Tyramine-conjugated alginate hydrogels as a platform for bioactive scaffolds

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Abstract: Alginate-based hydrogels represent promising microenvironments for cell culture and tissue engineering, as their mechanical and porous characteristics are adjustable toward in vivo conditions. However, alginate scaffolds are biocer and thus inhibit cellular interactions. To overcome this disadvantage, bioactive alginate surfaces were produced by conjugating tyramine molecules to high-molecular-weight alginate using the carbodiimide chemistry. Structural elucidation using nuclear magnetic resonance spectroscopy and contact angle measurements revealed a surface chemistry and wettability of tyramine-alginate hydrogels similar to standard cell culture treated polystyrene. In contrast to stiff cell culture plastic, tyramine-alginate scaffolds were found to be soft (60–80 kPa), meeting the elastic moduli of human tissues such as liver and heart. We further demonstrated an enhanced protein adsorption with increasing tyramine conjugation, stable for several weeks. Cell culture studies with human mesenchymal stem cells and human pluripotent stem cell-derived cardiomyocytes qualified tyramine-alginate hydrogels as bioactive platforms enabling cell adhesion and contraction on (structured) 2-D layer and spherical matrices. Due to the alginate functionalization with tyramines, stable cell–matrix interactions were observed beneficial for an implementation in biology, biotechnology, and medicine toward efficient cell culture and tissue substitutes. © 2018 The Authors. Journal of Biomedical Materials Research Part A published by Wiley Periodicals, Inc.. J Biomed Mater Res Part A: 107A: 114–121, 2019.

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INTRODUCTION

Controlling the interface between biomaterials and biological entities such as proteins and cells represents the key for sophisticated cell cultures, implants, and disease models for drug screening. In the last decades, the design of biomaterials was widely investigated to direct cellular behavior.1,2 Besides the surface characteristics of the biomaterial such as its wettability,3 charge,4 chemistry,5 and topography,6 the materials mechanical stability7 and porosity8 were defined as crucial parameters for cell–matrix interactions. The mechanical microenvironment of the cells affects cellular processes like adhesion,9 gene expression,10 and differentiation.11 Nevertheless, standard cell culture processes are still mainly carried out on stiff plastic neglecting the significance of the cell–matrix interface.

To meet the requirements of the complex, native cell environment soft and porous materials with adjustable mechanical properties are needed. Here, alginate as a widely applied biomaterial in tissue engineering12–14 represents a promising cell microenvironment, as this biocompatible polymer is able to form hydrogels with adaptable mechanical characteristics and porosities.15 However, native alginate inhibits cell adhesion due to its repelling anionic surface and lacking cell interaction sites.16 Hence, this study aims to address cell–alginate interactions via the functionalization of the cross-linked alginic scaffold (ALG).

Standard cell culture surfaces are mainly based on polystyrene (PS) which were modified by sulfuric acid treatment and physical methods (corona discharge, gas plasma, or irradiation) to introduce hydroxyl, phenolic, and carboxylic groups to the carbon backbone.17 In comparison, alginate consists of 1,4-glycosidically linked mannuronic and guluronic acids bearing hydroxyl and carboxylic functional groups. Consequently, the surface chemistry of standard cell culture plastic can be simulated via the introduction of phenolic groups to the alginate scaffolds. Tyramine, a biogenic...
amine occurring in plants and animals, owns a phenolic component and a terminal amino group which represents a suitable binding site for the amidation with alginate using the carbodiimide chemistry. In previous studies, tyramine was coupled to a wide range of natural polymers such as hyaluronic acid,\textsuperscript{18} dextran,\textsuperscript{19} and alginate\textsuperscript{20,23} to induce an enzyme-mediated gelation using horseradish peroxidase (HRP) and hydrogen peroxide.\textsuperscript{18–20,22,23} These bioresponsive systems were mainly applied for encapsulations\textsuperscript{10,20,22} and 3D printing\textsuperscript{21} as \textit{in situ}-forming, injectable hydrogels.

In contrast, this study targets the functionalization of ALG with tyramine to enable a stable cell adhesion to soft, bioactive scaffolds applicable for the demands of future tissue engineering approaches in, for example, personalized medicine.

**MATERIALS AND METHODS**

**ALG fabrication**

High-molecular-weight alginites extracted from the brown algae \textit{Lessonia nigrescens} (LN) and \textit{Lessonia trabeculata} (LT) (Alginotec, Riedenheim, Germany) were applied for scaffold fabrication as 0.7% (w/v%) solutions in isotonic, 0.9% sodium chloride solution (NaCl; B. Braun, Melsungen, Germany), and 1:1 mixtures to adjust a defined mannuronate/guluronate ratio. Planar alginate layers were used as scaffolds positioned on round standard cell culture treated polystyrene-based coverslips (Thermanox™, Ø 13 mm, Thermo Fisher Scientific, Dreieich, Germany) and dishes (Ø 35 mm, Corning, New York, NY) treated with poly-L-lysine (pLL; Sigma-Aldrich, Taufkirchen, Germany) in phosphate-buffered saline (PBS; Gibco, Thermo Fisher Scientific, Darmstadt, Germany) for 30 min at 37°C. The alginate solution (LN/LT 1:1, 0.7% [wt %/vol %]) were placed on the treated, dried plastic surfaces and gelled with 20 mM barium chloride solution (BaCl\textsubscript{2}) for 20 min at room temperature (RT). The obtained alginate layers were washed three times with NaCl and stored at 4°C until usage.

Structured, planar alginate surfaces were manufactured using a μ-contact printer (GeSiM mbH, Rossendorf, Germany; see Supporting Information, Fig. 1). The alginate solution was placed on an agarose gel (4% [wt/vol %]; Sigma-Aldrich, Taufkirchen, Germany) containing 100 mM BaCl\textsubscript{2} and structured for 3 min by a stamp based on polydimethylsiloxane (PDMS; Sylgard 184 elastomer kit, Sigma-Aldrich, Taufkirchen, Germany). The structured hydrogel was transferred to a pLL-treated plastic dish, post-cross-linked with 20 mM BaCl\textsubscript{2} for 10 min, washed three times with NaCl, and stored at 4°C until usage.

**Spherical ALG.** In terms of spherical scaffolds, alginate was processed as described previously.\textsuperscript{24} Briefly, the alginate solution was dispersed into droplets using a coaxial air stream and polymerized while dropping into the aqueous 20 mM BaCl\textsubscript{2}. After 20 min of cross-linking, spherical ALG were washed three times with NaCl and stored at 4°C until usage.

**ALG modification**

To introduce bioactive entities, alginate’s carboxylic groups were activated by aqueous carbodiimide chemistry resulting in the conjugation of \textit{N}-hydroxysuccinimide (NHS; Sigma-Aldrich, Taufkirchen, Germany). The obtained alginate-NHS esters were subsequently coupled with tyramine hydrochloride (Sigma-Aldrich, Taufkirchen, Germany) in various quantities (2.5, 7.5, and 12.5 mg/cm\textsuperscript{2} surface area) for 24 h at RT. Matrigel-coated surfaces were produced by the incubation of tyramine-conjugated alginate scaffolds (ALGTYR) with Matrigel (8.7 μg/cm\textsuperscript{2} surface area, Corning, NY) for 15 h at RT. Final modified ALG were washed three times with NaCl.

**Nuclear magnetic resonance (NMR) spectroscopy**

For structural elucidation of the (modified) ALG, NMR spectroscopy was applied. Ultrasonically, freeze-dried alginate samples were dissolved in D\textsubscript{2}O (7.5 mg/mL) containing sodium-3-trimethylsilylpropionate-2,2,3,3-d\textsubscript{4} (TSP) as an internal reference and analyzed with 128 scans and water suppression (presaturation) at 80°C. 1H NMR spectra were recorded using a Bruker DRX 500 (Bruker, MA) and processed (integrations and deconvolutions) using the software TopSpin.

**Wettability analysis using captive bubble technique**

Contact angles were measured using an OCA device with a conventional goniometer contact angle apparatus with an additional bracket for flat samples (SHC 20), a glass cuvette (GC 40), an upward curved dispensing needle (SNC 052/026), a digital image capture, and data analysis software (SCA 20) (all from DataPhysics Instruments, Filterstadt, Germany) at RT. Three seconds after placing an air bubble (3 μL) onto the hydrogel surface, which is surrounded with NaCl, a digital image was captured and analyzed using the SCA 20 software. To measure contact angles after protein adsorption, the (modified) ALG were incubated in DMEM/F-12 (Gibco, Thermo Fisher Scientific, Darmstadt, Germany) supplemented with 10% (v/v%) fetal calf serum (FCS; Sigma-Aldrich, Taufkirchen, Germany) at 37°C for 24 h and were subsequently washed with NaCl before analyzing.

**Zeta potential measurements**

Zeta potentials of (modified) spherical ALG (Ø 50 μm, 10 cm\textsuperscript{2} total surface area) suspended in PBS were measured at 25°C in a capillary cell (Malvern Instruments, Worcestershire, England) using the Zetasizer Nano-ZS PNX702 (Malvern Instruments, Worcestershire, England). Surface charges after protein adsorption were analyzed after the incubation (24 h) of spherical ALG in DMEM/F-12 supplemented with 10% FCS at 37°C and three washing steps with NaCl before analyzing.
**Compression tests**

(Modified) ALG (Ø 13 mm, height: 1.4 mm) surrounded with NaCl were compressed until 35% of strain with a deformation rate of 0.5 mm/s using TAKTplus (Stable Micro Systems, Godalming, United Kingdom). The elastic moduli were calculated as the slope of the stress–strain curve (10–30% of strain).

**Protein adsorption assays**

**Ellman’s reagent.** The sulphydryl groups of adsorbed proteins were determined using 5,5′-dithio-bis-(2-nitrobenzoic acid) (DTNB, Sigma-Aldrich, Taukirchen, Germany). Therefore, the (modified) ALG were incubated in DMEM/F-12 supplemented with 10% FCS at 37°C for 24 h and subsequently washed with PBS. After the addition of 0.1 mM DTNB solution and 2 min of incubation at RT, the absorbance of the test samples was measured at 412 nm with the Infinite F200 microplate reader (Tecan, Maennedorf, Switzerland).

**Immunofluorescence staining.** Spherical ALGTYR were treated with laminin (1.0 μg/cm² surface area; L2020, Sigma Aldrich, Taukirchen, Germany) dissolved in DMEM/F-12 for 4 h at RT and subsequently washed with PBS. Immunostaining against laminin were carried out using a primary antibody against laminin (rabbit polyclonal; Abcam, Cambridge, United Kingdom) and the secondary antibody Alexa Fluor 488 (goat anti-rabbit IgG H + L; Thermo Fisher Scientific, Dreieich, Germany). After washing steps with PBS images were taken immediately as well as after 7 and 14 days of storage at 4°C using a fluorescence microscope (Nikon Eclipse TS100; Nikon Instruments Europe, Amsterdam, The Netherlands).

**Cell culture**

**Human mesenchymal stem cells (MSCs).** Cell culture of MSCs (Wharton’s jelly, PromoCell GmbH, Heidelberg, Germany) was performed in standard cell culture treated flasks (75 cm² growth area; Corning, NY) using DMEM/F-12 supplemented with 10% FCS, 100 units/mL penicillin/streptomycin and 1 ng/mL basic fibroblast growth factor (all from Gibco, Thermo Fisher Scientific, Darmstadt, Germany) as culture media. MSCs were passaged using 0.05% trypsin/EDTA (Gibco, Thermo Fisher Scientific, Darmstadt, Germany) once a week or at confluence of ~80% with a seeding density of at least 1000 cells/cm² for inoculation.

**Human pluripotent stem cell-derived cardiomyocytes (hiPSC-CMs).** hiPSC-CMs (Cor.4U) were cultured using the Cor.4U medium (both from AxioGenesis, Cologne, Germany), which was changed 1–2 times per week before cell experiments. Cell detachment was carried out according to the instructions of the manufacturer.

**Cell adhesion and beat frequency studies**

MSCs (2.5 × 10⁶ cells/scaffold) were seeded onto PS, ALG, and ALGTYR surfaces and the respective adhesion (number adherent cells) was quantified manually after 6 and 12 h of inoculation via live cell imaging (1 image/15 min) using the Biostation IM (Nikon Instruments Europe, Amsterdam, The Netherlands).

hiPSC-CMs (2.5 × 10⁵ cells/scaffold) were seeded onto planar and structured, Matrigel-coated ALGTYR surfaces and cultivated for 21 days. Their adhesion behavior was documented after 1, 3, 7, 14, and 21 days using the microscope Eclipse TE 2000 (Nikon Instruments Europe, Amsterdam, The Netherlands). To study the beat frequency of adherent cells, hiPSC-CMs (4000 cells/cm²) were cultured for 8 days on spherical ALGTYR coated with different Matrigel quantities (0, 2.2, 4.3, 6.5, and 8.7 μg/cm² surface area). The respective beat frequencies were determined manually analyzing the image sequences.

**Statistical evaluation**

Graphical illustration of data and statistical analyses were performed using OriginPro and IBM SPSS Statistics. Differences between groups were considered significant by p < 0.05 and were evaluated with univariate analyses of variance (ANOVA) with simple contrasts.

**RESULTS**

**Scaffold preparation and characteristics**

Tyramine was conjugated reproducibly to ALG via aqueous carbodiimide chemistry. The modification process was confirmed by the structural elucidation using ¹H NMR spectroscopy [Figs. 1 and 2(A)]. The spectra of ALG illustrate the typical homo- and heteropolymeric block fractions of alginites consisting of mannuronate and guluronate. After the amidation between ALG and tyramine using EDC and NHS, the aromatic protons of tyramines phenolic group were present within the ALGTYR spectra (6.88–7.23 ppm). Without using EDC/NHS, tyramine did not bind to ALG excluding an unspecific reaction.

The captive bubble method was used to characterize the wettability of the investigated surfaces [Fig. 2(B)]. Cell culture-treated PS were found to be hydrophilic (39.8 ± 0.9°). ALG exhibited contact angles of 46.7 ± 0.5°. The ALG functionalization with carbodiimide and different tyramine quantities resulted in contact angles of 56.2 ± 0.9° (EDC/NHS), 51.1 ± 0.5° (2.5 mg TYR/cm²), 43.9 ± 2.2° (7.5 mg TYR/cm²), and 38.8 ± 1.3° (12.5 mg TYR/cm²). The surface charge of the scaffolds was analyzed using zeta potential measurements [Fig. 2(C)]. PS and unmodified ALG surfaces exhibited highly negative charges (−33.4 ± 2.5 and −23.5 ± 2.9 mV, respectively). The conjugation of tyramines to ALG resulted in increased zeta potentials. However, zeta potentials were reduced with increasing tyramine content.

Besides the surface characteristics, the mechanical stability of the fabricated scaffolds was tested using compression studies [Fig. 2(D)]. ALG possessed an elastic modulus of 40.5 ± 4.7 kPa. The modification with EDC/NHS resulted in stiffer gels (78.8 ± 29.2 kPa). However, with increasing tyramine conjugation (2.5, 7.5, and 12.5 mg TYR/cm²), the hydrogel scaffolds became softer (81.7 ± 5.8, 71.0 ± 14.4, and 60.1 ± 14.9 kPa, respectively) compared to carbodiimide-treated ALG.
Protein adsorption to tyramine-conjugated alginate surfaces

To study the interaction between the prepared scaffolds and proteins, the protein adsorption was evaluated in terms of protein content using Ellman’s reagent and stability via immunofluorescence staining against laminin. In addition, the impact of proteins on the scaffolds characteristics was determined. As shown in Figure 3(A), protein adsorption increased with tyramine conjugation to ALG. In comparison to PS, proteins adsorbed up to five times more on ALGTYR (12.5 mg TYR/cm²). Immunofluorescence staining against laminin showed that the adsorption of laminins to ALGTYR was stable for at least 14 days of observation [Fig. 3(B)]. Due to the attached proteins, the surface characteristics altered. After protein adsorption, the wettability of all surface were equalized at 34.9 ± 0.9° [Fig. 2(B)]. Moreover, similar surface charges were found (-12.1 ± 1.5 mV) [Fig. 2(C)]. However, protein coating did not affect the mechanical stability of ALGTYR scaffolds [Fig. 2(D)].

Cell behavior of MSCs and hiPSC-CMs

ALGTYR surfaces were examined in cell culture experiments with MSCs and hiPSC-CMs in comparison to PS and ALG. No cell adhesion was observed on unmodified ALG surfaces [Fig. 4(A/D)]. The cells remained round in suspension. In contrast, MSCs adhered and spread with high adhesion rates on PS and ALGTYR surfaces [Fig. 4(B–D)]. To exclude that intermediates of the tyramine coupling process to ALG triggered the cell adhesion, MSCs were cultivated using the hanging droplet method for 8 days on spherical ALG with different modifications (Supporting Information, Fig. 2). Only ALGTYR microcarriers showed MSC adhesion.

To further validate the potential of the modified scaffolds, hiPSC-CMs were applied exemplarily as cells that require a more complex microenvironment. Therefore, ALGTYR surfaces were coated with the protein mixture Matrigel. hiPSC-CMs were seeded onto planar and structured modified scaffolds and attached to both soft scaffolds. During the 21 days of cultivation, the cells adhered without a certain pattern on planar surfaces [Fig. 4(E)]. However, providing structured surfaces using μ-contact printing resulted in the orientation of the hiPSC-CMs according to the pattern of parallel lines within 3 days [Fig. 4(F)]. The beating frequencies of the hiPSC-CMs appeared to be independent from the surface structure [Fig. 4(G)]. Nevertheless, the provided structures resulted in a more uniform contraction of the hiPSC-CMs in comparison to those on planar surfaces. Moreover, the cultivation of hiPSC-CMs was successfully translated from 2D layer to spherical matrices (Supporting Information, Fig. 3). Coating the spherical ALGTYR microcarriers with 2.2 μg Matrigel/cm² surface area was sufficient for stable cell adhesion and contraction similar to hiPSC-CMs spheroids.

DISCUSSION

Alginate-based hydrogels represent promising biomaterials due to their biocompatibility and adjustable mechanical and porous characteristics. As native alginate inhibits cell adhesion, functionalization strategies are needed to address cell–matrix contacts. Hence, this study aimed to enable stable cell adhesion via the conjugation of tyramine to ALG mimicking the surface chemistry of standard cell culture-treated PS.

Therefore, carbodiimide chemistry was used to initialize the amidation between tyramine and ALG. The reproducible introduction of phenolic groups to ALG was confirmed using 1H NMR spectroscopy (Fig. 1). In consequence, ALGTYR bearing hydroxyl, carboxyl, and phenolic groups possess functional units comparable with the chemical composition of PS. Without the presence of EDC/NHS, no covalent binding of tyramine to ALG takes place. In contrast, the esterification of ALG with NHS to active intermediates results in scaffolds with increased hydrophobicity and surface charge due to the substitution of the carboxyl OH group with succinimidyl units (Fig. 2). Moreover, compression studies revealed an increased stiffness of the NHS-conjugated alginate gels indicating a functionalization process not only on the scaffold surface but also of the porous bulk toward denser networks. After that, the succinimidyl units of the active NHS-alginate scaffolds are substituted by tyramine resulting in increasing hydrophilicity and negative surface...
charge due to the introduced, phenolic OH groups. Here, the coupling of 12.5 mg tyramine to 1 cm² ALG creates a surface wettability comparable to cell culture treated PS surfaces (PS: 39.8 ± 0.9° and 12.5 ALGTYR: 38.8 ± 1.3°). In contrast to stiff PS (Young’s modulus = 3 × 10⁹ Pa²⁵), ALGTYR scaffolds are porous and soft hydrogels (60–80 × 10³ Pa) meeting the mechanical stabilities of native tissues (e.g., human liver: 20–60 × 10³ Pa²⁶ and human heart: 10–150 × 10³ Pa²⁷).

Moreover, protein adsorption was found to be enhanced with increasing tyramine conjugation. As tyrosine interactions...
are known to stabilize protein assemblies, the proteins tyrosine residues interact with ALGTYR resulting in an increasing content of adsorbed proteins which attached firmly for several weeks. As a consequence of the protein coating, various surfaces exhibit the wettability and surface charges of the protein layer. Hence, different protein compositions are applied for cell cultivation instead of diverse material characteristics. These protein-based binding sites are crucial to promote cellular interactions. In contrast to existing strategies to bioactivate alginates via the conjugation of RGD or YIGSR motifs, ALGTYR scaffolds allow noncovalent and universal surface coatings with adhesion molecules of different sources (e.g., serum proteins, Matrigel, laminins, etc.) adjustable in terms of protein type and quantity. As a consequence, an appropriate presence of proteins on the alginate hydrogel surface is created to induce cell adhesion compared to the previous noncovalent mixing of proteins with the alginate sol before gelation.

Subsequently, MSCs and hiPSC-CMs were used to examine the bioactivity of native and modified scaffolds in comparison to PS. Cell culture experiments revealed the bioinertness of unmodified alginates. Although proteins adsorb to the material, no cell adhesion occurred on ALG. The bioinert characteristic of ALG is attributed to tremendous conformational changes of adsorbed domains to denatured proteins. Active NHS intermediates also prevent cell adhesion, as carbodiimide are known to be cytotoxic.

The functionalization to ALGTYR results in bioactive scaffolds on which cells like MSCs and hiPSC-CMs adhere. In contrast to the work of Zhu et al., which revealed no adhesion of NIH 3T3 mouse fibroblasts to the gel surface of their prepared HRP-mediated alginate-tyramine hydrogels, adsorbed proteins on ALGTYR surfaces in this study enable stable cell adhesion and proliferation comparable to PS-based cell culture. Moreover, the soft, bioactive ALGTYR matrices support the functional cell performance like the beating of hiPSC-CMs (see Supporting Information, Movie JBMRA_suppmov1_PS for contraction of hiPSC-CMs on PS coated with Matrigel and Movie JBMRA_suppmov2_ALGTYR for contraction on ALGTYR surfaces with Matrigel coating at day 7). Uniform hiPSC-CMs contractions become feasible due to the alignment of the cells using μ-contact printing. In contrast to stiff cell culture plastic, ALGTYR hydrogels can be compressed by the cellular tension while mimicking the in vivo mechanical properties and cell alignment. The stable protein adsorption and cell–matrix interactions on soft alginates surfaces qualify ALGTYR hydrogels as a promising platform for bioactive scaffolds applicable in stem cell culture and tissue engineering. For an enhanced understanding of the established cell–matrix interactions, future studies involve gene expression analyses and in vivo evaluations. Here, the differentiation potential and metabolic profiles of the seeded (stem) cells and the biointegration in vivo are of particular interest.

**FIGURE 4.** In comparison to native alginate scaffolds (ALG, A) and standard cell culture treated polystyrene surfaces (PS, B) tyramine-alginate scaffolds (ALGTYR, C) were examined in cell culture experiments with MSCs (A–D) and hiPSC-CMs (E–G). ALGTYR surfaces coated with Matrigel (E,F) were found to be a bioactive surface supporting cell adhesion and contraction of hiPSC-CMs. Providing structured surfaces using μ-contact printing resulted in oriented contraction of hiPSC-CMs according to the pattern of parallel lines. Differences of groups were considered significant by $p < 0.05$ (*).
CONCLUSION

Tyramines were conjugated reproducibly to alginate hydrogels using carbodiimide to overcome the bioinertness of alginate scaffolds and establish stable cell–matrix interactions. Due to the introduction of the tyramines phenolic units to the alginate backbone, a surface chemistry comparable to standard cell–culture plastic PS was created. As a consequence, ALGTYR hydrogels exhibit similar wettability to PS and enable robust protein adsorption to soft and porous scaffolds possessing mechanical characteristics of human tissues. Cell culture studies with MSCs and hiPSC-CMs revealed stable cell adhesion and contraction on (structured) 2-D layer and spherical matrices. Hence, the functionalization of bioinert alginate-based hydrogels with tyramines represents an efficient technique to produce bioactive scaffolds beneficial for robust cell culture and tissue engineering. According to the generated data, ALGTYR hydrogels bear the potential to provide a versatile surface for tissue engineering. According to the generated data, ALGTYR cell-based models, and tissue substitutes.

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DISCLOSURE

The authors declare that they have no conflict of interest.

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