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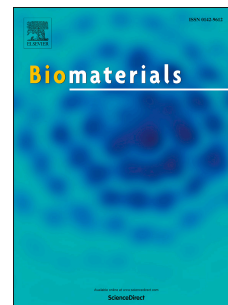
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Sulfated polysaccharide-based scaffolds for orthopaedic tissue engineering

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Abstract

Given their native-like biological properties, high growth factor retention capacity and porous nature, sulfated-polysaccharide-based scaffolds hold great promise for a number of tissue engineering applications. Specifically, as they mimic important properties of tissues such as bone and cartilage they are ideal for orthopaedic tissue engineering. Their biomimicry properties encompass important cell-binding motifs, native-like mechanical properties, designated sites for bone mineralization and strong growth factor binding and signalling capacity. Even so, scientists in the field have just recently begun to utilise them as building blocks for tissue engineering scaffolds. Most of these efforts have so far been directed towards *in vitro* studies, and for these reasons the clinical gap is still substantial. With this review paper, we have tried to highlight some of the important chemical, physical and biological features of sulfated-polysaccharides in relation to their chondrogenic and osteogenic inducing capacity. Additionally, their usage in various *in vivo* model systems is discussed. The clinical studies reviewed herein paint a promising picture heralding a brave new world for orthopaedic tissue engineering.

1. Introduction

Orthopaedic diseases are the second largest contributor to disability worldwide and are expected to grow rapidly in the foreseeable future due to the aging population.[1] They include debilitating diseases such as osteoarthritis, tendinopathies, osteoporosis, as well as skeletal and joint fractures.[2, 3] The current approaches for addressing this grand challenge rely on various prosthetic, allograft and autograft-based strategies. Even though the prosthetic-based interventions have shown exciting results in recent years, they still face major shortcomings such as suboptimal long-term outcomes, the need for revision surgeries and risk of infection.[4] Allograft and autograft strategies on the other hand impose their own limitations including the possibility of disease transmission, insufficient autologous resources, rejection of allograft tissue and potential need for immunosuppression therapies.[5] To overcome these hurdles a great variety of tissue engineering approaches have been proposed over the years (Figure 1).[3, 6]

The grand goal of tissue engineering is to generate artificial tissues with the capacity to bring normality back to dysfunctional tissues by replacing them with more functional ones.[4] The tissue engineering paradigm involves scaffolds combined with potent cell sources and suitable biochemical signals [7], which together can promote the formation of

new organs and tissues.[8] Ideally, these scaffolds emulate key physical and molecular features of the native extra cellular matrix (ECM) in order to facilitate cell attachment, proliferation and differentiation and ultimately new tissue growth (Figure 1).[9] The key in this regard is to provide the cells with a native-like milieu with the capacity to guide them into tissue specific phenotypes.[10-14] Generally speaking, bioactivity is included into scaffolds by using: I) insoluble signals, such as bio-ceramics and carbon-based nanocues [15], II) introducing growth factors and other biological moieties into the scaffold matrix [16], or III) by incorporating cell adhesion and differentiation promoting oligopeptides (such as the cell binding RGD peptide [17, 18]).

While all of these methods have shown promise in the synthesis of bioactive scaffolds, they still face certain limitations in the clinic. For instance, i) some insoluble signals such as carbon-based nanomaterials can cause a foreign body response that can facilitate tissue fibrosis [19, 20], ii) growth factors often face issues such as loss of bioactivity, low tissue penetration and dosage-dependent toxicity [21] and iii) many of the bioactive oligopeptides do not facilitate the needed intracellular signalling pathways for optimum tissue generation; even though a number of proteins (such as fibronectin[22, 23], collagen[24], osteopontin,[25] vitronectin[26] and fibrinogen[27]) stimulate much more robust intracellular signalling than bioactive oligopeptides[28-31] they are limited by either foreign body responses from the host or in some cases high cost and low scalability. For these reasons, native-like and abundant biopolymers with inherent bioactivity have attracted much attention in biomaterials science. In particular, sulfated polysaccharides are by now widely recognized for their ability to bind to important cell receptors to facilitate cell adhesion, proliferation and differentiation.[32, 33] They can also bind to and signal a number of important growth factors such as fibroblast, vascular endothelial and bone morphogenetic protein growth factors for controlled growth factor release; and they can improve growth factor bioavailability by protecting them against proteinase degradation.[32, 34-37]

In simple terms, sulfated polysaccharides can be classified under three distinct categories including i) sulfated GAGs, ii) marine sulfated glycans and iii) chemically sulfated polysaccharides. While the first two categories are inherently sulfated polysaccharides, the third one consists of non-sulfated polysaccharides that are chemically modified with various sulfating agents. Regardless, the bioactivity of sulfated polysaccharides depends on factors such as degree of sulfation and sulfation pattern.[35, 38] For instance, hyaluronic acid (HA)/collagen type I matrices were shown to inhibit osteoclast differentiation and resorption,

largely dependent on degree of sulfation of HA.[39] To this end, highly sulfated HA was capable of improving bone regeneration in *in vitro* and *in vivo* models.[40-42] In other studies, an intimate link between sulfation pattern and chondrogenesis has been proposed.[43] For example, it was shown that chondroitin sulfate (CS) rich in 4,6-O-disulfated disaccharides, had a higher potential to upregulate the expression of important chondrogenic biomarkers when compared to other CS derivatives containing either 4- or 6-O-sulfated disaccharides.[43]

Accordingly, sulfated polysaccharides have been rapidly picked up by scientists in the field in order to manufacture more bioactive scaffolds that can facilitate better skeletal tissue regeneration.[44-54] These scaffolds were made via various fabrication methods - such as casting, electrospinning and 3D printing - from either individually sulfated polysaccharides or in combination with other biopolymers. Generally speaking, the scaffolds have been used in two different ways to assist osteogenesis or chondrogenesis: i) in combination with growth factors to facilitate differentiation of cells via controlled release of growth factors, or ii) in the absence of any growth factors by solely relying on intermolecular interactions with important cell-membrance receptors.[55, 56]

This paper, reviews the most recent progress in sulfated polysaccharide-based scaffolds for skeletal tissue engineering, with particular focus on bone and cartilage tissue engineering. Specifically, three different groups of sulfated polysaccharides - sulfated GAGs, marine sulfated glycans and chemically sulfated polysaccharides, and their usage as building blocks in orthopaedic scaffolds are reviewed; since these polysaccharides present the most promising avenues in this field. This review also highlights the ability of these scaffolds to direct progenitor cells into either chondrogenic or osteogenic differentiation. Finally, application of these scaffolds in various preclinical studies related to mending bone and cartilage defects along with more complex osteochondral lesions are reviewed, as such studies are of utmost importance for bridging the current gap between the laboratory and the clinic.

2. Naturally Sulfated Polysaccharides

Sulfated polysaccharides can be derived from the ECM of animal tissues in the form of sulfated GAGs or from plants such as marine algae in the form of alginate, carrageenan, fucoidan and ulvan (Figure 2). The sulfate groups in the abovementioned biopolymers can also be chemically conjugated to the sugar backbones of non-sulfated molecules such as HA,

chitosan, alginate and cellulose. Along these lines, this section is divided into three subsections dealing with sulfated GAGs and polysaccharides derived from natural sources as well as sulfated polysaccharides that are custom-made in the laboratory. Notably, the wide variety of sulfated polysaccharides reviewed can display differing bioactivity depending on the sulfate position and degree.

2.1 Glycosaminoglycans (GAGs)

Sulfated GAGs are present in the ECM, cellular membrane and intracellularly within eukaryotes (Figure 2). They therefore, play an essential role in modulating extracellular and intracellular interactions. In simple terms, GAGs can be defined as negatively charged heteropolysaccharides, whose disaccharide units are comprised of repeating disaccharide units of a uronic acid (iduronic or glucuronic acid) and an amino sugar (glucosamine or galactosamine). Based on their disaccharide composition, they are grouped into four different families including heparin/heparan sulfate, chondroitin/dermatan sulfate, keratan sulfate and HA. While heparin, heparan, chondroitin, dermatan and keratan sulfate are sulfated and post-translationally synthesised via attachment to a core protein, HA is non-sulfated and synthesised at the cell surface without a protein core. Importantly, GAGs can differ significantly from one another in terms of bioactivity and structural complexity depending on their specific biosynthesis pathway and source of derivation.[57]

Heparin and Heparan Sulfate

Heparin is a highly sulfated GAG only produced by connective tissue mast cells and exclusively decorates the protein core of serglycin. [58] In contrast, heparan sulfates (HS) decorate intracellular, ECM and cell surface proteoglycans and are produced by almost all cell types where they play important roles in a wide range of physiological processes including cell proliferation and differentiation, immune responses, as well as angiogenesis.[59-62] Both heparin and HS are composed of repeating disaccharide units of either iduronic or glucuronic acid and glucosamine units but with less iduronic acid and less overall sulfation in HS compared to heparin. Importantly HS does not contain sulfation at the C3 position and does not possess anti-coagulant activity.[63-65] Both heparin and HS interact with a large number of proteins, (including heparin-binding growth factors), which together with their cell signaling role, make them ideal choices for scaffolding materials.[61]

Heparin has been widely explored in tissue engineering owing to its ease of supply being used clinically as an anticoagulant and is often used as an analogue of HS.[66-68] Heparin

and HS bind to a range of proteins via electrostatic interactions that are controlled by its three-dimensional structure, anionic nature and sulfation patterns. Heparin is known to enhance the osteogenic potential and bioavailability of bone morphogenetic protein-2 (BMP-2) through its binding, stabilization and presentation to cells.[69-71] Indeed, in a study by Hettiaratchi, Miller [72] it was shown that methacrylated heparin microparticles could bind high quantities of BMP-2, vascular endothelial growth factor (VEGF) and fibroblast growth factor-2 (FGF-2), which in turn could stimulate alkaline phosphatase (ALP) activity in skeletal myoblasts (C2C12) and increase the cell division rate. Notably, such heparin microparticles typically demonstrate better presentation of growth factors when compared to gelatin microparticles and soluble heparin; something which has been speculated to arise from heparin's higher charge density.[72] Similarly, PLGA microspheres when functionalised with both heparin and BMP-2, could significantly up regulate MG-63 osteosarcoma cell differentiation as seen through the enhanced expression of osteocalcin (OCN) and osteopontin (OPN), whilst simultaneously increasing both ALP activity and deposition of important bone minerals.[73]

However, heparin's anticoagulant capacity can hinder bone regeneration through antithrombin III activation, which can prevent the accumulation of various tissue regenerative growth factors and cytokines in the defected bone region. Thus, the lesser charged negatively charged HS could be a more useful bioactive supplement. To this end, Bramono, Murali [74] compared the osteogenic potential of heparin and HS from various sources; as regulators of BMP-2 activity, and found that heparin could up regulate BMP-2 induced osteogenic differentiation of C2C12 cells in the short term, however they did not observe any significant BMP-2 stimulated bone matrix mineralisation after 14 days. Interestingly, HS delivered BMP-2 in a prolonged and sustained manner, at more physiologically relevant concentrations whilst retaining its osteogenic activity (when compared to heparin). This was thought to be associated with the higher growth factor binding and signaling capacity of HS compared to heparin which enables the more efficient presentation and signaling of osteogenic ligands to their cell associated signaling receptors.[75] HS has also been shown to regulate other growth factors in the transforming growth factor beta (TGF- β) superfamily. For instance, Chen, Wang [76] showed that, in the presence of TGF- β 3, HS was capable of inducing chondrogenic differentiation of human MSCs whilst activating important TGF- β related signaling pathways. Similarly, heparin in combination with a self-assembling peptide (RAD 16-I) could drive adipose-derived stem cells (ADSCs) into the chondrogenic lineage as

evidenced by collagen type II up regulation; a phenomenon that was speculated to arise from heparin's affinity towards VEGF.[77] More recently, a biphasic silk fibroin biomaterial incorporating heparin was reported to increase growth factor retention and thereby preventing the undesired initial burst-like release that is so common in many traditional scaffolds.[78] Interestingly, the incorporation and controlled release of TGF- β 2 and GDF5 (growth differentiation factor 5) into the scaffold up-regulated chondrogenic markers, including SOX9, aggrecan and collagen type III (Figure 3).

In summary, several studies have demonstrated the versatility of heparin and HS to efficiently deliver and preserve the function of important chondrogenic and osteogenic growth factors. As mentioned, the prominent anticoagulant capacity of heparin can diminish the accumulation of growth factors and cytokines in a bone defect site and subsequently hinder tissue regeneration. HS, the less sulfated heparin analogue, on the other hand holds promise as an alternate delivery vehicle without such undesirable side effects. In this regard, HS has already showed promise at permitting sequestration and controlled local delivery of growth factors resulting in an improved bone and cartilage matrix production Overall, HS and heparin-based biomaterials will in the authors opinion soon move beyond their current usage in anti-coagulant treatments, and towards growth factor and cytokine delivery vehicle for bone and cartilage tissue regeneration.

Chondroitin Sulfate

Chondroitin sulfate (CS) is the most abundant GAG found in vertebrate and invertebrate ECM and decorates intracellular, ECM and cell surface proteoglycans. It is a linear polysaccharide composed of repeating disaccharide units of glucuronic acid and galactosamine that can be sulfated at carbon's 2 on the glucuronic acid, and 4 and/or 6 on the galactosamine, which provide heterogeneity in structure.[79] Aggrecan is the major CS proteoglycan in cartilage that binds to HA to form aggregate structures that have a high water retention capacity and provide the hydrodynamic weight bearing properties of cartilage.[80] CS has been shown to stimulate the synthesis of aggrecan, HA, glucosamine and collagen II, as well as preventing chondrocyte apoptosis and degradation of cartilage by inhibiting ECM degrading enzymes. Accordingly, CS has been extensively explored for cartilage repair and chondrogenic differentiation of stem cells.[81] For a more in-depth analysis of the influence of CS hydrogels on stem cell fate the reader is referred to an excellent review given recently by Farrugia, Lord [82]

A number of recent studies have harnessed the abovementioned biomimicry properties of CS in cartilage tissue engineering with exciting outcomes. For instance, a study by Levett, Melchels [83] aimed to enhance chondrocyte behaviour in gelatin methacrylate-based (GelMA) hydrogels by incorporating GAGs including hyaluronic acid methacrylate (HAMA) and CS methacrylate (CSMA) into the hydrogels; both separately and together. Interestingly, they found that the integration of HAMA enhanced chondrocyte re-differentiation and improved matrix distribution, whereas CSMA showed marginal improvements over both the GelMA control and GelMA/HAMA/CSMA triple composite. This means that HAMA positively influences bioactivity and the mechano-physiological properties of GelMA hydrogels when compared with CSMA. Although, HA provides the biochemical cues for chondrogenesis, it was shown that the inclusion of CS in the HA hydrogels can upregulate mRNA expression of chondrogenic markers, while decreasing expression of the hypertrophic markers that are normally associated with HA hydrogels.[85] Additionally, incorporation of CS into HA hydrogels led to an increase in GAGs accumulation both *in vitro* and *in vivo*. Similar results were observed by Costantini, Idaszek [86] during bioprinting of bone marrow derived hMSCs in a composite matrix containing GelMA, HAMA and CS amino ethyl methacrylate (CSAEMA). In the absence of HAMA, the ratio of collagen II/collagen I and collagen II/collagen X increased suggesting neocartilage formation, whereas differentiation towards hypertrophic cartilage was observed with HAMA alone. This may be due to the stiffness increase from 59 kPa (GelMA/CS) to 100 kPa (GelMA/CS/HAMA), as MSC differentiation is sensitive to interface stiffness.[87, 88] In summary, they concluded that the chemical composition, network density and stiffness of the 3D microenvironment in combination play a role in determining the chondrogenic potential of MSCs, with CS showing the most promising cartilage regenerative capacity.

CS has also been employed together with other biopolymers such as polyethylene glycol (PEG), chitosan, and alginate to constitute bioactive scaffolds for cartilage tissue engineering.[47, 89-94] In a noteworthy example, with the aim of evaluating the effect of CS sulfation degree on its interaction with positively charged growth factors, researchers made two different types of scaffolds composed of poly(ethylene glycol)-diacrylate (PEG-DA) with either CS or desulfated CS.[90] *In vitro* experiments demonstrated that the release of a positively charged model protein (histone) from hydrogels containing desulfated CS resulted in an increased histone release when compared to a hydrogel containing normal CS, suggesting that sulfation alone plays an essential role in modulating protein interactions with

GAG hydrogels, and thereby also the growth factor release profile. Interestingly, MSCs in hydrogels containing desulfated CS had significantly higher expression of collagen II and aggrecan by day 21 in chondrogenic medium, compared to PEG control scaffolds or CS containing scaffolds. This was speculated to arise from the augmented TGF- β 1 pull-down from culture media caused by the presence of CS in the hydrogels.

In another study, a biomaterial composed of chitosan and CS was used for cartilage tissue engineering.[47] The *in vitro* results with a pre-chondrocyte cell line (ATDC5) showed that chitosan/CS induced a higher collagen II/collagen I ratio (a characteristic of hyaline cartilage formation) after 21 days, when compared to pristine chitosan. Furthermore, the collagen X expression in chitosan/CS showed an increase after 21 days compared to pristine chitosan scaffolds, indicating that these scaffolds can drive ATDC5 cells into a hypertrophic state. CS has also been used in combination with alginate to form porous scaffolds for chondrogenesis of hMSCs.[94] After 14 days, it was shown that under chondrogenic conditions total collagen and GAG contents were higher in cells seeded onto CS-containing scaffolds as compared to the CS-free ones.

Apart from cartilage tissue engineering, CS has been used to promote osteoblast adhesion for bone tissue engineering.[95] In this respect, Vandrovcová, Douglas [96] coated PLGA with collagen I with and without CS. Results, indicated that CS improved both the osteoconductivity and osteoinductivity of the (osteoblastic) MG-63 cell line, observed through the increased proliferation and upregulation of osteocalcin, as compared to pristine collagen I coatings. Similarly, titanium implants have also been coated with CS/collagen[97] or CS,[98] as sulfated GAGs are known to bind calcium and calcium phosphates such as hydroxyapatite [99]. The former compared three forms of CS (4-sulfated CS (CS A); 6-sulfated CS (CS C) and dermatan sulfate (CS B)), and found that both CS A and CS B stimulated local osteoblast adhesion. We also note, that the study by Dudeck, Rehberg [98] demonstrated a synergistic effect between CS and hormone replacement therapy in an osteoporotic rat model, and thus indicates that CS scaffolds could open new therapies for osteoporosis.

In summary, CS has been used in conjunction with biopolymers to form more functional composite biomaterials that can facilitate both chondrogenesis and osteogenesis. When used with cartilage forming cells, it has been seen that the inclusion of CS increases the expression of collagen II, while facilitating a more hyaline-like cartilage formation, as a result of enhanced binding with growth factors and integrin-mediated cell-matrix interactions. the CS

structure, and specifically the location of the sulfates on the CS backbone, directly influences its ability to bind to cells and direct their differentiation. Therefore, CS holds great promise for skeletal tissue engineering since it can both have an impact on chondrogenesis and bind to important components of the hard phase of bone; all because of its many sulfate groups.

2.2 Marine sulfated Glycans

Over 70% of the earth's surface is inundated by oceanic environments, rich in biodiversity. Among these marine organisms lies algae and seaweed that are abundant with bioactive compounds of use in the field of biomedicine owing to their numerous health benefits stemming from their anti-inflammatory, anti-cancer, anticoagulant and immunomodulatory properties.[89, 100, 101] Although seasonal disparities can influence their overall composition,[102] their sustainable cultivation is not constrained by climate as with various terrestrial plant species. Notably, some of these algae are also made up of simple sugars (monosaccharides) joined by glycosidic bonds (Figure 2) that resemble GAGs and they can promote protein binding and cell growth without giving rise to immunogenicity. As with other GAG-like polymers, the bioactivity of sulfated marine sugars depends on their composition, molecular weight, degree and location of sulfate groups. The three most prevalent marine-based sulfated polysaccharides currently used in biomedicine are carrageenan, fucoidan and ulvan, derived from red, brown and green algae respectively.

Carrageenan

In simple terms, Carrageenan's (CARs) can be described as linear and water-soluble anionic-sulfated polysaccharides. They are derived from red algae of the class *Rhodophyceae* and identified based on their disaccharide sulfation. They have previously successfully been exploited in bone and cartilage tissue engineering applications, due to their thermoreversible gelling behaviour in the presence of non-toxic cations, as well as their ability to facilitate bone apatite formation.[103-111]. As a noteworthy example, Popa, Caridade [103] demonstrated that kappa (κ) - CAR hydrogels were able to support the proliferation and chondrogenic differentiation of encapsulated ADSCs. Following 21 days in culture they also observed an increase in hydrogel storage modulus and viscoelastic properties possibly related to the ECM deposition from the cells. Additionally, the mechanical properties of the hydrogel, following compression were observed to be in the range of native human cartilage. In another study, Oliveira, Silva [112] investigated how variations in the primary structure of CARs can influence bone mineralisation. They compared the osteogenic properties of three

different CAR sugar backbones, kappa (κ), iota (ι), and lambda (λ), within a chitosan/polycaprolactone (PCL)-based scaffold. In this respect, it was demonstrated that bone apatite formation varies significantly between different CAR species. Specifically, of the three CARs employed, the ι -variant demonstrated significantly higher biomineralization, possibly due to an increased affinity for various bioactive compounds from the osteogenic media as a result of higher sulfur, oxygen and nitrogen content within its sugar-like backbone. In a similar vein, the osteogenic capacity of a composite containing ι -CAR/chitosan/gelatin was recently explored.[113] Here, the researchers found that the inclusion of gelatin with its native RGD peptides and chitosan with its favorable cationic and osteogenic properties,[114] into the CAR hydrogel network, promoted the osteogenic differentiation of ADSCs. Notably, they found that the inclusion of a 10 wt % ι -CARs significantly increased the alkaline phosphatase activity of encapsulated cells when compared to the composites containing 0, 5 and 15 wt % of ι -CAR. Correspondingly, an osteogenic-specific histology assay suggested that the 5 and 10 wt % ι -CAR-based composites caused higher mineral deposits following a 28-day *in vitro* study than the other groups. In another recent investigation, κ -CAR was blended into biodegradable polyesters to consummate a biocompatible scaffold for bone tissue engineering.[51] Interestingly, the authors found that – like the other studies reviewed herein – the presence of κ -CAR could facilitate the formation of nanosized apatite crystals when compared to pure polyesters, which instead gave rise to non-native-like and larger microsized crystals. Of interest, the introduction of κ -CAR in the polyester material also enabled tailored degradability. In a related study, Liang, Wang [52] found that the expression of cartilage specific genes (SOX9, collagen II and aggrecan) were up regulated with increasing CARs concentrations within chitosan, when compared to pristine chitosan. They also showed that CARs promoted cellular responses such as adhesion, viability and proliferation in the composite hydrogel. These benefits were attributed to the chemical similarities between CARs and CS, which is widely recognized for its chondrogenic capacity.

The thermoreversible and thixotropic gelling behaviour of κ -CAR under physiological conditions also makes them suitable as injectable hydrogels for cartilage tissue engineering, as evidenced by a recent study by Rocha et al.[115] Specifically, in this study, it was found that ADSC-laden κ -CAR hydrogels cultured in TGF- β 1 supplemented growth media did not induce chondrogenic differentiation, though when used with chondrogenic medium, the cells developed a spherical, chondrogenic-like phenotype. Likewise, immunohistochemical

analysis revealed increased collagen II deposition following the integration of TGF- β 1 in the κ -CAR hydrogels under chondrogenic conditions, suggesting the production of cartilage-specific proteoglycans. Interestingly, the heated gelling conditions did not elicit thermal stress on encapsulated hASCs following live-dead staining, justifying their potential future use for *in situ* forming hydrogels for cartilage tissue engineering.

Fucoidan

Fucoidan is a sulfated polysaccharide derived from the cell-wall matrix of brown seaweed. It contains a substantial amount of L-fucose and sulfate ester groups which varies from species to species.[101] The species that is most frequently used in the field, is - *Fucus vesiculosus* - which typically gives rise to Fucoidan consisting of 1,2- α -fucose, with its sulfate groups primarily located at C4 position.[116] Interestingly, fucoidan has been shown to interact with transforming growth factor (TGF)- β 1, which was speculated to be associated with its heparin-like chemical structure,[117] and like the CARs, fucoidan can also facilitate bone-like apatite formation.[118] Specifically, it was demonstrated that the addition of fucoidan promoted osteocalcin and ALP production whilst supporting human bone marrow stromal cells (hBMSC) growth. The increase in ALP was indicative of initial osteogenic differentiation, which happened after a rapid cell division (a well-known stage in osteogenic differentiation of stromal cells in culture). Interestingly, they also found that fucoidan could more than double the compressive strength of the scaffolds from 191 ± 5 KPa to 414 ± 3 MPa, something that could come to use later, due to the intimate link between cartilage/bone formation and biomaterial stiffness.[119] In another study, Puvaneswary, Raghavendran [50] developed a porous fucoidan scaffold to influence bone mineralisation and apatite formation. These scaffolds promoted hBMSC attachment, proliferation and differentiation. Though the lengthy process of mineralisation was not significant, upregulation of collagen I under osteogenic conditions demonstrated osteogenesis within the fucoidan composite. Additionally, Runt-related transcription factor-2 (RUNX2) and osteonectin (ON) were significantly upregulated compared to the chitosan only hydrogel.

Owing to the TGF- β -binding properties of fucoidan, it was also exploited for cartilage tissue engineering applications. For instance, Karunanithi, Murali [120] studied the chondrogenesis of encapsulated hMSCs within a fucoidan-alginate composite. The results revealed that hMSCs cultured in chondrogenic medium supplemented with fucoidan expressed a higher level of chondrogenic markers (including tenascin-C, SOX9, collagen II, aggrecan and cartilage oligomeric matrix protein). In addition, the cultures expressed a

significantly lower level of hypertrophy markers (including Col X and Runx2), when compared to alginate hydrogels. Further more, cells encapsulated in the fucoidan-alginate hydrogel produced a higher GAG content at day 21 (when compared to alginate hydrogels), which is a widely recognized indicator of mature chondrocyte phenotype. Thus fucoidan may enhance the chondrogenic differentiation of stem cells due to its affinity to various growth factors, such as TGF- β 1. Likewise, cell condensation – a hallmark for chondrogenic differentiation - were observed in this study, which puts further emphasis on the promise that Fucoidan holds in cartilage tissue engineering.

Ulvan

Ulvan is a lightly branched anionic-sulfated polysaccharide, which is derived from the cell wall of green algae; and consist of sulfated rhamnose, iduronic and glucuronic acids.[121] The ulvan sugar share a chemical similarity with GAGs, due to its glucuronic acid and sulfate groups.[89, 122] As with the previously investigated marine glycans, ulvan has been used in combination with chitosan to produce osteogenic coatings for titanium implants. To this end, coatings seeded with 7F2 osteoblasts showed complete confluency after 6 days; something significantly different as compared to cells seeded on pure ulvan or pure chitosan. From this point-of-view ulvan/chitosan composite promoted the attachment and proliferation of 7F2 osteoblasts while maintaining the cell morphology and viability. In a related study by Dash, Samal [123] ulvan was used for bone tissue engineering applications. Purposely, the group introduced methacrylate groups to the ulvan backbone to further increase the physiological stability of the hydrogel through UV-crosslinking. Hydrogels were incubated with ALP at varying concentrations to gauge mineralisation capacity, as mineralisation is known to promote bioactivity through the formation of chemical bonds with surrounding bone tissue after implantation. The lowest methacrylated-ulvan group, saw the highest concentration of ALP resulting in pre-osteoblast cells differentiating towards an osteogenic lineage, as interpreted from increased ALP activity and a reduction in cell proliferation.

Overall, these naturally sulfated marine glycans have seen limited use thus far in orthopaedic tissue engineering applications. Since they're known to have chemical compositions that mimic several ECM-based GAGs and proteoglycans there's no doubt they could be used to drive the R&D engine of the next-generation of biomaterials for orthopaedic tissue engineering. Especially, their strong affinity towards a wide range of tissue regenerative growth factors makes them ideal growth factor delivery vehicles, which in turn further improve their tissue regeneration capacity. Additionally, their high abundance and

sustainability along with reduced immunogenicity strongly advocates their promise in the broader field of tissue engineering.

2.3 Chemically sulfated

The biological properties of sulfated polysaccharides from mammalian and plant-based sources are vast. In fact, their bioactivity is a function of molecular size, type of sugar-backbone and sulfate content[124] However, naturally-derived polysaccharides typically give rise to batch-to-batch variations, which further hinders the reproducibility of their ensuing biophysical properties.[125, 126] As a result, in an effort to produce sulfated polysaccharides with more specific and controllable functional properties, researchers have started to chemically manipulate non-sulfated polysaccharides such as HA, chitosan, alginates and cellulose. Controlled chemical sulfation of these polysaccharides can be achieved through various surface immobilisation strategies including chemical binding[127] and electrostatic assembly.[128] Modifying or combining these polysaccharides with sulfate groups could exploit their native chondrogenic or osteoblastic potential whilst prolonging growth factor delivery to promote proliferation and differentiation of tissue specific stem cells, as well as circumventing shortcomings such as hypertrophy or rapid enzymatic scaffold degradation.[129]

Hyaluronic acid (HA)

HA is a naturally occurring GAG, that has been widely utilised in tissue engineering as it possesses cell surface receptors such as CD44 that enable cell binding,[130] and is immunoneutral at the same time.[131] Indeed, the CD44-based cell binding receptor has been utilised and shown to increase chondrogenesis.[83, 132] Various groups have also studied the effect of modifying the HA with sulfate groups, to enable sustained growth factor delivery through improved growth factor binding. For instance, Xu, Jha [133] investigated the effect of decorating HA with heparin. It was seen that when MSCs were seeded onto a HA-heparin hydrogel with BMP-2 present, there was significant upregulation of mRNA and key chondrogenic genes including collagen II, SOX9 and aggrecan, as compared to pristine HA. These improvements can be attributed to the heparin subgroups that contain sulfate groups, which were seen to have a higher binding capacity for BMP-2. Importantly, a sustained release profile over 13 days was observed, compared to pristine HA which displayed an initial burst release profile.

In a similar vein, Jha, Yang [134] combined HA with perlecan, a sulfated HS proteoglycan. Here, the HA-perlecan hydrogel exhibited the ability to bind significantly more BMP-2 as compared to HA alone and promoted chondrogenesis. Likewise, Srinivasan, McCoy [135] combined HA with HS and demonstrated a targeted and controlled delivery of BMP-2 for cartilage tissue engineering. For bone tissue engineering HA-based hydrogels have been used in conjunction with heparin for BMP-2 delivery *in vitro* and *in vivo*. [136] In this study a rapid burst release of BMP-2 in non-heparin hydrogels was observed, with sustained release only seen in heparin containing hydrogels, which in turn maintained the osteogenic potential of BMP-2 over 28 days. Another study by Hintze, Miron [137] compared HA, sulfated HA and CS hydrogels, and found that, native HA, low sulfated HA and CS showed low affinity for all TGF- β isoforms. Specifically, the highly-sulfated HA had the greatest affinity for TGF- β 1 and TGF- β 2 but not TGF- β 3. [138]

Overall, HA has proven to be a favorable material for various tissue engineering applications as it contains the important CD44 receptor and is capable of binding to important tissue regenerative growth factors. Some studies in the field also suggest that by decorating HA with sulfated materials such as heparin, perlecan and CS, it is possible to significantly increase its affinity towards important growth factors for skeletal tissue engineering as well as delaying their release in a controlled manner.

Chitosan

Chitosan is a non-sulfated, linear polysaccharide with a semi-crystalline and biodegradable nature. It's typically derived from chitin extracted from insects, crustaceans and fungi (Figure 2). Chitosan is known to have intrinsic antimicrobial properties against fungi and bacteria. [139] The molecular weight of chitosan ranges from 300 – 1000 kD and it is composed of glucosamine and N-acetyl glucosamine linked by β (1–4) glycosidic bonds. [140] Notably, chitosan behaves as a polycation under acidic conditions, and thus is capable of forming hydrogels in the presence of polyanions and polyelectrolytes. Additionally, the degradability of chitosan directly relates to its degree of crystallinity and can thus be tailored to correspond to the targeted tissue. [141]

To even further improve the already impressive biological properties of chitosan, tissue engineers have recently tried to modify its polymeric backbone with sulfate groups. For instance, Cao, Werkmeister [142] transformed chitosan into 2-N, 6-O-sulfated chitosan (2,6SCS); and demonstrated that this particular sulfated chitosan is useful for sustained and dose-dependent BMP-2 delivery among many sulfated variants. [142] In a follow-up study

they made a comparison between BMP-2-gelatin (G)-based scaffolds, BMP-2 loaded 2,6SCS chitosan nanoparticles (BMP-2/NPs) incorporated into these gelatin scaffolds (BMP-2/S-NP/G) and a BMP2-2,6SCS-G composite. To this end, the authors found that the BMP-2/S-NP/G variant could significantly prolong the growth factor release and up-regulate *in vitro* ALP activity as compared to the other variants (Figure 4); something which was thought to be associated with the synergistic action of released BMP-2 and the unique material properties of 2,6SCS sulfated nanoparticles.[143] Interestingly, the addition of nano-particles also had an impact on the mechanical properties of the scaffold, thereby significantly prolonging its degradation time, to create an optimal condition for balancing scaffold removal with the deposition of fresh bone tissue. Building on these results, a recent approach by Pan, Chen [144] demonstrated that 2,6SCS can also be used to improve the angiogenic and osteogenic capacity of BMP-2, confirmed both on a protein and genetic level. In another recent study, Cao et al. used 2,6SCS in combination with poly(lactide-co-glycolide) (PLGA), to manufacture a composite scaffold (S-PLGA). Here they demonstrated that the BMP-2 binding efficiency within the PLGA scaffold could increase almost 10-fold in the presence of 2,6SCS. The release profiles of BMP-2 were 30% slower in S-PLGA scaffolds as compared to pristine PLGA. In the same study, BMSC cells showed an elongated and spindle-shaped morphology when interacting with the hydrophilic surface of S-PLGA.[128] Additionally, these cells were seen to circumvent Noggin inhibition, a BMP antagonist that binds extracellular BMP-2, which in turns inhibits important receptor interactions ultimately leading to reduced osteogenic capacity. Modification of the chitosan backbone with arginine yields a water-soluble molecule that is able to interact efficiently within the biological environment in contrast to the acid soluble starting material. Sulfate modification of this molecule has been achieved at the 2N as well as C2, C3 and C6 positions on the chitosan backbone.[145, 146] These sulfated derivatives bind and signal members of the fibroblast growth factor family replicating the activities of HS. While chitosan-arginine has been reported to promote an osteogenic phenotype in primary human fetal chondroblasts in the absence of osteogenic medium, sulfated chitosan-arginine promoted a chondrogenic phenotype in these same cells.[145] These data demonstrate how subtle changes in sulfation affect cell phenotype and can direct stem cell differentiation.

In summary, the high abundance of chitosan in nature along with its favorable biocompatible and biodegradable properties makes it an attractive biomaterial for skeletal tissue engineering. The modification of chitosan with sulfate groups can further improve the

already amazing bioactivity of this material. Indeed, the controlled introduction of sulfate groups onto chitosan's backbone can expand its use as a potential coagulator and a growth factor delivery vehicle.[144] Interestingly, the cationic nature of chitosan enables negative GAGs and proteoglycans to easily be incorporated into such scaffolds to promote better tissue regeneration. What's more, sulfated chitosan is in many ways structurally similar to GAGs, and thus share many of the same biological properties; as its capable of modulating both cell morphology and function – two important hallmarks of cell proliferation and differentiation.[147, 148] Overall, these exciting biomaterial properties of chitosan justify it's continued usage as a novel biomaterial in orthopaedic tissue engineering applications.

Alginate

Alginate is a sustainable polysaccharide derived from brown algae (Pheophyceae) and less frequently from gram-negative bacteria (Azotobacter and Pseudomonas sp.). Alginates are linear-anionic polymers with favorable biocompatibility for various tissue engineering applications (Figure 2).[149, 150] Notably, alginate has the capacity to form ionic hydrogel networks through chelation with divalent cations, such as Ca^{2+} , broadening its use towards drug delivery[151]. Additionally, due to the innate adhesive and tailorable shear thinning viscoelastic properties of alginate it has found widespread use in bioprinting applications.[152-154] As with other plant-based hydrogels, alginate does not natively support cell adhesion and has been described as a “blank slate” by many engineers in the field.[155] Even still, alginate can be customised through sulfation and peptide modifications to control the phenotypes of encapsulated osteoblasts,[156] chondrocytes[157] and hMSCs.[158]

Alginate sulfation based on sulfur trioxide (SO_3) [159] and sulfuric acid[160] treatments have been widely used over the years. In this regard, a number of studies have shown that such sulfated alginates can retain growth factors and promote chondrogenesis through various cellular signaling pathways;[161] and for these reasons they are considered as heparin analogues (Figure 5). Along these lines, Mhanna, Kashyap [162] employed an SO_3 /pyridine method of alginate sulfation for cartilage tissue engineering. In this study, the formation of ionic networks was restricted to a degree of sulfation of 0.8 (per monosaccharide unit), as higher degrees of sulfation (2.6) did not facilitate hydrogel formation, possibly due to strong electrostatic forces and/or steric effects between adjacent polymers. Interestingly, they found that sulfation maintained the proliferative capacity as well as phenotype of encapsulated chondrocytes, in contrast to previous studies showing initial dedifferentiation in a non-

sulfated hydrogel microenvironments.[163-165] The introduced sulfate groups also influenced Ras homolog gene family member A (RhoA) activity, which is known to be associated with chondrocyte proliferation and differentiation[166]; though the expression of collagen I and collagen II as well as proteoglycan synthesis was not significantly impacted. In another study, a bio-ink made up of sulfated alginate was used for sustained delivery of BMP-2 and osteogenesis of osteoblast cells.[45] The results showed that bio-inks exhibited an improved retention of BMP-2 in 3D-printed scaffolds. Most importantly, *in vitro* cell printing experiments revealed enhanced proliferation, as well as, osteogenesis in the hydrogels containing alginate-sulfate compared to the control bio-ink (made from pristine alginate) as indicated by increased level of ALP activity and calcium deposition. The results suggested that sulfated alginate bio-inks induced higher level of osteogenesis, by increasing the stability and retention of the loaded BMP-2.

Thus, sulfated alginate-based scaffolds are promising alternatives to mammalian derived GAGs due to their biocompatibility, low immunogenicity, protein retention capacity and the great variety of readily implementable gelling and functionalisation strategies that can improve their bioactivity. Their extensive and continued use will definitely empower researchers with the knowledge to better understand the regulatory role of sulfated-alginate in extracellular and intracellular interaction, something, which hopefully will lead to their more frequent use in skeletal tissue engineering in the foreseeable future.

Cellulose

Cellulose is the most abundant natural polysaccharide available in the world.[167, 168] Its chemical structure consists of unsubstituted, linear glucose homosaccharide with six available hydroxyl groups. Intriguingly, it has been seldom used in tissue engineering, potentially due to difficulties in hydrogel assembly caused by solubility inadequacies.[169] The sulfation of cellulose can improve solubility, through the disruption of intermolecular hydrogen-bonds[170] to potentially broaden its applicability towards various tissue engineering applications.[171]

One study by Huang, Molina [172] explored the use of sulfated cellulose scaffolds for cartilage tissue engineering. Initially MSC induction media was spiked with a fully sulfated form of sodium cellulose (NaCS) leading to a significant upregulation of collagen II and aggrecan. In the same study, NaCS was combined with gelatin to develop scaffolds through electrospinning. Interestingly, the scaffolds with the lowest concentration (0.1%) of NaCS added to induction media resulted in the highest production of collagen II both on a protein

and genetic level after 56 days of culture. Additionally, cells on the 1% and 5% NaCS/Gelatin-based scaffold showed low collagen X production, suggesting higher NaCS may result in a reduced propensity towards hypertrophy. These higher sulfate concentrations may have an inhibitory effect on chondrogenesis because of irreversible growth factor-biomaterial bindings, which in turn can comprise the release and delivery of TGF- β 3 to the targeted cells.[173] The same group took this a step further and introduced partially sulfated cellulose (pSC) into gelatin hydrogels instead, and discovered an enhanced expression of chondrogenic markers (collagen II/collagen I ratio, aggrecan and SOX9) upon increasing pSC concentration in the scaffolds, indicating the potential of pSC as a scaffolding material for cartilage tissue engineering.[174]

For these reasons, cellulose sulfate is an interesting vehicle for growth factor delivery in cartilage tissue engineering and could have broader uses in the foreseeable future due to its abundance, sustainability and reduced immunogenicity. Specifically, the backbone sulfation of cellulose allows for precise control over the sulfation pattern and sulfation degree, and thereby enables the biological properties of such scaffolds to be fine-tuned in a customizable manner. The range of available chemical modifications can also pave the way for tuneable mechanical and pharmaceutical properties, and could thereby potentially enable an even greater variety of biomaterials. [175] [176]

3. Tissue engineering

While sulfated polysaccharides have been shown to successfully act as delivery vehicles for growth factors in an *in vitro* environment, their ability to elicit this response in an *in vivo* model needs to be evaluated as well. Indeed, many tissue engineering approaches have shown significant benefits in *in vitro* studies yet when they progress to animals models they show some limitations.[29] Understanding, whether the successful *in vitro* strategies also show promise in an *in vivo* setting, is therefore critical to successfully translate tissue engineering strategies from the laboratory and into the clinic. This section, highlights recent advances in translating the hard tissue regenerative potential of scaffolds made from sulfated polysaccharides in various animal models both alone, in combination with various growth factor or with other biopolymers (Table1).

3.1 Bone

The number of people at risk of bone fractures has grown steadily in most parts of the world due to the ageing population. In 2015 around 160 million people worldwide experienced a bone fracture; a number that is expected to double to 320 million by the end of 2040.[177] Traditional clinical therapies for mending bone fractures rely on various forms of casts to fixate the broken fracture to enable the native bone to heal itself on its own terms, however, native bone displays a restrictive regenerative capacity, that is haunted by a number of challenges including non-anatomical reduction of the fracture, a-vascular necrosis, as well as non-union and mal-union fracture healing.[178] These issues are more prevalent in older people and will thus grow steadily in the near future as the median lifetime is expected to increase significantly in the coming decades. Autologous bone grafts are commonly utilized to promote osteoconduction and osteoinduction in bone defects to avoid the abovementioned scenarios. While these grafts have shown some promise for healing bone defects, they require multiple invasive surgeries and are limited by low availability and donor site morbidity associated with relocating native bone tissue from the patient's own bone and into the defect site.[179] Allografts on the other hand are limited due to the lack of available donor tissues and unwanted foreign body responses; and bone implants in some cases do not facilitate sufficient bone healing and therefore revisions surgeries are common with this methodology.[180]

For these reasons, a number of bone tissue engineering strategies have emerged to address this critical challenge by delivering the promise of a better method to mend bone defects.[181] As such, these approaches rely on developing synthetic bone tissues by combining 3D biomaterials with stem cells either exogenously or by recruiting them from native bone-tissue in a post-implantation scenario. The 3D biomaterials have the potential to drive stem cells into bone-like cells that under the right conditions can form mature tissues either in the laboratory or within the body depending on which one of the abovementioned strategies has been employed (Figure 1). However, many of the tissue engineered scaffolds explored to date have not reached this full potential and in many cases fall short of the performance of autografts.[182] A number of studies, including those by Wang and Yeung [179] and Lee, Silva [16] suggest that such results could be related to the uncontrolled release of growth factors that collaterally interfere with untargeted cells. As sulfated polysaccharides can bind and regulate the signalling of a number of important growth factors they are likely to be essential components of next-generation biomaterials for bone tissue engineering.

Indeed, sulfated polysaccharides are considered one of the most important biological and mechanical components of the native ECM of hard tissues.[183] They have therefore in recent years emerged as new and promising building blocks for bone tissue engineering scaffolds.[184] Heparin is one of the most widely employed sulfated polysaccharides in this respect, due to its ability to capture, stabilize and present growth factors to bone progenitor cells in a controllable manner. For instance, Yang, La [185] developed heparin-conjugated fibrinogen (HCF) injectable scaffolds for orthotopic *in vivo* models (hind limb muscle pockets in rats) to control the release of BMP-2, with the aim of enhancing the new bone formation. Initially, *in vitro* experiments with osteoblast cells showed that BMP-2 released from the HCF hydrogels induced a significantly higher level of ALP activity, when compared to BMP-2 released from the fibrin hydrogels, indicating that BMP-2 released from HCF is bioactive and long-term delivery of BMP-2 is advantageous over short-term delivery for bone regeneration. *In vivo*, this prolonged activity ultimately translated itself into significant improvements in bone mineralization when compared with pristine fibrin scaffolds. Notably, by using heparin, they were able to obtain a similar amount of new tissue formation with lower concentrations of BMP-2 than previously reported in the literature.[186] However, some studies have reported that exogenous heparin under certain circumstances reduces the bioactivity of osteogenic biomolecules and can thus compromise the bone healing process, by inhibiting the binding of BMP-2 to the BMP receptor. What's more, the potent anticoagulant activity of heparin is, by many in the field, thought to be counterproductive for bone growth.[187]

To address these issues, sulfated chitosan, has been used as an alternative due to its good biocompatibility and similar growth factor binding ability as heparin without the abovementioned native biological issues associated with heparin.[188] In this direction, Zhou, Qian [189] synthesized BMP-2 loaded chitosan with varying degrees of sulfation and compared their responses *in vivo*. These *in vivo* results revealed that the most sulfated chitosan-based scaffold was the best promoter of BMP-2 bioactivity and could even surpass the bone regeneration capacity of heparin-based scaffolds. Similarly, Lü, Bai [190] and Lü, Bai [190] developed a self-healing, biocompatible and injectable dual cross-linked CS-based hydrogels for *in vivo* delivery of BMP-4. This hydrogel was crosslinked through both diels-alder (DA) and acylhydrazone bonds; and the authors used these bonding schemes to fine-tune various hydrogel properties such as rigidity and degradation. Through this sophisticated crosslinking scheme they were also able to manufacture a superior hydrogel, which could

prevent excessive hydrogel swelling *in vivo*; and thereby prevent poor stem cell differentiation and tissue regeneration.[191] In both instances, histology staining's demonstrated new bone formation in the BMP-4 loaded hydrogel samples after 12 weeks, with controls primarily stimulating fibrous tissue growth. Additionally, initial sproutings of blood vessels were observed. In another noteworthy study, Kim, Lee [183] evaluated the inclusion of UV-crosslinked methacrylated CS (MeCS) in PEGDA hydrogels at various concentrations in terms of their bone regenerative properties within the body (Figure 6). Specifically, these scaffolds were implanted in critical sized calvarial defects (4mm diameter) in six-week-old female mice (n = 4) for up-to eight weeks. Interestingly, scaffolds containing the highest concentration of CS induced the most effective bone formation evidenced by larger bone mineralization density. This was speculated to arise from the ability of the sulfate groups within CS to bind to calcium ions and facilitate the formation of fresh hydroxyapatite; one of the most important components of the mineral phase of bone. Additionally, Hematoxylin, Eosin and Masson's trichrome staining's also showed significant improvements in bone tissue formation with increasing CS concentration.

Although, a wide range of sulfated polysaccharides have been studied in the literature, these biomaterials are seldom employed in clinical treatments due to the lack of more standardized clinical studies.[192] Indeed, a number of important parameters such as the size of the bone defect, the place of the defect, the implanted cell type, and implantation time needs to be considered to fully unravel the bone tissue engineering potential of such scaffolds. Unfortunately, these parameters have not been studied enough to turn this promising strategy into a clinical therapy which can benefit the many sufferers of bone disorders.[192] Consequently, more in-depth *in vivo* studies are necessary to validate the efficiency of sulfated polysaccharides for bone tissue engineering, and to identify the best combination to use in the clinic.

3.2 Cartilage

The primary cause of cartilage damage within the body is due to osteoarthritis (OA) in articular cartilage. The clinical treatment for OA is currently suboptimal as the “state-of-the-art” surgical approaches are limited in terms of their efficacy and high invasiveness. First stage interventions include arthroscopy, which involves the flushing and removal of damaged cartilage and meniscus.[193] For more severe cases, the implantation of autologous osteochondral graft (mosaicplasty) into the defect site and surgical drilling into the

subchondral bone (microfracturing) can be employed.[194] However, unfortunately both measures are controversial as they often result in fibrous cartilage rather than native articular cartilage.[195] For the most severe cases, extremely invasive and costly total knee replacements can be performed.[196] Notably, these measures are aimed at slowing the impact of OA without actively regenerating native cartilage.

Recently, techniques such as stem cell therapy have been used to regenerate cartilage tissue, by injecting regenerative cells into the damaged region.[44, 49, 176, 197] This technique is limited by low cell retention and a low cell viability, caused by the shear-forces that cells experience when passing through the thin injection needle. It also does not provide the cells with a 3D microenvironment to properly differentiate them into the required tissues. The usage of hydrogels can provide a mechanical shield during the needle-injection phase and provide a suitable 3D microenvironment for guiding cells into the desired cell phenotypes in a post-injection scenario. Especially, sulfated hydrogels hold great promise in this respect, since they display high affinity towards important growth factors for cartilage regeneration; and in many ways resemble – CS - one of the most important components of the native cartilage ECM. Indeed, such biopolymers have recently been used to develop scaffolds with the capacity to deliver growth factors such as BMP-2 and TGF- β 3 in a sustainable manner to significantly improve the cellular performance of chondrocytes.[76, 77] In another related study by Han, Wang [198] a mussel inspired CS-based hydrogel was created for enhanced adhesion between graft and native cartilage tissue (Figure 7). Specifically, the inclusion of CS promoted an upregulation of chondrogenic differentiation markers such as aggrecan and collagen II. The scaffolds were also evaluated in a full thickness defects (diameter: 3.5 mm; thickness: 5 mm) in the patella groves in the right legs of white rabbits (n = 8). Following a three-month implantation period, the scaffolds showed significantly higher tissue formation in terms of Modified O'Driscoll and International Cartilage Repair Society grading scores.

In another study Feng, Lin [53] conjugated sulfate groups onto the backbone of methacrylated hyaluronic acid (MeHA) in order to deliver growth factors in a articular cartilage rodent (n = 10) model in a controlled and sustainable manner (Figure 8). Typically, HA is degraded rapidly by hyaluronidases *in vivo* and lacks high protein binding affinity. They found that the introduction of sulfate groups reduced the degradation and deformation of hydrogel scaffolds and promoted cartilage matrix deposition, as indicated by immunohistochemical stainings of collagen II and CS, following 4 weeks *in vivo*.

Additionally, the sulfated-HA in combination with hMSCs was capable of attracting and retaining supplemented TGF- β 1, and thereby promoting chondrogenesis and suppressing hypertrophy. Overall, the paper by Feng, Lin [53] demonstrates that sulfated HA hydrogels enable the generation of high quality neocartilage via intra-articular injection.

The abovementioned studies on using sulfated polysaccharides for cartilage regeneration clearly demonstrate the great promise that they hold for the field of cartilage tissue engineering. Indeed, considering the importance of cell therapy in treating acute cartilage injuries, sulfated polysaccharides can be ideal candidates for biodegradable scaffolds to temporarily support the chondrocytes until they are replaced by matrix components synthesized from the implanted cells. Collectively, the use of such scaffolds is expected to reduce chondrocyte leakage from the transplant site, provide a more homogeneous chondrocyte distribution, and lessen graft hypertrophy.[199] Regardless, in order to fully explore the potential of such scaffolds in cartilage tissue engineering, we need to consider important parameters such as lesion location and damage size, activity level and patient's age. These parameters are by many in the field considered the important parameters when it comes down to choosing the right cartilage repair techniques and controlling the outcome of the treatment.[200] Finally, the biomaterials utilised in *in vivo* cartilage tissue engineering need to demonstrate appropriate biomechanical and biochemical cues without triggering immune responses. Therefore, biomaterials and cell therapy techniques should also be compared to 'gold standard' techniques such as microfracture and grafting in order to accurately gauge their efficacy *in vivo*. The continued investigations into the usage of sulfated polysaccharides as growth factor delivery vehicles is also needed to fully elucidate their potential as tissue engineering scaffolds for cartilage regeneration.

3.3 Osteochondral

Defects that impact both the articular cartilage and the underlying subchondral tissues are termed osteochondral defects. Such lesions are caused by tissue degradation from aging, sports injuries or severe cases of osteoarthritis. They typically result in joint instability, significant discomfort for the patient and loss of patient mobility. Much like cartilage, osteochondral defects can be treated through microfracturing, allografting and mosaicplasty, or even total knee replacements, however, all of these therapies unfortunately have similar issues as those briefly mentioned in the previous section.[201] The abovementioned tissue

engineering approaches could remedy these shortcomings by recapitulating the highly hierarchal structure of osteochondral defects.

In this direction, Zhou, Zhang [202] recently combined silk fibroin with CS to develop a composite scaffold that could mend osteochondral defects in a rabbit animal model. Indeed, this composite material produced greater neo-tissue formation and improved structural restoration compared to the pristine silk scaffold at 6 and 12 weeks as evident from an International Cartilage Repair Society histological analysis (Figure 9). Additionally, when analysed *in vitro*, the composite scaffold was seen to maintain better chondrocyte morphology compared to the silk scaffold alone, in combination with a higher expression of SOX9, collagen II, aggrecan and lower expression of TNF- α 2 (an important inflammation marker) (Figure 9). In a similar vein, Liao, Qu [203] implanted a biomaterial composite consisting of methacrylated CS and poly(ethylene glycol) methyl ether- ϵ -caprolactone-acryloyl chloride (MPEG-PCL-AC) incorporated with graphene oxide, into full-thickness osteochondral defects (thickness: 3mm, diameter: 4mm, n = 27) in the hind limbs of rabbits. When combined with chondrocytes, the scaffold was seen to improve chondrocyte morphology, integration, and subchondral bone formation. Notably, this strategy could rapidly induce the formation of both new and thicker cartilage tissue as compared to a cell-free scaffold.

In a recent study, Lee, Luo [204] combined HS with a hyaluronic acid hydrogel for osteochondral repair in a rabbit model. Accordingly, osteochondral defects treated with the composite hydrogel showed higher MOCART and ICRS I scores compared to the hyaluronic acid gel alone group, indicating improved filling of the defects and integration with surrounding host tissue. In addition, MRI analyses showed more intact surface and higher subchondral bone formation in defects treated with the composite hydrogel (as compared to HA gels alone). The lower amount of new bone formation in HA alone treated group may be due to insufficient calcification of newly formed ECM in the absence of HS. Most remarkably, regenerated hyaline cartilage in the chondral layer was only observed following treatment with the composite hydrogel. This finding can be attributed to enhanced subchondral bone regeneration as articular cartilage growth relies on the sufficiency of subchondral bone for mechanical support and nutrition.

Another noteworthy study used a heparin immobilised polycaprolactone (PCL)/Pluronic F127 scaffold combined with TGF- β 2 and BMP-7 to facilitate even more cartilage tissue formation as compared to PCL/Pluronic scaffolds alone. However, no significant histological

differences following implantation into large (diameter = 6mm, depth = 3mm) distal femur defects in rabbits (n = 12) was seen in this study.[205] Finally, Re'em, Witte [206] recently created a bilayer scaffold with alginate-sulfate incorporating both TGF- β 2 and BMP-4. This scaffold was subsequently implanted into subchondral defects (diameter = 3mm, depth = 3mm) in the femur of rabbits. Encapsulated hMSC's were successfully differentiated into both osteoblasts and chondrocytes at respective layers over 4 weeks, confirming the controlled release of the growth factors. Additionally, the cartilage–bone interface formation remained the same in hMSC incorporated scaffolds, indicating that native cells were able to migrate into the scaffolds and sense the biological cues spatially present in there, and respond accordingly by differentiating to the appropriate cellular lineage.

History has shown that applying promising laboratory strategies to animal models is not always as successful. Even a rudimentary understanding, through the use of pilot studies, of the *in vivo* efficacy of such techniques can create a much more efficient process for producing novel, viable tissue engineering solutions. For these reasons, sulfated-scaffolds for osteochondral tissue engineering are also beginning to be translated into *in vivo* environments. Most often, these materials are used in composites to capitalise upon the benefits of multiple materials and to develop the hierarchical scaffolding architecture needed for optimal osteochondral repair. To this end, the effects of growth factor delivery and improved cellular performance observed in *in vitro* studies appear to translate into *in vivo* outcomes. Additionally, the studies reviewed here indicate that sulfated polysaccharide do not elicit any significant inflammatory responses when implanted *in vivo*, confirming that they indeed are suitable biomaterials for osteochondral tissue engineering.

4. Conclusion and future directions

Tissue engineering has shown tremendous potential in several facets of biomedicine, particularly in skeletal tissue engineering. With the ongoing development of novel sulfated biomaterials along with sophisticated *in vitro* culturing systems tissue engineering will enhance our capacity to recapitulate bone and cartilage regeneration through the sustained delivery of relevant growth factors. Overwhelmingly, the most commonly studied and successful naturally sulfated biomaterials include CS and heparan sulfate and its analogues. The benefits that these naturally sulfated ECM components provide can be chemically incorporated into non-sulfated biomaterials. Specifically, HA and chitosan sulfation allows for the controlled binding and release of growth factors in a localised environment. The use

of composite materials in tissue engineering is omnipresent and can capitalise upon the benefits of multiple materials. These four materials, CS, Hep/HS, HA and chitosan, can be easily utilised in a composite system, where the scaffold can provide cells with controlled, prolonged and protected growth factor delivery. Though, the translational capacity of animal-derived sulfated biomaterials is limited *in vivo* due to immunogenicity, further exploration into plant-derived substrates could be a worthy endeavour. Intriguingly, as these materials don't have specific enzymes for degradation their use could potentially extend growth factor delivery beyond the body's native capacity. Many areas within the vibrant field of tissue engineering could readily benefit from the utilization of sulfated biomaterials as a vehicle for providing growth factors to the target tissues to elicit improved cellular performance both *in vitro* and *in vivo*.

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Figures and Figure captions

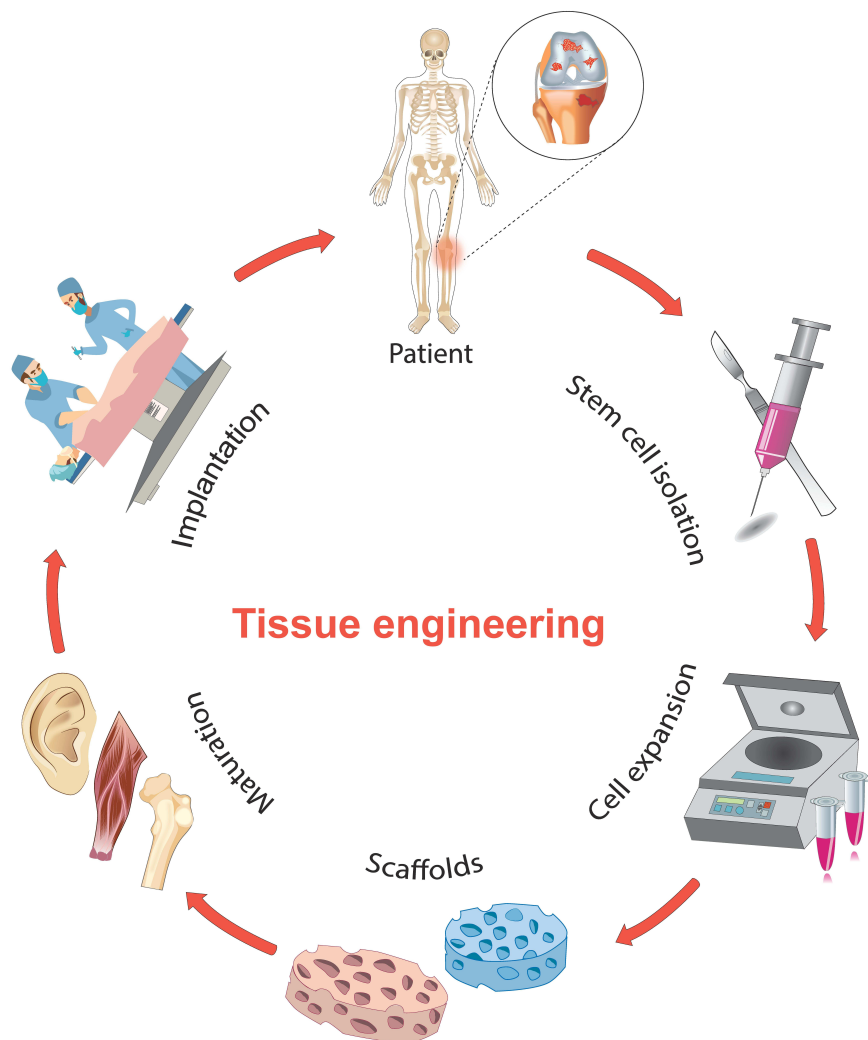


Fig. 1: A schematic showing the core-principles behind tissue engineering.

Fig. 2: Some of the most important sulfated polysaccharides reviewed herein have been highlighted in this figure along with their derivation source and chemical structure.

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Fig. 3: The growth factor retention capacity of heparin. (a) Schematic showing the design principle behind the biphasic silk fibroin scaffold used in the study. (b) The heparin loading efficiency and its release profile from the scaffold is displayed here. (c) The sustained release of TGF- β 2 and GDF5 is displayed here. Crosslinked heparin significantly delayed the growth factor release. Modified from [78], with permission from Elsevier, Copyright 2018.

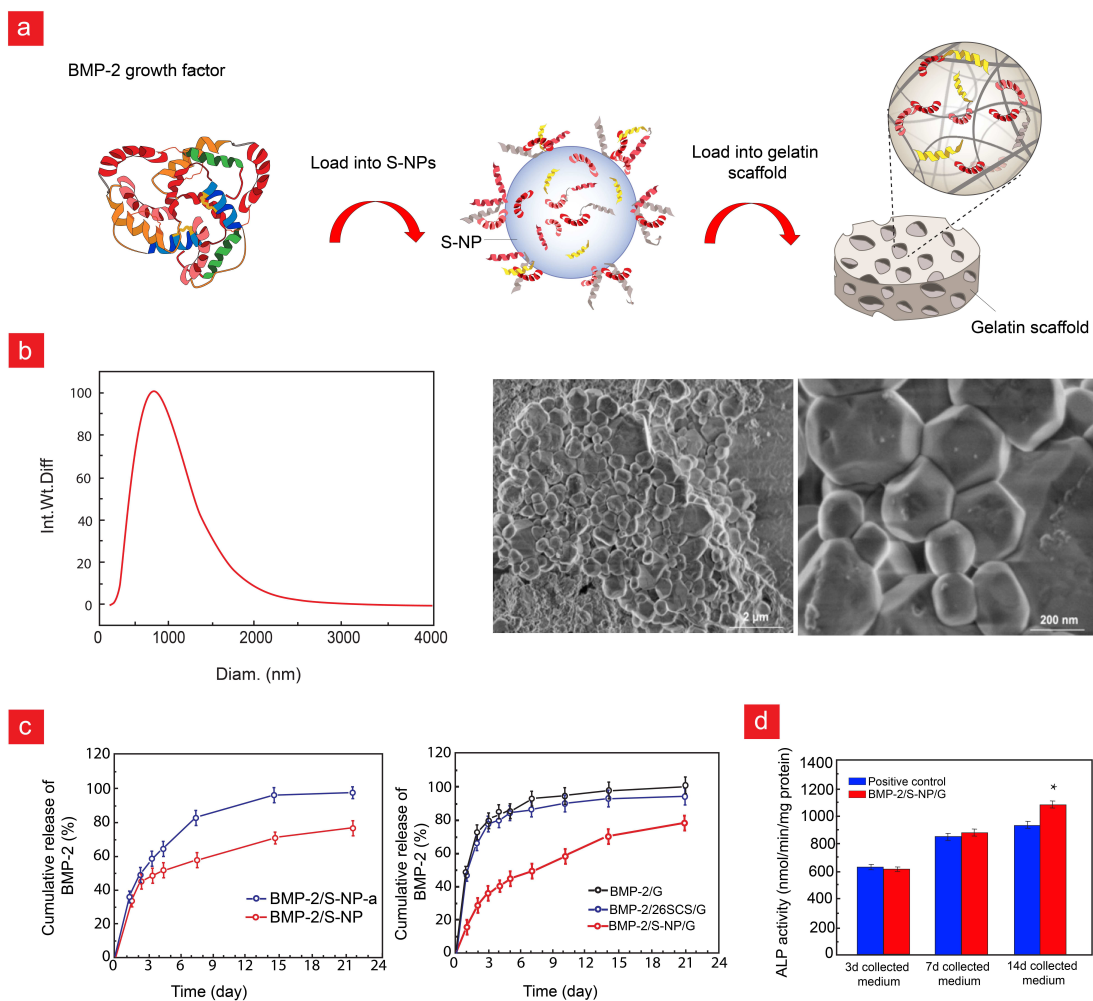


Fig. 4: The growth factor retention capacity of sulfated chitosan. (a) Schematic showing the design principle behind the S-NP incorporated gelatin scaffolds. (b) The size distribution and scanning electron microscopy images (SEM) of the S-NP's are displayed here. (c) The sustained release of BMP-2 from the scaffolds employed in this study is displayed here.

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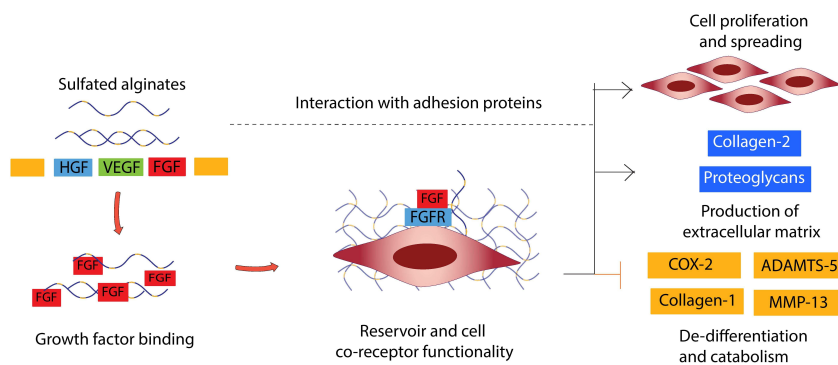


Fig. 5: A schematic showing the growth factor bind properties of sulfated alginates and their ability to promote chondrogenesis through important signalling pathways. Modified from [161], with permission from MDPI, Copyright 2017.

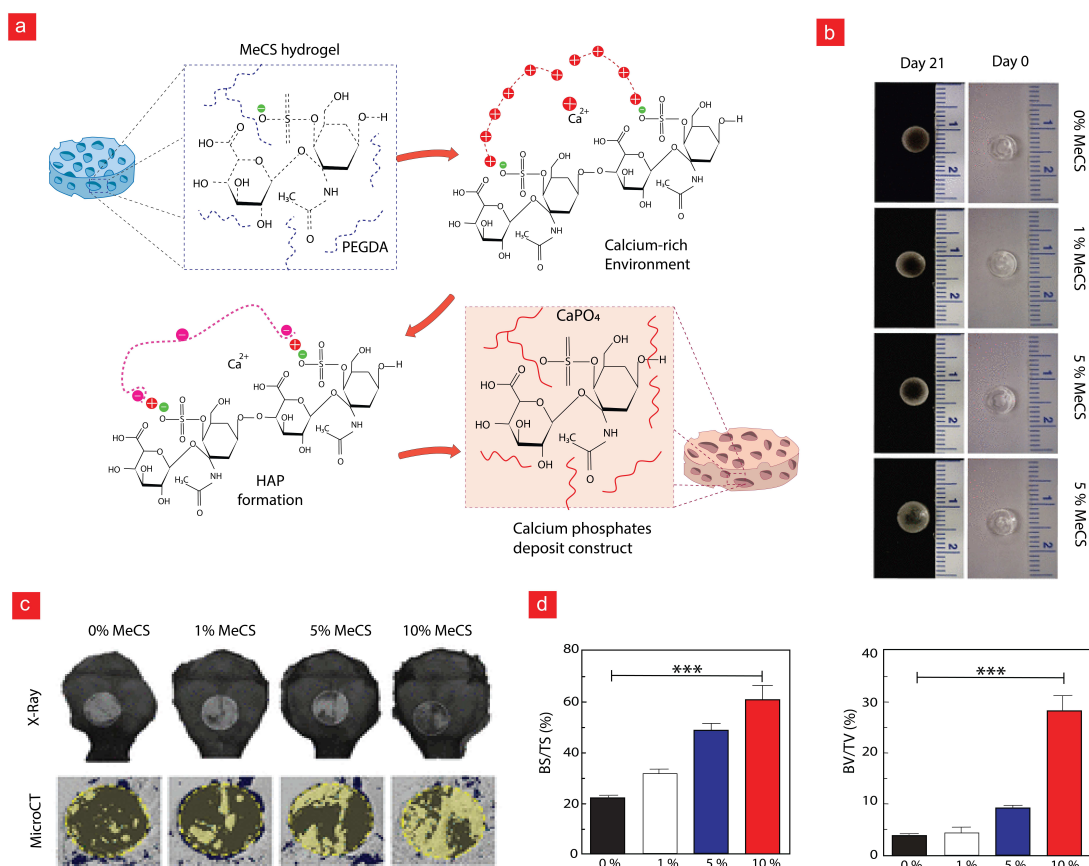


Fig. 6: A chondroitin (CS)-based scaffold for bone tissue engineering. (a) The manufacturing of the PEGDA-MeCS hydrogel and its hydroxyapatite (HAP) formation capacity is shown here. (b) The calcification and HAP formation of the cell-laden hydrogels after 21 days are shown here through photographic images of the hydrogels at relevant time points. (c) The bone regenerative capacity of the respective scaffolds incorporating different concentrations of CS was quantified through Micro-CT analysis after 8 weeks of implantation. (d) The bone area (BS/TS) and bone volume (BV/TV) were also calculated and are displayed here. Adapted with permission from [183]. Copyright (2017), American Chemical Society.

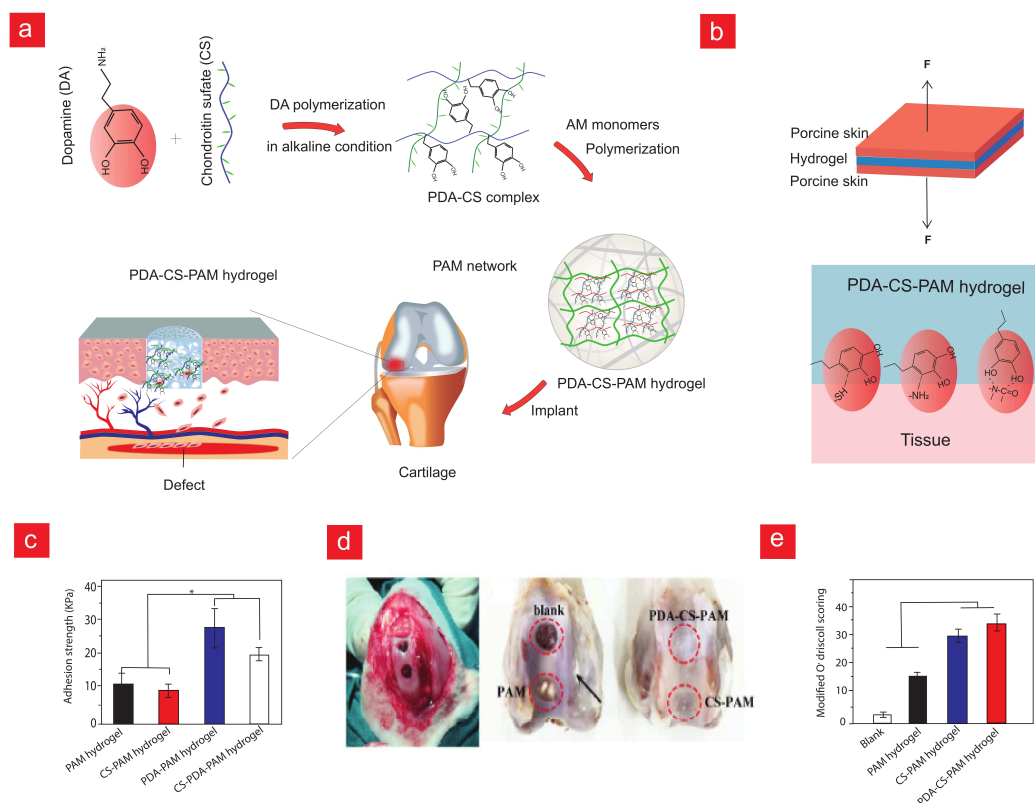


Fig. 7: A tissue adhesive CS-based scaffold for cartilage tissue engineering. (a) The CS-based scaffold was made tissue adhesive by polymerizing dopamine (DA) and acrylamide (AM) into it. (b) The tissue adhesive properties of the scaffold was mediated by the many amino groups present on PDA and PAM. (c) The adhesion strength of the various manufactured scaffolds towards porcine skin is shown here. (d) The cartilage regenerative potential was highest for the PDA-CS-PAM hydrogel. (e) This was further validated by analysing the Modified O' driscoll scoring for the implanted scaffolds after 3 months of implantation. Adapted with permission from [198]. Copyright (2018). American Chemical Society.

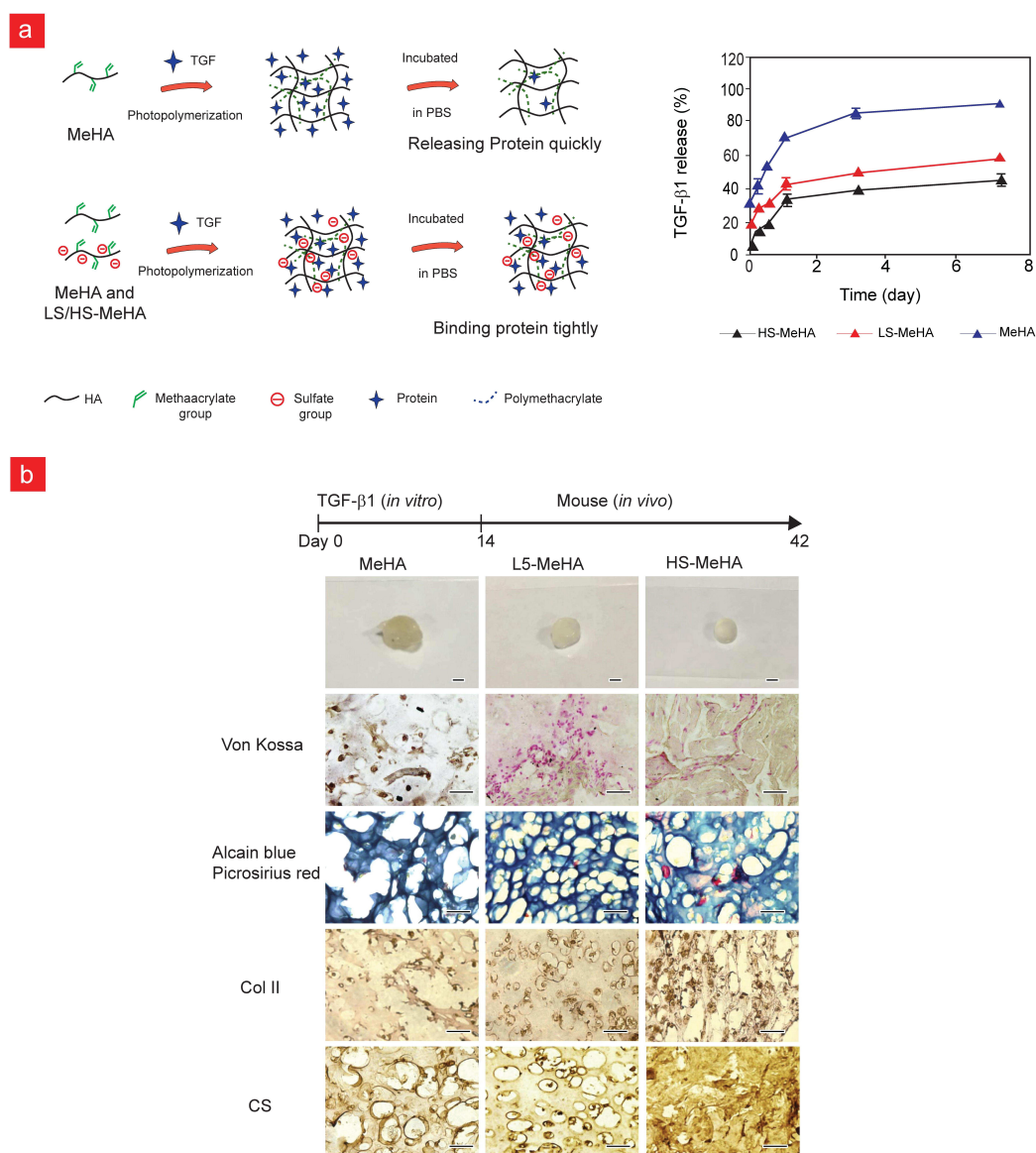


Fig. 8: A sulfated hyaluronic acid scaffold for cartilage tissue engineering. (a) The manufacturing scheme behind the scaffolds are shown here, where LS-MeHA and HS-MeHA are short for low sulfated and high sulfated methacrylated hyaluronic acid (HA), respectively. (b) The TGF- β 1 retention capacity of the various scaffolds employed in the study is shown here. (c) Histological staining of the respective hMSCs-laden scaffolds after 42 days of implantation. Modified from [53], with permission from Elsevier, Copyright 2017.

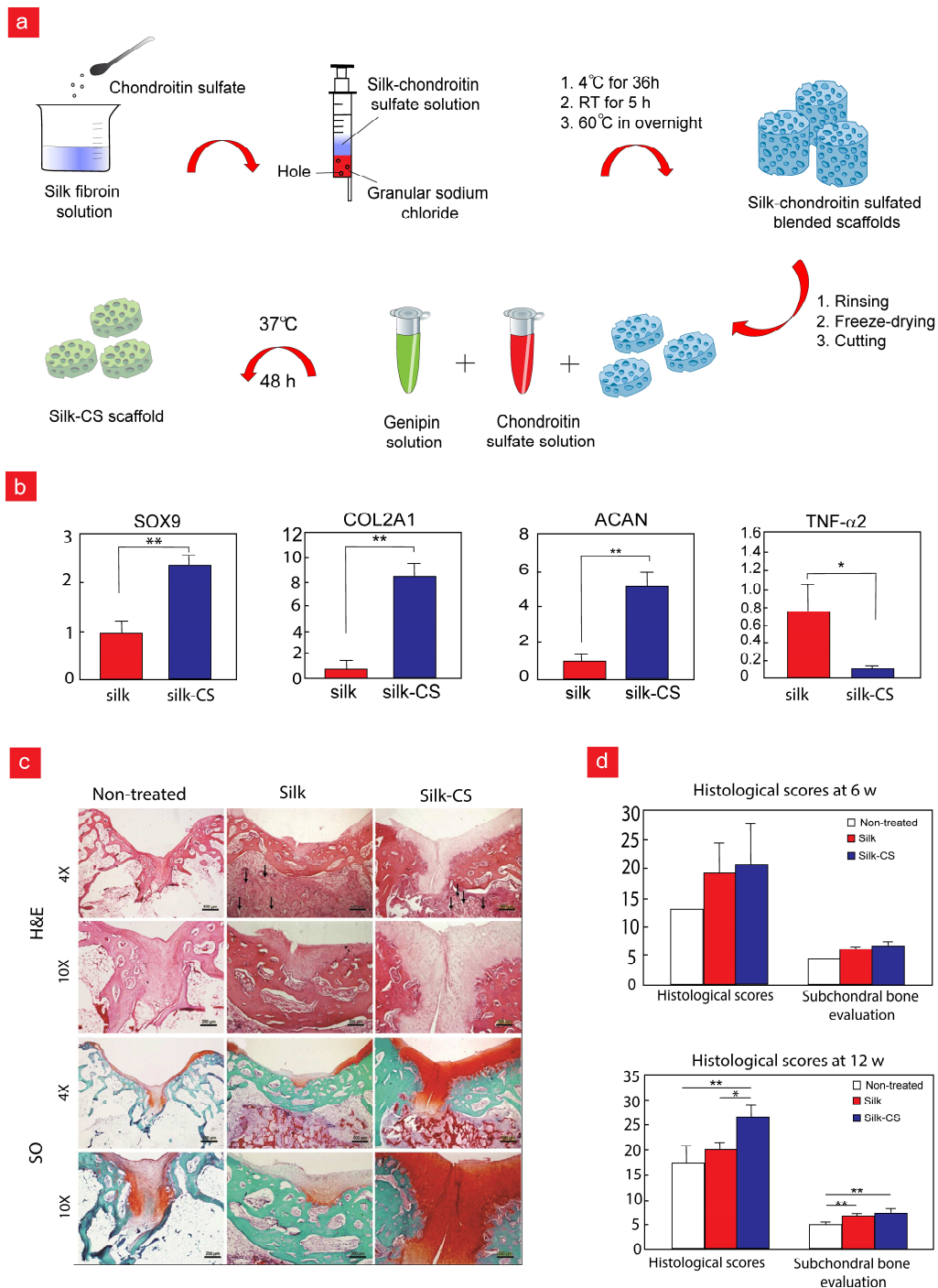


Fig. 9: A Silk-CS-based scaffold for osteochondral tissue engineering. (a) The manufacturing process behind the Silk-CS scaffold is shown here. (b) The chondrogenic and anti-inflammatory capacity of the Silk-CS was quantified from expression of relevant gene markers. (c) Histological evaluation of the scaffolds after 12 weeks of implantation. H&E is short for hematoxylin and eosin and SO for Safranin O. (D) The histological scores for subchondral bone formation was evaluated after 6 and 12 weeks. Modified from [202], with permission from Elsevier, Copyright 2017.

Table 1: Summary of sulfated polysaccharide-based materials used for various skeletal tissue engineering applications.

Sulfated polysaccharide Type	Scaffold Type	Growth factor(s) Type	Study type	Results	Ref.
<i>Heparin</i>	PLGA microsphere	BMP-2	Bone <i>In vitro</i> <i>MG-63 cells</i>	Enhanced osteogenesis, whilst simultaneously increasing both ALP activity and deposition of important bone minerals	[73]
<i>Heparin</i>	Biphasic silk fibroin discs	TGF- β_2 & GDF5	Cartilage <i>In vitro</i> <i>Adipose-derived mesenchymal stem cells</i>	Controlled release of TGF- β_2 and GDF5 from the scaffold up-regulated chondrogenic markers	[78]
<i>Heparan sulfate (HS)</i>	Collagen sponges	BMP-2	Bone <i>In vitro</i> <i>C2C12 cells</i> & <i>In vivo</i> <i>Rat ectopic model</i>	<i>In vitro</i> , delivered BMP-2 in a prolonged and sustained manner. <i>in vivo</i> , combination of BMP-2 with HS resulted in 2-fold more bone volume formation than BMP-2 treatment alone	[74]
<i>Heparan sulfate (HS)</i>	Hyaluronic acid hydrogel	-	Osteochondral <i>In vivo</i> <i>osteochondral defect in rabbit</i>	Improved filling of the defects and integration with surrounding host tissue.	[204]
<i>Chondroitin sulfate (CS)</i>	Hyaluronic acid methacrylate hydrogel	-	Cartilage <i>In vitro</i> <i>Human mesenchymal stem cells</i>	The inclusion of CS in the HA hydrogels can upregulate mRNA expression of chondrogenic markers, while decreasing expression of the hypertrophic markers	[85]
<i>Chondroitin sulfate (CS)</i>	Chitosan membrane	-	Cartilage <i>In vitro</i> <i>Pre-chondrocyte cells (ATDC5)</i>	chitosan/CS induced a higher collagen II/collagen I ratio (a characteristic of hyaline cartilage formation) after 21 days, when compared to pristine chitosan	[47]
<i>Chondroitin sulfate (CS)</i>	PLGA films coated with collagen I	-	Bone <i>In vitro</i> <i>MG-63 cells</i>	CS improved both the osteoconductivity and osteoinductivity of the (osteoblastic) MG-63 cell line, observed through the increased proliferation and upregulation of osteocalcin, as compared to pristine collagen I coatings	[96]
<i>Chondroitin sulfate (CS)</i>	Injectable fibrinogen	BMP-2	Bone	<i>In vitro</i> , BMP-2 released from the HCF hydrogels	[185]

			<i>In vitro</i> osteoblasts & <i>In vivo</i> hind limb muscle pockets in rats	induced a significantly higher level of ALP activity, when compared to BMP-2 released from the fibrinogen hydrogels. <i>In vivo</i> , BMP-2 loaded HCF hydrogels showed significant improvements in bone mineralization when compared with pristine fibrinogen scaffolds	
Chondroitin sulfate (CS)	PEGDA hydrogel	-	Bone <i>In vitro</i> Tonsil mesenchymal stem cell (hTMSC) & <i>In vivo</i> Calvarial Defect in mice	<i>In vitro</i> , induced osteogenesis differentiation of encapsulated hTMSC. <i>In vivo</i> , cell laden scaffolds containing the highest concentration of CS induced the most effective bone formation	[183]
Chondroitin sulfate (CS)	Polyacrylamide (PA) hydrogel	-	Cartilage <i>In vitro</i> Chondrocytes & <i>In vivo</i> Critical-sized cartilage defects of a rabbit	<i>In vitro</i> , the inclusion of CS promoted an upregulation of chondrogenic differentiation markers. <i>In vivo</i> , the scaffolds showed significantly higher tissue formation in terms of Modified O'Driscoll and International Cartilage Repair Society grading scores	[198]
Chondroitin sulfate (CS)	Silk fibroin	-	Osteochondral <i>In vitro</i> Chondrocytes & <i>In vivo</i> Osteochondral defect on the femoropatellar groove in rabbit	<i>In vitro</i> , higher expression of chondrogenic markers, compared to pristine silk scaffolds. <i>In vivo</i> , produced greater neo-tissue formation and improved structural restoration compared to the pristine silk scaffold at 6 and 12 weeks	[202]
Carrageenan (CAR)	PCL/chitosan membranes	-	Bone <i>In vitro</i> saos-2 cells	Among three different CAR sugar backbones, kappa (κ), iota (ι), and lambda (λ), ι -variant demonstrated significantly higher biomineralization	[112]
Carrageenan (CAR)	Chitosan hydrogel	-	Cartilage <i>In vitro</i> Pre-chondrogenic ATDC5 cells	expression of cartilage specific genes were up regulated with increasing CARs concentrations within chitosan, when compared to pristine chitosan	[52]
Fucoidan	Chitosan/TCP scaffold	-	Bone <i>In vitro</i> human bone marrow stromal cells (hBMSC)	Addition of fucoidan promoted osteocalcin and ALP production whilst supporting hBMSC growth	[118]
Fucoidan	Alginate	-	Cartilage	Encapsulated cells expressed	[120]

	hydrogel		<i>In vitro</i> <i>Human mesenchymal stem cells</i>	a higher level of chondrogenic markers and produced a higher GAG content. Cells also expressed a significantly lower level of hypertrophy markers, when compared to alginate hydrogels	
<i>Ulvan</i>	Methacrylated ulvan hydrogel	-	Bone	The lowest methacrylated-ulvan group, showed the highest ALP activity	[123]
<i>Sulfated hyaluronic acid</i>	Heparin decorated hyaluronic acid hydrogel	BMP-2	Cartilage <i>In vitro</i> <i>Murine mesenchymal stem cells</i>	Improved BMP-2 delivery and chondrogenic differentiation when compared to pristine hyaluronic acid	[133]
<i>Sulfated hyaluronic acid</i>	Perlecan-(domain I) decorated hyaluronic acid hydrogel	BMP-2	Cartilage <i>In vitro</i> <i>Murine mesenchymal stem cells</i>	Exhibited the ability to bind significantly more BMP-2 as compared to HA alone and promoted higher level of chondrogenesis	[134]
<i>Sulfated hyaluronic acid</i>	Sulfated methacrylated hyaluronic acid	TGF- β_2	Cartilage <i>In vitro</i> <i>Human mesenchymal stem cells</i> & <i>In vivo</i> <i>Rat osteoarthritis model</i>	<i>In vitro</i> , sulfated methacrylated HA hydrogels promote the chondrogenesis and suppresses the hypertrophy of encapsulated hMSCs. <i>In vivo</i> , intra-articular injections of the sulfated HA hydrogels averted the cartilage abrasion and hypertrophy in the animal osteoarthritic joints.	[53]
<i>Sulfated chitosan</i>	GelMA hydrogels loaded with sulfated chitosan nanoparticles	BMP-2	Bone <i>In vitro</i> <i>Human mesenchymal stem cells</i> & <i>In vivo</i> <i>critical-sized segmental defect in rabbit</i>	<i>In vitro</i> , this scaffold could significantly prolong the growth factor release and up-regulate ALP activity as compared to the pristine GelMA hydrogels loaded with BMP-2. <i>In vivo</i> , this scaffold provided a higher repair rate and better integrity of the healed bone as compared to the pristine GelMA hydrogels loaded with BMP-2.	[142]
<i>Sulfated chitosan</i>	PLGA scaffolds	BMP-2	Bone <i>In vitro</i> <i>C2C12 cells</i>	Improved BMP-2 adsorption and prolonged release process, increased ALP activity and cell attachment	[128]
<i>Sulfated alginate</i>	Alginate hydrogel	-	Cartilage <i>In vitro</i> <i>Chondrocyte cells</i>	Sulfation maintained the proliferative capacity as well as phenotype of encapsulated chondrocytes.	[162]

Also, enhanced chondrocyte proliferation and differentiation

<i>Sulfated alginate</i>	Alginate scaffold	BMP-2	Bone <i>In vitro</i> <i>MC3T3-E1 osteoblast cells</i>	Prolonged release of BMP-2, and enhanced osteogenesis of encapsulated cells.	[45]
<i>Sulfated cellulose</i>	Gelatin scaffold	-	Cartilage <i>In vitro</i> <i>Human mesenchymal stem cells</i>	Enhanced chondrogenesis compared to scaffolds made from pure gelatin.	[172]