure could be achieved by integrating self-assembling or phase separation processes with 3D printing techniques. Nanofibrous hydrogels, which were self-assembled from amphiphilic peptides, were used as bioink for printing 3D scaffolds with a predesigned pattern [57]. Thirdly, the release of growth factor from electrospun scaffolds could be envisioned. Growth factors which can induce angiogenesis are often required to promote tissue formation [58]. However, direct injection of soluble growth factors into the defect site will compromise the biological effect of growth factors due to a rapidly diffusion away from the injection site. Thus, scaffolds are used as drug release systems to maintain sustained release of growth factors. An ideal engineered release system for growth factor should resemble their natural release system in the human body. Lastly, developing new materials is also essential for designing bioactive scaffolds. The function and properties of scaffold are mainly determined by its materials chemistry. It would be attractive to design new materials which could induce stem cell differentiation, via high throughput screening of chemistry. It would be interesting to develop new materials that would not evoke immune response and inflammation in vivo.

Although much joint effort by scientists from multiple disciplines is still needed for the development of nanofibrous scaffolds in different regenerative medicine applications, it can be foreseen that nanofibrous scaffolds could approach “off-the-shelf” surgically implantable constructs in the near future.
Reference


Valorisation

Scientific knowledge does not, by itself, result in widespread implementation and positive social impacts. The research from the laboratory must be translated into practical use. This process, which is about creating impact through transfer of scientific knowledge, is also called ‘valorisation of knowledge’ or ‘technology transfer’. Valorisation has been defined by Netherlands Proteomics Centre as “The process of value-creation out of knowledge, by making this knowledge suitable and available for economic or societal utilisation and to translate this into high-potential products, services, processes and industrial activity”[1]. The research in this dissertation is on fabrication of bioactive scaffolds for regenerative medicine. Some of the findings show high potential for valorisation, especially, the one in chapter 5 where a novel technique is developed to fabricate nanofibrous scaffolds with tailored patterns. In the following sections, we will focus on elaborating the possible valorisation options for chapter 5. First, I will start the discussion from a social and clinical relevance, and subsequently I will discuss about the novelty concept and potential application.

8.1. Social and clinical relevance

Tendon is a tough bundle of fibrous connective tissue that transfer forces between bone and muscle [2, 3]. Like low vascularized tissue of articular cartilages, tendons are characterized with low cell density, low oxygen and nutrient requirement, and poor healing capacity [2, 4]. The injury of tendon is common. For example, the yearly incidence for Achilles tendon is roughly 18 per 100,000 in the world and this number is thought to increase rapidly due to the growth of sports participation and ageing of the world population [5]. Given their poor self-regenerative capacity, surgical management is the only recourse for large tendon injury[4]. Unfortunately, current surgical reparative approaches which are often associated with auto- or allografts replacement, could not fully restore their functional and structural properties. Recently, tissue engineering has been established as a promising strategy for tendon regeneration. Scaffolds are considered as one of the most important principle elements of tissue engineering.
Currently, electrospinning has been extensively used to fabricate scaffolds for tendon repair due to its capacity to continuously generate nanofibers. However, the traditional electrospinning technique has difficulty in fabricating scaffolds with buckled patterns, which are of important topographical cues in the development of connective tissue, including tendon. Thus, we invented a new technique to overcome this issue.

8.2. Novelty of the concept

Chapter 5 describes a new method of making wavy fibers, comprising: (a) depositing electrospun fibers on a surface of a thermally shrinkable material, which material is shrinkable in at least one direction, wherein the deposited fiber at least partially adheres to the shrinkable material; (b) shrinking the shrinkable material in at least one direction, thereby buckling the fiber. The technique presented here is simple and low cost compared to prior arts [6, 7]. Firstly, the deposited fibers will attach to the thermal shrinkable materials automatically without extra bonding process. Secondly, extra fixating agents or methods are not required for retaining the wavy pattern. To the best of our knowledge, this is the first time we present a technique that is capable to generate buckled patterns on individual fibers, shifting to buckling the whole fibrous mesh through configuring fiber depositing time. Another notable advantage of the technique described in chapter 5 is that wavy patterns could be regulated in a wide range of options, such as verifying the orientation of depositing fibers, using a second layer of a water-soluble polymer to change the amplitude and frequency of the wavy patterns, and varying the geometry shape and kind of shrinkable materials. The pilot cell biology study demonstrated that the obtained wavy scaffolds promoted cellular penetration, which is one of the major concerns in terms of electrospun scaffolds for tissue engineering applications. In addition, pilot studies indicated that the obtained wavy scaffolds promoted TGF-β signaling expression that is a key signaling pathway involved in the development of several connective tissues including tendon.

8.3. Potential application

The results described in this dissertation may open doors to the development of fabricating functional scaffolds for tissue regeneration, especially where wavy topographical cues were needed. The successful translation of this technique described here toward society, firstly and most importantly, may benefit a great variety of patients who need to recover from tendon injury. The following approaches could be adopted to put this novel technique in the market. Firstly, it is worth filing a patent application for this
technique; secondly, a startup company could be formed to aim at producing scaffolds for tissue engineering. However, before producing the final products, further studies on growing more specific cells (such as tendon cell lines) on the obtained wavy scaffolds should be performed, followed by validation in a functional animal model. Thirdly, this technique could be sold to the market directly, for example, to those companies that are already active with electrospinning products.
Reference


In this thesis, a number of novel strategies to design and fabricate instructive scaffolds for the regeneration of skeletal tissues are presented. Chapter 1 describes recent advanced in technologies for manufacturing nanofibrous scaffolds with a structure resembling the native ECM. Strategies for improving cell-scaffold interaction were also illustrated before discussing nanofibrous scaffolds for tissue engineering and regenerative medicine applications. From the review of literature proposed in chapter 1, it appears that traditional nanofibrous scaffolds do not display instructive properties, despite their physical resemblance to the native ECM. Thus, there is a great need for fabricating bioactive fibrillar scaffolds, which are capable to instruct cellular functions like adhesion, proliferation, and differentiation. Such bioactive scaffolds could be constructed via tailoring their physical properties (e.g. surface topography, mechanical properties, and structural features), and chemistry.

Chapter 2 shows functional electrospun fibers were fabricated from a new class of poly(butylene succinate)-based (PBS-based) polyesters containing either butylene thiodiglycolate (BTDG) or butylene diglycolate (BDG) sequences. Controllable properties including mechanical strength, wettability, and hydrolysis, could be achieved for these scaffolds through manipulating the molecular architecture and introducing an etheroatom along the polyesters backbone. Polyesters containing BDG sequences were more hydrophilic compared to those containing BTDG units. The higher hydrophilicity of polyesters led to a higher degradation rate. The polymer chemistry of the scaffolds also showed to have an important effect on cell-scaffold interactions. Here, the pure PBS scaffolds were beneficial to stimulate early hMSCs osteogenic differentiation. With the introduction of ether units (BDG) along its polymer backbone, the scaffolds showed enhanced mineralization properties. Having the presence of BTDG functional sequences in the polymer chains, scaffolds supported hMSCs chondrogenesis, while inhibiting their osteogenic differentiation. Tailoring the chemistry composition of nanofibrous scaffold could, therefore, serve as an appealing strategy for designing bioactive scaffolds in the regenerative medicine field.

Apart from the chemistry point of view, the structure and texture are also essential for functional scaffolds. In chapter 3, yttrium stabilized zirconia (YSZ) ceramic nanofibrous
scaffolds were fabricated via electrospinning a mixture of its precursors and polyvinylpyrrolidone (PVP), followed by annealing of the obtained green fibers. These ceramic nanofibrous scaffolds behaved like a paper, having high flexibility and good fatigue behavior at macroscale, which is not common seen in traditional ceramic mats. Yet, the scaffolds retained their stiffness at the microscale. The surface roughness and grain size of YSZ nanofibers were dependent on the annealing processing parameter. Cell biology results demonstrated that the osteogenic differentiation and mineralization of seeded hMSCs was supported by the nanofibrous structure of YSZ scaffolds, in contrast to the known bioinert behavior of bulk YSZ. In addition, a better mineralization of hMSCs was observed on the scaffolds with a higher surface roughness. Altogether, these flexible ceramic scaffolds presented highly potential application for bone regeneration.

As a logical consequence of the findings of chapter 3 related to the effect of surface topography on mineralization, the influence of surface roughness on cell behaviors were further investigated. Whereas the influence of surface roughness and specific surface topographies on hMSCs activity is known in a number of two-dimensional substrates, limited knowledge is present when translating these studies to 3D scaffolds. In chapter 4, the surface nanoroughness of individual fibers in PEOT/PBT electrospun scaffolds was varied via adjusting the relative humidity in the spinning chamber during electrospinning. The obtained scaffolds had the same polymer chemistry, similar porosity and pore size, and comparable fiber dimensions, yet a different surface roughness on individual fibers. The scaffolds having a rougher surface roughness of 71 nm had a higher protein adsorption than those having a surface roughness of 14 nm. To investigate the influence of the surface roughness of individual fibers on cellular activities, hMSCs were seeded in scaffolds and cultured in osteogenic and chondrogenic differentiation media for 21 days. The higher surface roughness (Ra=71.0 nm) promoted the expression of some osteogenic genes including osteopontin, bone morphogenetic protein 2, and runt-related transcription factor 2, while the lower surface roughness (Ra=14.3 nm) showed to better up-regulate other osteogenic genes, such as bone sialoprotein, collagen type I alpha 1, and osteocalcin. The lower surface roughness (Ra=14.3 nm) was more beneficial for chondrogenic genes expression at day 7. In summary, this research proposes a promising approach to fabricate electrospun scaffold with tailored topographical cues for skeletal differentiation of hMSCs.

The fibrous structure of native tissue has a complex and tissue dependent organization. For example, wavy patterns are observed in many connective tissues, while the collagen
fiber orientation in cartilage, tendons, or ligaments is anisotropic. Thus, traditional electrospinning, which normally produce non-woven meshes, could not fairly meet all the biomimicry requirement for fabricating scaffolds in regenerative medicine. As such, new electrospinning techniques were described in chapter 5 and chapter 6. Chapter 5 offers a simple and effective technique to fabricate wavy or crimped fibrous scaffolds. A wide range of crimped or wavy patterns could be achieved via tuning the processing conditions, including the orientation of the deposited fibers, additional sacrificial polymer layers, and the kind of thermally-shrinkable films. To the best of our knowledge, this research for the first time demonstrated the capability to generate multiscale crimped patterns spanning from fibers to fiber meshes through controlling the thickness of fiber layers. Cellular infiltration is one of the main concerns for electrospun scaffolds regarding their application in regenerative medicine. Here, hMSCs showed a more even cellular penetration in wavy electrospun fiber meshes compared to the non-wavy fiber meshes. Given the obtained wavy patterns mimicking the configuration observed in connective tissues, a mink lung epithelial cell line harboring a TGF-β reporter construct was used to examine the effect of such topographical cues on TGF-β signaling expression, which is a key signaling pathway involved in the development of several connective tissues. Compared to the non-wavy scaffolds, the wavy scaffolds promoted the expression of TGF-β signaling.

Although solution electrospinning technique has been widely used for varied application, its inherent working principle of exploiting jet instability causes also the difficulty to deposit fibers in an accurate manner. This may hinder the further application of electrospinning where more hierarchical organization of nanofibers is needed. As such, chapter 6 shows a new technique for solution electrospinning with the capability to make electrospinning jet follow a stable path from the tip of the spinneret needle to the collector plate. Complex patterns and 3D multiscale structures with controllable porosity could be easily produced using this technique in contrast to traditional electrospinning. Using this new technique, a scaffold that mimics articular cartilage with its distinct multi-zone morphology was fabricated. To estimate their potential application in cartilage tissue engineering, hMSCs was seeded in scaffolds in chondrogenic differentiation medium for 21 days. SEM analysis, histological and immunostaining results demonstrated that the scaffold directed tissue organization and fibril matrix orientation in a zone-dependent way. Moreover, gene expression results revealed that chondrogenic markers including Sox9 and aggrecan were significant upregulated in the scaffolds compared to traditional
electropsun mesh. Overall, the research here provided a simple way to fabricate customized fibrous scaffolds for regenerative medicine, especially those for which anisotropy is of importance.

In chapters 7, themes are summarized and discussed in relation to the results presented in this thesis and a few possible ideas for future work. Finally, chapter 8 describe the valorization potential of the research presented in this thesis.
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Honglin Chen
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Honglin Chen was born and raised in a beautiful village of Putian, China. In the fall of 2004, he began his studies at Minjiang University where he received training in in the principle of Chemical and Chemistry Engineering. In 2008, he continued his studies in Department of Chemistry at East China Normal University, Shanghai. After three years, he graduated with a Masters of Polymer Chemistry and Physics. His dissertation for Master degree is entitled “electrospun chitosan-polycaprolactone materials and their application for tissue engineering”. In the fall of 2011, he started his doctoral studies in the Tissue Regeneration group of Clemens van Blitterswijk, University of Twente, Netherlands, under the daily supervision of Prof. Lorenzo Moroni and Dr. Roman Truckenmüller. He gained his knowledge in designing scaffolds for guidance skeletal differentiation of human mesenchymal stromal cells, as well as his age. In the December of 2014, the Tissue Regeneration Group found its bright future in Maastricht University. He decided to move from Enschede to Maastricht following his supervisors and continued to pursue his PhD degree in the Complex Tissue Regeneration.
This thesis is based on the following publications:

**Peer Reviewed Papers:**

Honglin Chen, Matteo Gigli, Chiara Gualandi, Roman Truckenmüller, Clemens van Blitterswijk, Nadia Lotti, Andrea Munari, Maria Letizia Focarete, Lorenzo Moroni. Tailoring chemical and physical properties of fibrous scaffolds from block copolyesters containing ether and thio-ether linkages for skeletal differentiation of human mesenchymal stromal cells. *Biomaterials.* 2016; 76: 261-272.


Honglin Chen, Xiaobin Huang, Minmin Zhang, Febriyani Damanik, Mattew B. Baker, Anne Leferink, Huipin Yuan, Roman Truckenmuller, Clemens van Blitterswijk, Lorenzo Moroni. Tailoring surface nanoroughness of electrospun scaffolds for skeletal tissue engineering. (In submission).

Honglin Chen, Danielle Ferreira Batista, Roman Truckenmuller, Clemens van Blitterswijk, Lorenzo Moroni. Tailoring Crimp Pattern of Electrospun Fibers by using thermal shrinkage film. (manuscript in preparation).

Honglin Chen, Afonso Malheiro, Paul Wieringa, Carlos Mota, Roman Truckenmuller, Clemens van Blitterswijk, Lorenzo Moroni. Fabrication of scaffold mimicking the architecture of articular cartilage using direct-write electrospinning. (In submission).

**Book Chapter:**

**Other Publications:**


**Conference Abstracts:**

Honglin Chen, Xiaobin Huang, Minmin Zhang, Febriyani Damanik, Roman Truckenmuller, Clemens van Blitterswijk, Lorenzo Moroni, Tailoring surface nanoroughness of electrospun scaffold for skeletal tissue engineering. Netherlands society for biomaterials and tissue engineering (NBTE) conference(Oral, Lunteren, Netherlands, 2015)

Honglin Chen, Danielle Ferreira Batista, Roman Truckenmuller, Clemens van Blitterswijk, Lorenzo Moroni. Tailoring Crimp Pattern of Electrospun Fibers by using thermal shrinkage film. European Society for Biomaterials (ESB) Conference(Oral, Liverpool, UK 2014)

Honglin Chen, Chiara Gualandi, Matteo Gigli, Marialetizia L. Focarete, Nadia Lotti, Roman Truckenmuller, Clemens V. Blitterswijk, Lorenzo Moroni. Bioactive Elastomers for
Osteogenic Differentiation of Human Mesenchymal Stem Cells. Termis-EU Conference (Poster, Genova, Italy 2014).