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Type, density, and presentation of grafted adhesion peptides on polysaccharide based hydrogels control pre-osteoblast behavior and differentiation

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ABSTRACT:
In this work, cell-responsive polysaccharide hydrogels were prepared by a simple procedure based on the sequential bioconjugation and crosslinking of the polysaccharide backbone with bioactive peptides and poly(ethylene glycol)-bis(thiol) (PEG-(SH)\textsubscript{2}), respectively. Using thiol-ene reactions, we successfully functionalized hyaluronic acid (HA) and carboxymethylcellulose (CMC) with short and long peptides (5-mer and 15-mer derivatives, respectively) derived from adhesive proteins of bone extracellular matrix. The resulting HA-peptide and CMC-peptide
conjugates with varying degrees of substitution were then carefully characterized by $^1$H NMR spectroscopy to precisely control the peptide density into the hydrogels crosslinked with PEG-(SH)$_2$. Pre-osteoblast seeded on the hydrogels with controlled identical stiffness spread in a manner that was strongly dependent on ligand density. Surprisingly, increasing the density of the adhesive peptide anchors did not result in a plateau of initial cell spreading but rather in a bell-shaped cell response which varies with the nature of both polysaccharide backbone and functional peptide. Placing the cells under optimal conditions for cell/hydrogel interaction, we showed that in HA hydrogels, the polysaccharide moiety is not solely a passive scaffold that presents the active peptides but is an active player in cell microenvironment to control and sustain cell activity.

Keywords: hyaluronic acid, hydrogel, thiol-ene, cell-adhesion, osteoblasts

INTRODUCTION

Bio-instructive, -specific, and -responsive hydrogels mimicking the extracellular matrix (ECM) environment have emerged out of the traditional landscape of inert biomaterials.$^{1-3}$ They are often made of natural polysaccharides that act as scaffold for cellular guidance and wound healing while they may also be subjected to biodegradation and can be replaced by bona fide extracellular matrix over time. In addition, their biophysical properties such as compliance can be easily tuned to match the physiological cellular environment and have been of broad use such as in vitro culture and maintenance of stem cells, or direct stem cell specification.$^{4-5}$ Indeed, changes in tissue and organ stiffness are frequently symptoms of diseases such as cancer.$^6$
Hyaluronic acid is a high molar mass (6-7000 kg/mol) linear glycosaminoglycan found in soft tissues such as connective, epithelial and neuronal tissues, and synovial fluid as well. It consists of a repeating disaccharide unit made of N-acetyl-D-glucosamine and D-glucuronic acid residues. This polysaccharide can be obtained with a high degree of purity (pharmaceutical grade) on a large scale by bacterial fermentation which makes it very attractive for designing soft materials in the biomedical field. Various functional groups such as sulfate, (meth)acrylate, alkyl chains but also host molecules, and peptides can be introduced along the polysaccharide backbone in order to control gelation or compliance and trigger cell specific interactions with cell surface receptors. Therefore this polysaccharide has been widely used to produce a variety of tailor made hydrogels. Other hydrogels relying on polysaccharides that are not present in human tissues (carboxymethylcellulose, chitosan, alginates…), may be chosen for their longer resistance to in vivo environment while keeping their biocompatibility.

The cell surface receptors integrins are viewed as major players of cell adhesion to the extracellular matrix and transmit signals to cells upon occupancy with extracellular ligands as well as acting as mechano-transduction sensors. Their roles have been emphasized in many biological processes such as development, immune response, migration, and hemostasis. In addition, on mammalian cells, four major classes of HA cell receptors have been described: CD44 standard or variants, their co-receptor RHAMM, layilin, and ICAM-1. So far, little is known about the role of HA receptors in cell signaling as well as their putative crosstalk with other well-established partners of cell/ECM interactions such as integrins, although recent reports have described some interplay between HA and integrin substrate in controlling cell behavior. On the other hand, many cell types can attach and spread onto HA hydrogels in a stiffness-dependent manner. This indicates that beside integrins, HA receptors may act as mechano-sensors as well. However this adhesion is cell type dependent and is weaker than integrin
mediated adhesion. This is in contrast to non-mammalian polysaccharide-based gels which are generally resistant to cell-adhesion. Such hydrogel matrices are often used to provide a blank canvas in which peptides containing consensus sequences recognized by integrins such as the Arg-Gly-Asp (RGD) sequence are immobilized, thereby providing a defined chemical environment to regulate cell functions. Naturally derived alginate is an example of such hydrogel in which ligand density was found to be a critical regulator of adhesion, spreading, proliferation, and differentiation of pre-osteoblasts. Yet, for properly probing cell responses to these soft matrices, polysaccharide hydrogels with independently tunable stiffness and biochemical ligand density must be carefully designed. We have previously shown that ene-functional polysaccharides esterified with pentenoic anhydride can be rapidly cross-linked with di-functional thiols via thiol-ene photochemistry under physiological conditions. Cell-responsive polysaccharide hydrogels were thus prepared by a simple procedure based on the sequential bioconjugation and crosslinking of the polysaccharide backbone with a bioactive peptide and poly(ethylene glycol)-bis(thiol), respectively. By varying the initial reactant stoichiometry as well as the nature of the biomolecules, this strategy offers enormous possibilities in the design of reproducible, highly tunable gels of which the cell-adhesive and mechanical properties are independently controlled.

In this work, we used this procedure to investigate cell adhesion, spreading, and proliferation in short term culture as well as differentiation in long term culture of pre-osteoblasts on hydrogels depending on their polysaccharide component (hyaluronic acid vs carboxymethylcellulose) as well as on peptide type and density. The peptides used are derived from adhesive proteins present in the ECM of bone, namely fibronectin (Fn) and bone sialoprotein (bsp). They consist of GRGDS (a fragment from Fn) and Ac-CGGNGEPRGDTYRAY-NH₂ (a 15-mer derived from bone sialoprotein, and here denoted as bsp-RGD). These two RGD peptides were selected
because they have shown promise for use in orthopedic tissue engineering applications by improving the adhesion and function of a number of osteoblastic cell lines.\textsuperscript{30-31} While several HA-based hydrogels with tunable stiffness and RGD-ligand density have been reported in the literature,\textsuperscript{32-36} the potential regulatory role of presentation of adhesion peptides in terms of polysaccharide backbone, type and density on 2D cell culture has never been examined in detail. Herein, we demonstrated ability to form HA-based gels that either permit or inhibit cell spreading according to the ligand density at fixed stiffness. Moreover, by comparing cell responses to HA-based gels with those to CMC-gels, at optimal peptide density allowing to support efficient spreading of pre-osteoblasts, HA-peptide hydrogels versus CMC peptide hydrogels allowed a better differentiation of pre-osteoblasts into osteoblasts (using increased mineralization capabilities as a read-out). Therefore in addition to peptide-induced integrin signaling, the HA backbone triggers an additional signal that was required for differentiation.

MATERIALS AND METHODS

Materials. Bacterial hyaluronic acid under the sodium salt form ($M_w = 120$ kg/mol) was purchased from HTL (Javené, France). The sample of carboxymethylcellulose (CMC, marketed under the name Blanose®, grade 7LF PH, DS = 0.7, $M_w = 70$ kg/mol) was kindly provided by Hercules (France). The molecular weight distribution and the weight-average molecular weight of the HA and CMC samples were determined by size exclusion chromatography using a Waters GPC Alliance chromatograph (USA) equipped with a differential refractometer and a light scattering detector (MALLS) from Wyatt (USA); the solutions were injected at a concentration of 1 mg/mL in 0.1 M NaNO$_3$. The polydispersity index of the samples is $M_w/M_n \sim 1.5-2$. Pentenoic anhydride, $N,N$-dimethylformamide (DMF), phosphate buffer saline (PBS, pH ~ 7.4), 2-$(N$-morpholino)ethanesulfonic acid (MES), (3-aminopropyl)trimethoxysilane (APTS), $N$-ethyl-$N'$-((3-
dimethylaminopropyl)carbodiimide hydrochloride (EDC), N-hydroxysulfosuccinimide (sulfo-NHS) were purchased from Sigma-Aldrich-Fluka (L'isled'Abeau, France). 2-hydroxy-1-[4-(2-hydroxyethoxy)phenyl]-2-methyl-1-propanone (Irgacure 2959) was kindly provided by Ciba Specialty Chemicals (Basel, Switzerland). All chemicals were used without any further purification.

The peptides GRGDS, GRGDE, Ac-CGGNGEPGRGDTYRAY-NH$_2$ and Ac-CGGNGEPGETYRAY-NH$_2$ with >85% purity (from manufacturer HPLC analysis), were obtained from GeneCust Europe (Luxembourg). Poly(ethylene glycol)-bis(thiol) (M$_n$ = 3400 g/mol) was purchased from PEGWorks (USA). All chemicals were used without any further purification. The water used in all experiments was purified by aElgaPurelab purification system, with a resistivity of 18.2 MΩ·cm.

**Synthesis of pentenoate-modified HA (HA-p).** HA (0.30 g, 0.75 mmol) was dissolved in ultrapure water (15 mL) at 4 °C and the resulting mixture was kept at 4 °C under continuous stirring overnight for complete dissolution. DMF (10 mL) was then added dropwise in order to have a water/DMF ratio of (3:2, v/v). Pentenoic anhydride (0.411 g, 2.25 mmol) was added while maintaining the pH between 8 and 9 (by adding 0.5 M NaOH) for 4 hours. The reaction was kept at 4 °C under continuous stirring for one night. After this time, NaCl was added to the reaction mixture to have a NaCl concentration of 0.5 M. The polymer was precipitated by addition of ethanol (water/ EtOH (v/v) ratio of 2:3). After removal of the supernatant, the precipitate was successively washed with mixtures of water/EtOH (3:7, 1:4, v/v) and finally dissolved in ultrapure water for a final purification by diafiltration (ultramembraneAmicon YM10) with ultrapure water. The purified product was recovered by freeze-drying and characterized by $^1$H NMR spectroscopy. The degree of substitution (DS, the average number of moles of substituents per repeating unit of the polysaccharide) was determined to be 0.50 ± 0.05.
Synthesis of pentenoate-modified CMC. CMC (0.30 g, 1.34 mmol) was dissolved in ultrapure water (15 mL) at 4 °C and the resulting mixture was kept at 4 °C under continuous stirring overnight for complete dissolution. DMF (10 mL) was then added dropwise in order to have a water/DMF ratio of (3:2, v/v). Pentenoic anhydride (0.980 g, 5.36 mmol) was added while maintaining the pH between 8 and 9 (by adding 0.5 M NaOH) for 4 hours. The reaction was kept at 4 °C under continuous stirring for one night. After this time, NaCl was added to the reaction mixture to have a NaCl concentration of 0.5 M. The polymer was precipitated by addition of ethanol (water/ EtOH (v/v) ratio of 2:3). After removal of the supernatant, the precipitate was successively washed with mixtures of water/EtOH (3:7, 1:4, v/v) and finally dissolved in ultrapure water for a final purification by diafiltration (ultramembraneAmicon YM10) with ultrapure water. The purified product was recovered by freeze-drying and characterized by $^1$H NMR spectroscopy. The degree of substitution was determined to be 0.20 ± 0.02.

Synthesis of HA-peptide and CMC-peptide conjugates.

Pentenoate-modified HA or CMC (0.100 g) was dissolved in ultrapure water (10 mL). 50 µL of an aqueous solution of Irgacure 2959 (10 mg/mL) was then added to the polysaccharide solution to obtain a final photoinitiator concentration of 5 % (w/v), followed by thepeptide containing a thiol function. The amount of the peptide was adapted to the desired degree of substitution (see Table 1). The mixture was exposed to light with a UV intensity of 20 mW/cm$^2$ for 5 min under stirring. The product was purified by diafiltration (ultramembraneAmicon YM10) with ultrapure water and was recovered by freeze-drying. The degree of substitution of the polysaccharide-peptide conjugates was determined by $^1$H NMR.

NMR spectroscopy
The $^1$H and $^{13}$C NMR spectra of the polysaccharide derivatives dissolved in deuterium oxide were performed at 25 or 80 °C (depending of the viscosity of the solution) using a Bruker AVANCE III HD spectrometer operating at 400 MHz ($^1$H) and at 100 MHz ($^{13}$C). All $^1$H NMR spectra were recorded by applying a 45° tip angle for the excitation pulse, and a 10 s recycle delay for accurate integration of the proton signals. The $^{13}$C NMR spectra were recorded by applying a 45° tip angle for the excitation pulse, and a 3 s recycle delay. All 2D experiments were acquired using 2K data points and 256 time increments. Chemical shifts are given relative to external tetramethylsilane (TMS = 0 ppm) and calibration was performed using the signal of the residual protons or carbons of the solvent as a secondary reference. Deuterium oxide was obtained from SDS (Vitry, France). Details concerning experimental conditions are given in the figure captions.

Photorheometry.

An AR2000Ex rheometer (TA Instruments Inc.) fitted with a UV-curing cell ($\lambda = 365$ nm) and an aluminum plate (diameter 19 mm) was used for the in situ measurement of the viscoelastic properties of the HA- and CMC-based hydrogels. Following deposition of 250 µL of a mixture of HA-peptide conjugate (or CMC-peptide conjugate), PEG-(SH)$_2$ and photoinitiator, the gap between the flat quartz plate and the aluminum plate was initially 0.7 mm (measuring ambient temperature). It was controlled during the experiments by maintaining the normal force at $0 \pm 0.1$ N. On each hydrogel, oscillatory time sweep and frequency sweep experiments were performed. They were carried out at 25 °C, with a film of silicone to avoid solvent evaporation. All the dynamic rheological data were checked as a function of strain amplitude to ensure that the measurements were performed in the linear viscoelastic region. In the oscillatory time sweep experiments, the storage modulus ($G'$) and loss modulus ($G''$) were measured during a period of
25-30 min at a fixed frequency of 1 Hz and a fixed deformation of 3.5 %. Typically, after
deposition of the solution of the mixture of polymers in PBS between the plates and equilibration
for 1 min, the solution was illuminated (λ = 365 nm) for 25-30 min at a fixed light power (20
mW/cm²) leading to gelation. All measurements were done in triplicate. The photoinduced
crosslinking reaction was performed by adjusting the PEG-(SH)₂ amount with respect to HA and
CMC to obtain hydrogels having a similar elastic modulus (G’) of 15000 ± 1500 Pa. The
thiol:CMC molar ratio used for gel formation was approximately two-times lower than the
thiol:HA molar ratio as the repeating unit of CMC is a monosaccharide whereas that of HA is a
disaccharide (see next section).

Hydrogel immobilization

For the cell culture experiments, hydrogels were covalently linked to the coverslips during photo-
crosslinking reactions by grafting pentenoate-modified HA or CMC on the coverslip prior to use.
The coverslip was modified with APTS as previously described. The modified slip with –NH₂
groups was immersed in 2 mL of a solution of pentenoate-modified HA or CMC (3 mg/mL in 0.2
M MES, pH 4.75). A solution of sulfo-NHS (30 g/L, 100 µL) and a solution of EDC (50 g/L, 100
µL) were subsequently added. The coverslip was gently stirred at room temperature for 4 h. The
slip was then rinsed with water and dried under a flow of nitrogen before use. The HA- and
CMC-based hydrogels were prepared by deposition of a solution of HA or CMC modified with
pentenoate groups and peptides (75 µL, [HA] = 30 g/L and [CMC] = 30 g/L) containing Irgacure
2959 (5 % w/v) and PEG-(SH)₂ (2.1 mg (0.61 × 10⁻³ mmol) for HA and 0.9 mg (0.26 × 10⁻³ mmol
for CMC), [SH]/[pentenoate] = 0.5 for HA and for CMC) to the glass slip. The mixture was then
coated with a glass slide using 0.4 mm-thick glass plate spacers and then exposed to UV-
irradiation (20 mW/cm², 20 min). Under such conditions, hydrogel disks with and without cell-
adhesive peptides ($G' = 15000 \pm 1500 \text{ Pa}$) were prepared. After preparation, hydrogels were transferred to cell culture media (Dulbecco's modified minimal essential medium, DMEM, Invitrogen) supplemented with 50 U/mL penicillin, and 50 mg/mL streptomycin and left for 12 h for equilibrium before cell culture studies. Media was refreshed after 12 h to remove any remaining monomer or initiator.

**Generation of preosteoblast cell line.**

To generate preosteoblast cell lines, primary osteoblasts (passage 2) were immortalized by infection with adenovirus expressing the large SV40 T antigen. Immortalized cells were cloned and each clone tested for its ability to differentiate as viewed alkaline phosphatase expression and aptitude to mineralize ECM, as previously described. They were cultivated under the standard conditions in DMEM supplemented in 10% fetal calf serum in a 5% CO$_2$ atmosphere.

**Cell labeling and quantification of cell spreading**

For quantifying the projected areas of spread cells, the cells were fixed with a solution of 4% (w/v) of paraformaldehyde in 0.2 M phosphate buffer (pH= 7.2) for 10 min at room temperature. After 3 washes in TRIS saline buffer (TBS), the cells were incubated in DMEM supplemented with 0.5% of Vybrant™ DIL cell labeling solution for 15 min at 37°C according to the manufacturer's instructions (Life Technologies, St Aubin, France). The cells were washed 3 times in PBS and mounted under a coverslip with 10 µL of mounting solution for epifluorescence observations. Observation were carried out with an Olympus BX 51 microscope (Olympus Europe Hamburg, Germany) equipped with a Plan NeoFluar 20X (N.A. 0.5) objective. Areas were determined with Metamorph software (Molecular Devices, Sunyvale, CA USA) at a magnification of 200X. Statistical analyses were carried out with R software (The R Project for Statistical Computing, [http://www.r-project.org/](http://www.r-project.org/)).
EdU cell proliferation assay

DNA synthesis detection in proliferating cells was based on the incorporation of 5-ethynyl-2-deoxyuridine (EdU) and its subsequent detection by a fluorescent azide through a Cu(I)-catalyzed [3 + 2] cycloaddition reaction.\(^{41}\) Cells were grown on hydrogel substrates for 24 h in DMEM supplemented with 10% fetal calf serum, penicillin, and streptomycin. EdU was added to the culture media at 10 μM (Stock 10mM), for 1-2 hour. After labeling, cells were washed with PBS. Cells were fixed by using a standard paraformaldehyde 4% for 10 min. Subsequently, cells were rinsed once with TBS and stained by incubating for 30 min with 0.5-1mL/well with 6 mL of afreshly prepared staining mix made of 110 mMTris pH 8.5, 1 mM CuSO\(_4\), 60mM ascorbic acid, 10μM Alexa azido (from stock solution in DMSO). After staining, the cells were washed three times with TBS with 0.2% Triton X-100. The fraction of fluorescent (dividing) cells was estimated on at least 10 independent frames.

Alizarin Red-S Protocol for Mineralization

Pre-osteoblasts 80-90% confluent were placed into osteogenic media made of regular DMEM medium with 10% FetalCloneII (Hyclone), supplemented with 10mM\(\beta\)-glycerophosphate, and 100μM of ascorbic acid. Culture was continued for 21 days. Mineralization was already observed at 10 days. The calcified mineral deposits were stained with Alizarin Red-S (AR-S) using the protocol developed by Gregory et al.\(^{42}\) Briefly, separate 12-well plates after 8, 11, 15 and 24 days in culture were washed gently with PBS and fixed in 10 % (v/v) formaldehyde for 15 min at room temperature. After washing twice with excess deionized water, the samples were immersed in 0.5 mL of 40 mM AR-S (pH 4.1) for 20 min with gentle shaking. Then, the samples were washed four times with excess deionized water for 5 min each with shaking. The wet samples were subsequently imaged using a stereo microscope Olympus SZX10v with an angled
illumination source (Schott K1500). For quantification, threshold of the staining on digital images allowed the measurement of mineralized areas using Metamorph software.

RESULTS

Synthesis of HA-peptide and CMC-peptide conjugates. To fabricate HA and CMC hydrogels (Figure 1), we first functionalized HA and CMC with pentenoate groups by reaction of the hydroxyl groups of the polysaccharides with 4-pentenoic anhydride as previously described. In these reactions, the feed molar ratios of anhydride to the polysaccharide repeat unit were adjusted in a way to obtain HA-pentenoate (HA-p) and CMC-pentenoate (CMC-p) derivatives having a degree of substitution (DS) of 0.50 and 0.20, respectively (see experimental section). The DS was approximately two-times lower in the case of CMC because the repeating unit of this polysaccharide is only composed of one sugar.

Then, the resulting HA-p and CMC-p derivatives were reacted with the 5-mer peptide GRGDS and the 15-mer peptide bsp-RGD via photoinduced thiol-ene reactions between the thiol of the terminal mercaptopropionic acid in GRGDS and the terminal cysteine in bsp-RGD and, the alkene function of pentenoate. In order to tune the cell-adhesive ligand density while maintaining a constant elastic modulus, the peptides were added in equivalents of no greater than 50% of the total number of alkene groups of HA and CMC. In this way, the double bonds after peptide conjugation are in sufficient numbers to maintain the same crosslinking density while increasing the peptide density. Integrin specific cell response was controlled by grafting non-adhesion peptides containing the RGE sequence instead of RGD: GRGES (Gly-Arg-Gly-Glu-Ser) and bsp-RGE (Ac-Cys-Gly-Gly-Asn-Gly-Glu-Pro-Arg-Gly-Glu-Thr-Tyr-Arg-Ala-Tyr-NH2). Following this coupling step, the HA-peptide and CMC-peptide conjugates were purified by diafiltration and analyzed by 1H NMR to check their structural integrity and precisely determine their DS (see
Table 1 and $^1$H NMR spectra Supporting Information (Figures S1-S10)). The DS was calculated by digital integration of well-defined signals of the peptide and of the polysaccharides (Figures S3, S6, S8, S10). Of note, the DS values could be confirmed by analyzing the disappearance of the signals of the protons associated with the double bonds (Figures S3, S6, S8, S10). Comparison between the feed molar ratios of peptide to pentenoate and the DS of the modified polysaccharides (Table 1) indicated that the peptides were grafted on HA and on CMC in quite satisfactory yields (66-91 %). The final crosslinking reaction step was performed by adjusting the PEG-(SH)$_2$ amount at a fixed amount of HA or CMC. The HA- and CMC-based hydrogels were prepared using, respectively, a thiol:HA disaccharide repeating unit molar ratio of 0.25 and a thiol:CMC monosaccharide repeating unit molar ratio of 0.10 in order to obtain hydrogels having a similar elastic modulus (G’) of 15000 ±1500 Pa. This value was chosen based on our previous results showing the ability of pre-osteoblasts to attach and partially spread on a HA-based hydrogel having an elastic modulus of 17000 Pa.

**Pre-osteoblast spreading.** Preliminary biological studies were performed to investigate the effect of the nature of the peptide ligand and its density as well as of the polysaccharide backbone on cell spreading at constant stiffness. To this end, pre-osteoblasts were seeded on the hydrogels prepared from the HA and CMC derivatives listed in Table 1 as well as from HA-p and CMC-p, and cultured overnight in DMEM supplemented with 10% fetal calf serum. Cell spreading was analyzed by statistical measurements of the cell projected area using Vybrant™ DIL staining and Metamorph image analysis software (see experimental section).

Figure 2A shows that the HA hydrogel lacking peptides was able to sustain some but limited cell spreading (in good agreement with previous reports$^{25}$) while under identical conditions, pre-osteoblasts seeded on the gel derived from CMC-p remained rounded up. In contrast, significant spreading on CMC and HA gels was obtained by incorporation of the GRGDS and bsp-RGD
peptides. A marked difference in cell spreading efficiency could be observed for the two peptides
with hydrogels based on CMC hydrogels (Figures 2A and 2B); The bsp-RGD-endowed gels
sustained cell spreading in a more effective way. Surprisingly, for all hydrogels bell shaped
adhesion curves were obtained depending on peptide densities, and optimal peptide concentration
varied according to the polysaccharide backbone used and the peptide sequence. For instance,
optimal GRGD peptide density was higher than the bsp-RGD one to induce the same degree of
spreading (Figure 2B). Since the control GRGES or bsp-RGE peptides promote little or
nosignificant spreading when incorporated on HA and CMC, respectively, similarly to non grafted
gels (Figure 2B), one can conclude that the observed pre-osteoblast responses are
mostly dependent on the grafted peptide and likely due to integrin surface receptors.
The decrease in cell binding at high ligand concentrations may result from a negative effect of
neighboring ligands on binding as previously reported. Assuming that the cells can only access
the top 10 nm of the hydrogel, the peptide surface concentrations were found to be between 0.7
and 12.2 pmol/cm² (tableau 1). Rezania et al. determined the minimum concentration of bsp-
RGD required for spreading of rat-derived calvarial osteoblasts on quartz surfaces functionalized
with bsp-RGD to be ~ 0.6 pmol/cm². Although the actual minimum concentration of bsp-RGD
ligands required at the HA and CMC-based gel surfaces for spreading of pre-osteoblasts was not
identified, one can however note that the optimal concentration required for maximum spreading
on the bsp-RGD endowed HA- and CMC hydrogels is 4.5 fold the limit literature value.
All together, these results highlight that by exploiting the design flexibility of the characteristics
offered by these biomimetic hydrogels, it may be possible to selectively engineer their biological
properties. The gels based on HA-11RGD and on HA-4bsp-RGD appeared to be the most efficient
substrates for promoting cell spreading after one night (marked in yellow in Figure 2B). The
CMC-based substrates were found to be effective in promoting cell spreading only if they
contained bsp-RGD ligands. Since the optimal cell/substrate interaction depends not only on both
the nature of the peptide used for functionalization and its surface concentration, but also on the
polysaccharide backbone, these experimental conditions must clearly be determined when
comparing the biological properties of hydrogels.

**Cell response proliferation and differentiation during long term culture on hydrogels.** To
better understand how the types of biomimetic modification and polysaccharide matrix might be
used for best biocompatibility, proliferation assays where carried out after a 24 h pre-culture and
measured by EdU staining of dividing cells (see experimental section). The results presented in
Figure 3, clearly indicated that at the optimal DS allowing to support efficient cell spreading,
both RGD and bsp-RGD triggered a high proliferation rate, and that RGD-grafted HA gels
allowed a better cell proliferation rate that the CMC counterpart. Analysis of matrix
mineralization in long term culture (a read-out of osteogenic differentiation) was carried out with
hydrogels that support optimized cell spreading, namely gel substrates based on HA-11RGD,
HA-4bsp-RGD and CMC-4bsp-RGD for 21 days. Observing surface-bound pre-osteoblasts
during this period, it was found that cells not bound to surfaces died after a few days, the others
were proliferating and formed a monolayer before stopping dividing and initializing
differentiation.

At different time points ECM mineralization was examined on representative cell cultures by
staining calcium deposition using Alizarin Red S (Figure 4). Significantly more mineral
deposition (red to dark brown staining depending on the amount of calcium) on the gel surfaces
was detected for HA-based gels after 10 and 21 days (Figure 4). This was confirmed by a
statistically analysis of 10 distinct fields for quantification of the areas of positively stained
deposits (Figure 5). Significantly larger areas of mineral deposits on HA-based gels compared to
the CMC-based hydrogel were observed after 10 and 21 days although mineralization nodules
were quite visible on CMC gels at 21 days. Furthermore, comparison between the HA-4bsp-RGD and HA-11RGD substrates indicates a higher degree of mineralization for the gel displaying bsp-RGD ligands. Overall these results indicate that HA-based hydrogels promote osteogenic development when combined with peptide ligands that facilitate integrin-mediated binding matrix mineralization. Considering the significant difference in the optimal DS of the two HA-peptide conjugates, it can clearly be concluded that the bsp-RGD-endowed hydrogel sustained cellular proliferation and mineralization in a more effective way.

DISCUSSION

In this study, we synthesized and characterized a new class of HA- and CMC-based hydrogels with independently tunable stiffness and adhesive peptide functionality and used these soft hydrogels to study the dependence of spreading and long-term function of pre-osteoblast cells on changes in matrix chemical composition at fixed gel stiffness. Taking advantage of the flexibility of thiol-ene reactions, we showed the ability to incorporate short peptides (5-mer derivatives) as well as long peptides (15-mer derivatives) in nearly 100% yields and precisely quantify their incorporation into hydrogels by a careful characterization of peptide-modified polysaccharides by $^1$H NMR spectroscopy. Pre-osteoblast seeded on the hydrogels spread in a manner that was strongly dependent on ligand density. Surprisingly, increasing the density of the adhesive peptide anchors did not result in a plateau of initial cell spreading but rather in a bell shaped cell response which varies with the nature of both polysaccharide backbone and functional peptide. When bsp-RGD was incorporated into HA and CMC gels instead of GRGDS, the optimal average peptide density for cell spreading was shifted to lower values, which can be attributed either to a better accessibility of the bsp-RGD possibly due the increased chain length, or to a better integrin signaling or/and clustering. It is therefore very important to determine key matrix design
relationships such as the effects of presentation of adhesion peptides in terms of type and density, nature of polysaccharide backbone used for gel formation and crosslink density leading to optimal cell response, before drawing conclusions on the biocompatibility of a specific hydrogel.

We also demonstrated that for pre-osteoblasts of which adhesion to fibronectin is mediated by significant interactions with both the α5β1 and αv integrins, hyaluronic acid synergizes with integrin occupancy not only to promote cell spreading under short term culture, but also osteoblast differentiation and mineralization under long term culture. This specific cell response to RGD-endowed HA gels suggests that there is an interplay between HA receptors and RGD-specific integrins (i.e. α5β1 and αvβ3). Although several investigators have previously reported on the development of HA-based hydrogels with tunable stiffness and ligand density, the likely involvement of HA receptors namely CD44 and RHAMM along with RGD-binding integrins has only recently been reported. Our work clearly shows that in HA hydrogels, the polysaccharide moiety is not solely a passive scaffold that presents the active peptides but is an active player in cell microenvironment to control and sustain cell activity.

In this regard, our strategy described here for synthesizing well-defined hydrogels that combine the intrinsic properties of mammalian polysaccharides and cell adhesive ligands isolated from extracellular matrix proteins, provides a versatile platform to get a better insight into the molecular mechanisms that underlie cell-specific responses to naturally-derived hydrogels with respect to key chemical and mechanical attributes.

CONCLUSIONS

The strategy for the synthesis of bioactive polysaccharide hydrogels described in this work represents a versatile platform to evaluate the effect of peptides, their density and
the polysaccharide matrix on cellular adhesion and function with minimal contribution from non-specific interactions. Using these substrates, the residues flanking the RGD sequence in peptides was shown to significantly influence the spreading and long-term function of pre-osteoblast cells. Future studies will focus on engineering gels with controlled porosity to promote cell infiltration into gels as a bioactive scaffold for osteogenesis. In addition, naturally-derived hydrogels with controlled biological interactions conferred by tissue-specific ligands may be applied to the engineering of a variety of other tissue types.

ACKNOWLEDGMENTS

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Supporting Information Available:

\(^1\)H NMR spectra of GRGDS, GRGDE, HA-RGD with different DS, two-dimensional \(^1\)H,\(^{13}\)C-HSQC spectrum of bsp-RGD, HA-bsp-RGD with different DS, CMC-RGD with different DS, CMC-bsp-RGD with different DS. This material is available free of charge via the Internet at http://pubs.acs.org.
**Figure 1.** Schematic representation of the synthesis of polysaccharide-peptide conjugates and their cross-linking to form hydrogels using thiol-ene photochemistry.

**Figure 2.** Pre-osteoblast attachment and spreading on HA- and CMC-based hydrogels as a function of presentation of adhesion peptides in terms of type and density. A) phase contrast observations carried out on living cells with a Zeiss 100M Axiovert microscope equipped with a plan 10X objective (N.A 0.25). Bar is 60 µm. B) cell spreading quantification. After cell fixation with paraformaldehyde, cells were labeled with Vybrant™ DIL lipophilic tracer. Observations of the fluorescent cells were carried out with an Olympus BX51 epifluorescent microscope equipped with a Plan NeoFluar 20X objective (N.A. 0.50). The images threshold value was adjusted to fit the cells limits and the projected cell areas were determined with Metamorph software. Statistical analyses and box plots were performed with the R software (n=60). The bottom and top of the box are the 25th and 75th percentile (the lower and upper quartiles, respectively), and the band near the middle of the box is the 50th percentile (median). The ends of the whiskers represent the lowest datum still within 1.5 Inter Quantile Range of the lower quartile, and the highest datum still within 1.5 Inter Quantile Range of the upper quartile. The DS of the HA-peptide and CMC-peptide conjugates is indicated above the boxes in red color. The DS values giving rise to the highest degree of spreading are marked in yellow.

**Figure 3.** Cell proliferation on peptide-modified HA and CMC hydrogels with DS promoting cell spreading: HA-11RGD, HA-4bsp-RGD and CMC-4bsp-RGD (see Figure 2) after 24 h of pre-culture. Error bar represents mean ± standard deviation. The analysis of 10 independent fields.
under each experimental conditions represented the analysis of more than 100 cells. Data on two experiments were pooled to allow student tests.

Figure 4. Digital photographs of representative Alizarin Red-S stained cell monolayers on peptide-modified hydrogels after 0, 10, and 21 days in culture in the osteogenic medium. The red to dark brown areas represent the calcified deposits. Color intensity reflects the amount of hydroxyapatite deposition. After 21 days the HA-based hydrogels showed higher degrees of mineralization than the CMC-based hydrogel. Bar is 200 µm.

Figure 5. Area quantification of the calcified ECM after 0, 10, and 21 days in culture on 10 independent fields. The HA-based hydrogels displayed significantly greater mineralization than the CMC-hydrogel on days 10 and 21. Error bar represents mean ± standard deviation.
### Table 1. List of peptide sequences and of polysaccharide (PS)-peptide conjugates prepared by thiol-ene chemistry

<table>
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<tr>
<th>PS</th>
<th>Peptide/Peptide sequence</th>
<th>[peptide]/[PS repeat unit]</th>
<th>DS (% peptide)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Name</th>
<th>Coupling yield&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Peptide surface density&lt;sup&gt;c&lt;/sup&gt;</th>
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<tr>
<td>HA-p</td>
<td>GRGDS</td>
<td>0.04</td>
<td>3</td>
<td>HA-3RGD</td>
<td>75</td>
<td>2.0</td>
</tr>
<tr>
<td>HS-CH₂CH₂CO-NH-GRGDS</td>
<td></td>
<td>0.07</td>
<td>6</td>
<td>HA-6RGD</td>
<td>85</td>
<td>4.1</td>
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<tr>
<td></td>
<td></td>
<td>0.12</td>
<td>11</td>
<td>HA-11RGD</td>
<td>91</td>
<td>7.5</td>
</tr>
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<td></td>
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<td>0.20</td>
<td>18</td>
<td>HA-18RGD</td>
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<td>12.2</td>
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<td>bsp-RGD</td>
<td>0.015</td>
<td>1</td>
<td>HA-1bsp-RGD</td>
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<td>0.7</td>
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<tr>
<td></td>
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<td>2</td>
<td>HA-2bsp-RGD</td>
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<td>CMC-3bsp-RGE</td>
<td>60</td>
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<sup>a</sup>determined from <sup>1</sup>H NMR spectroscopy; <sup>b</sup>calculated by dividing (0.01× DS) by the feed ratio [peptide]/[PS repeat unit];<sup>c</sup>calculated by assuming that the cells can only access the top 10 nm of the hydrogel.
Figure 1

HA-p or CMC-p

Photoinitiator
Water, RT

≡

SH

SA-p or CMC-p

Photoinitiator
Water, RT

≡

SH

SH

SH

SH

Photoinitiator
Water, RT

≡

SH

HA-p

HS-CH₂-CH₂-CO-NH-GRGDS (denoted as “RGD”)

or

Ac-CGGNGEPGRDTRAY (denoted as “bsp-RGD”)

or

HS-CH₂-CH₂-CO-NH-GRGES (denoted as “RGE”)

or

Ac-CGGNGEPGRGETRAY (denoted as “bsp-RGE”)

Pre-osteoblasts
Figure 2
Figure 3

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Edu Positive Nuclei (%)</th>
<th>p</th>
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<tbody>
<tr>
<td>HA-RGD</td>
<td>60 ± 5</td>
<td></td>
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<tr>
<td>CMC-RGD</td>
<td>40 ± 4</td>
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<tr>
<td>HA-BSP</td>
<td>70 ± 6</td>
<td></td>
</tr>
</tbody>
</table>

- p = 0.0040
- p = 0.0001
- n.s.
Figure 4

Day 0                      Day 10                      Day 21

HA-11GRGDS

HA-4bsp-RGD

CMC-4bsp-RGD
Figure 5

The graph shows the percentage of mineralized area over different days for three types of materials: HA-11RGD, HA-4bsp-RGD, and CMC-4bsp-RGD. The data is presented at Day 0, Day 10, and Day 21.
**Figure S1.** $^1$H NMR spectra (400 MHz, 3 mg/mL in D$_2$O, 25 °C) of GRGDS (a) and GRGDE (b).
Figure S2. $^1$H NMR spectra (400 MHz, 6 mg/mL in D$_2$O, 80 °C) of HA-18RGD (blue curve), HA-11RGD (pink curve), HA-6RGD (black curve) and HA-3RGD (green curve).
**Figure S3.** Complete and partial contour plots of a $^1$H,$^{13}$C-HSQC experiment performed on bsp-RGD (4 mg/mL in D$_2$O, 25 °C). The $^1$H and $^{13}$C DEPT135 NMR projection spectra of the peptide are shown along the horizontal and vertical axes, respectively.

**Figure S4.** $^1$H NMR spectra (400 MHz, 6 mg/mL in D$_2$O, 80 °C) of HA-12bsp-RGD (black curve), HA-7bsp-RGD (green curve), HA-4bsp-RGD (pink curve) and HA-2bsp-RGD (blue curve).
**Figure S5.** $^1$H NMR spectra (400 MHz, 6 mg/mL in D$_2$O, 80 °C) of pentenoate-modified CMC (black curve), CMC-6RGD (green curve) and CMC-3RGD (pink curve).
Figure S5. $^1$H NMR spectra (400 MHz, 6 mg/mL in D$_2$O, 80 °C) of pentenoate-modified CMC (green curve), CMC-9bsp-RGD (black curve) and CMC-4bsp-RGD (pink curve).
Figure S6. $^1$H NMR spectra (400 MHz, 6 mg/mL in D$_2$O, 80 °C) of HA-12bsp-RGD (black curve), HA-7bsp-RGD (green curve), HA-4bsp-RGD (pink curve) and HA-2bsp-RGD (blue curve).