Nanofibrillar hydrogel scaffolds from recombinant protein-based polymers with integrin- and proteoglycan-binding domains

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Abstract: This study describes the design, production, and testing of functionalized variants of a recombinant protein-based polymer that forms nanofibrillar hydrogels with self-healing properties. With a view to bone tissue engineering applications, we equipped these variants with N-terminal extensions containing either (1) integrin-binding (RGD) or (2) less commonly studied proteoglycan-binding (KRSR) cell-adhesive motifs. The polymers were efficiently produced as secreted proteins using the yeast Pichia pastoris and were essentially monodisperse. The pH-responsive protein-based polymers are soluble at low pH and self-assemble into supramolecular fibrils and hydrogels at physiological pH. By mixing functionalized and nonfunctionalized proteins in different ratios, and adjusting pH, hydrogel scaffolds with the same protein concentration but varying content of the two types of cell-adhesive motifs were readily obtained. The scaffolds were used for the two-dimensional culture of MG-63 osteoblastic cells. RGD domains had a slightly stronger effect than KRSR domains on adhesion, activity, and spreading. However, scaffolds featuring both functional domains revealed a clear synergistic effect on cell metabolic activity and spreading, and provided the highest final degree of cell confluency. The mixed functionalized hydrogels presented here thus allowed to tailor the osteoblastic cell response, offering prospects for their further development as scaffolds for bone regeneration.

Key Words: KRSR cell-adhesive domain, Pichia pastoris, protein-based polymers, self-assembled hydrogel, tissue engineering

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INTRODUCTION

In natural tissues, cells are surrounded by the extracellular matrix (ECM), which ensures the mechanical support and signaling necessary for regulation of cell metabolism and functions such as migration, proliferation, differentiation, and apoptosis.1–4 The ECM can be seen as a complex hydrogel containing multiple functionalities and bioactive groups at different length scales,5 which synergistically influence cell response. Mimicry of the properties and structure of the ECM remains a key objective in biomaterial design,2,6,7 which is why hydrogels are considered particularly promising candidates for use as scaffolds in tissue engineering.3,8–10

As highly hydrated polymers, hydrogels are usually biocompatible, noncytotoxic, well permeable to oxygen and nutrients, and often biodegradable.5,10 Hydrogels based on natural proteins and carbohydrates of the ECM, as well as a variety of synthetic polymers, have been investigated as biomaterials.5 Despite their generally weak mechanical strength, hydrogels have not only shown promise in soft tissue engineering but also in bone regeneration applications.11 In applications not requiring gap filling, the hydrogel must primarily facilitate delivery of cells and/or release of growth factors.11 Coating of surgical implants with hydrogels that provide biological cues, growth factors,
or antibiotics can promote osseointegration. Composite hydrogels complexed with ceramics or other additives have been devised to improve the mechanical properties, although no study has yet applied hydrogels in a load-bearing bone defect model. Injectable hydrogels, that is, hydrogels that form in situ by chemical or physical crosslinking, are also of interest. Interest in these systems, however, has not been as high due to their high potential to form gels that are not rigid enough to support the loads required by healing bones. This has been the primary reason why high-yield biotechnological production of functionalized variants of this protein-based polymer, and analyze the potential of the resulting fibrillar hydrogels for osteoblast cell culture. By mixing functionalized proteins, containing either RGD or KRSR motifs, and non-functionalized proteins in different ratios, hydrogel scaffolds were conveniently prepared with precisely controlled overall concentration of the two motifs. We show that this tailoring allowed to control osteoblast cell adhesion, activity, spreading, and formation of a confluent layer.

**MATERIALS AND METHODS**

**Construction of recombinant strains**

The production strain for triblock copolymer C2S4H-C2 has been described by us previously. Two variants of the corresponding gene were designed; one encoding N-terminal RGD-motifs (B(RGD)C2S4H-C2), and another encoding N-terminal KRSR-motifs (B(KRSR)C2S4H-C2). These genes and corresponding strains were constructed as follows.

A double-stranded adapter B(RGD) (short for cell-binding block with RGD motifs) with HindIII and Bsal overhangs was prepared by annealing of oligonucleotides

5'-AGCTTGAATTCGAGGATCTCTTGAGATCCGG
TGGAGGTTCTGGAGGTGGATCTGGTGGAAGGTGACTCACCAGG

TCTCG-3' and

5'-GCACCGAGACCTGGTAGCTCCTCCTTCCACCCACCAGATCCACC
TCCAGAACACCTCCAGGACCCACCAGGAAATCTCCCTTGCC

GAATTA-3'. This adapter encodes two GGGSPG motifs separated by a (GGSG)3 flexible spacer. Likewise a double-stranded adapter B(KRSR) (short for cell-binding block with KRSR motifs) encoding three repeats of the sequence PKRSRGGG was prepared by annealing of oligonucleotides

5'-AGCTTGAATTCGAGGATCTCTTGAGATCCGG
TGGAGGTTCTGGAGGTGGATCTGGTGGAAGGTGACTCACCAGG

TCTCG-3' and

5'-GCACCGAGACCTGGTAGCTCCTCCTTCCACCCACCAGATCCACC
TCCAGAACACCTCCAGGACCCACCAGGAAATCTCCCTTGCC

GAATTA-3'. These adapters were inserted into the HindIII/BSal sites of vector PMTL23-C2S4H-C2 resulting in vectors pMTL23-B(RGD)C2S4H-C2 and pMTL23-B(KRSR)C2S4H-C2, respectively. These vectors were then digested with EcoRI/NotI, and the released inserts were cloned into the EcoRI/NotI sites of Pichia pastoris expression vector pPIC9 (Thermo Fisher Scientific). The vectors pPIC9-B(RGD)C2S4H-C2 and pPIC9-B(KRSR)C2S4H-C2 thus obtained were linearized with Sall and used to transform P. pastoris GS115 by electroporation as described previously.

**Fermentation and protein purification**

Methanol fed-batch fermentations of the B(RGD)C2S4H-C2 and B(KRSR)C2S4H-C2 strains at 30°C and pH 3 were performed in 2.5-L Bioflo 3000 bioreactors (New Brunswick Scientific) as described previously for C2S4H-C2. At the end of fermentation, the cells were separated from the broth by centrifugation for 10 min at 10,000 × g (RT) in an SLA-3000 rotor (Sovall), and the supernatant was microfiltered.
Proteins were purified from the supernatant as previously described for \( C_2S_{16}C_2 \), with the exception that the acetone precipitation step was omitted. Shortly, proteins were selectively precipitated twice at 45% ammonium sulfate saturation and subsequently dialyzed, filter sterilized (0.2 µm), and lyophilized.

**SDS-PAGE, N-terminal protein sequencing, amino acid analysis, and carbohydrate determination**

The NuPAGE Novex system (Thermo Fisher Scientific) was used for SDS-PAGE, with 10% Bis-Tris gels, MES SDS running buffer, and SeeBlue Plus2 prestained molecular mass markers. Gels were stained with Coomassie SimplyBlue Safe-Stain (Thermo Fisher Scientific).

Purified proteins in solution were N-terminally sequenced via Edman degradation by Midwest Analytical (St. Louis, MO). Amino acid composition analysis after acid hydrolysis was performed by Ansynth Service B.V. (Roesendaal, The Netherlands). Protein purity was estimated by linear least-squares fitting to the obtained amino acid composition of (1) the theoretical composition of the respective pure protein and (2) the composition determined for host-derived proteins present in the medium.

Because polysaccharides are the major nonprotein contaminant in the extracellular medium of \( P. \) *pastoris* cultures, the carbohydrate content in the purified protein was determined using a phenol–sulfuric acid assay as previously described.\(^{37}\) Proteins contained <2% (w/w) of polysaccharides.

**Mass spectrometry**

Matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry was performed using an ultrafleXtreme mass spectrometer (Bruker). Samples were prepared by the dried droplet method on a 600 µm AnchorChip target (Bruker), using 5 mg/mL 2,5-dihydroxyacetophenone, 1.5 mg/mL diammonium hydrogen citrate, 25% (v/v) ethanol, and 1% (v/v) trifluoroacetic acid as matrix. Spectra were derived from 1000-shot (1 kHz) acquisitions taken at nonoverlapping locations across the sample. Measurements were made in the positive linear mode, with ion source 1, 25.0 kV; ion source 2, 23.3 kV; lens, 6.5 kV; pulsed ion extraction, 680 ns. Protein Calibration Standard II (Bruker) was used for external calibration.

**Atomic force microscopy**

Protein samples of (A) \( C_2S_{16}C_2 \), and (B) \( B^{KRSR}C_2S_{16}C_2 \) were prepared in a 1:1 ratio and dissolved in 10 mM HCl (final pH 2 to 3). The pH of the ~2 mL samples was adjusted to 7.4 ± 0.2 by addition of 0.1 M NaOH, using a pH meter with a micro electrode (Mettler Toledo InLab\(^{®}\) Micro). Milli-Q was then added to obtain protein solutions between 0.05% and 0.1% (w/v). After 24 h of incubation at room temperature, solutions were pipetted onto silica wafers and left for 20 min to allow fibril deposition. Wafers were subsequently rinsed with Milli-Q to remove salts. The samples were dried by removing the bulk of the solution with a tissue and/or using a mild stream of nitrogen. The dry samples were analyzed with a Nanoscope V (Veeco), in Scan Asyst mode, using a nonconductive silicon nitride probe (NP-10, Bruker) with a spring constant of 0.32 N/m. Images were recorded at a frequency of 0.997 Hz and processed with NanoScope Analysis 1.20 software (Veeco).

**Rheology**

Solutions of the purified proteins (A) \( C_2S_{16}C_2 \), (B) \( B^{KRSR}C_2S_{16}C_2 \), and (C) \( B^{KRSR}C_2S_{16}C_2 \) were prepared in 10 mM HCl and vortexed for 1.5 h using a multi-tube holder. The pH of the ~2 mL samples was adjusted to 7.4 ± 0.2 by addition of 0.1M NaOH, using a pH meter with a micro electrode (Mettler Toledo InLab\(^{®}\) Micro). PBS (phosphate buffered saline, pH 7.4, Sigma-Aldrich) was then added to reach a 2% (w/v) protein concentration. Solutions were loaded immediately into the rheometer (Physica MCR 501 Rheometer, Anton Paar) with a Couette CC10/T200 geometry (bob diameter 10.002 mm, cup diameter 10.845 mm). Each sample was placed in the measuring cell on top of 500 µL of perfluorinated fluid (Galden\(^{®}\) HT-70, Solvay Specialty Polymers) to decrease the sample volume, and covered with paraffin oil to avoid evaporation. Gel formation was analyzed by recording storage moduli over time under sinusoidal deformation (frequency (f) = 1 Hz and strain (γ) = 1%). After 15 h, gels were broken (f = 1 Hz and logarithmically increasing deformation γ = 0.1 to 100%), Self-healing behavior was then monitored by recording storage moduli under sinusoidal deformation as before. All measurements were performed at 37°C.

**Preparation of scaffolds for cell culture**

Freeze-dried proteins were separately dissolved at 4% (w/v) in 10 mM HCl for at least 2 h by vortexing with a multi-tube holder. Solutions were then mixed to obtain 1 mL samples of the required protein compositions. To allow simultaneous preparation of multiple scaffolds without pH measurement, one volume of 0.1 M NaOH was added to adjust the protein concentration to 2% (w/v) and to reach a pH above neutral in all samples. This ensures proper gel formation across samples independent of the precise pH, as self-assembly occurs at pH 6 or above.\(^{36}\) The protein solutions were immediately transferred to 48-well cell culture plates (250 µL per well) and left for 2 h at room temperature to allow full setting of the gels. These 2 h are a conservative approach, as gels already form (crossover point of storage modulus and loss modulus) within minutes (Supporting Information Figure 1). To ensure a pH value of 7.4 for cell seeding, the gels were further incubated in 0.5 mL of PBS for 3 h and then in 0.5 mL of cell culture medium (α-MEM, 22571, Thermo Fisher Scientific) with 10% (v/v) fetal bovine serum (FBS) for 15 h (with one medium change after 12 h).

**Cell pre-culture**

Cryo-preserved MG-63 cells (ATCC\(^{®}\) CRL-1427\(^{TM}\); LGC Standards GmbH), passage 90, were cultured for 6 days in proliferation medium (α-MEM, 22571, Thermo Fisher Scientific), supplemented with 10% (v/v) FBS, at 37°C, 95%
relative humidity and 5% CO₂. Prior to seeding, cells were washed twice with PBS, enzymatically detached for 5 min with trypsin-EDTA (0.25% (w/v) trypsin, 0.02% (w/v) EDTA), and resuspended at 36,000 cells/mL in osteogenic medium. The medium was composed of α-MEM supplemented with 10% (v/v) FBS, 100 U/mL penicillin and 10 µg/mL streptomycin (Thermo Fisher Scientific), 50 mg/L ascorbic acid (Sigma-Aldrich), 10 nM dexamethasone (Sigma), 10 mM β-glycerophosphate disodium salt hydrate (Sigma). Next, 500 µL of cell suspension was added per well (final seeding density: 20,000 cells/cm²). Scaffolds incubated in osteogenic medium without cells were included as controls. All samples were prepared in triplicate (n = 3). The entire study was run for 23 days in an incubator at 37°C, 95% relative humidity and 5% CO₂. The culture medium was refreshed twice a week for all samples.

Cell adhesion
Cell adhesion was analyzed 4 h after cell seeding. The medium was removed from a dedicated set of wells to take out nonadherent cells. Gel surfaces were imaged by randomly taking light microscopy pictures (n = 3 per sample) using an inverted phase-contrast microscope (Leica DMIL). Pictures were taken close to the center of the gels to ensure that the imaged area was relatively flat. The number of cells on the area of 2,048 µm x 1,536 µm was counted in manual mode using ImageJ 1.49o software. Cell adhesion was calculated as the number of cells per mm².

Cell metabolic activity
To assess cell metabolic activity, an alamarBlue® (Thermo Fisher Scientific) assay was performed on a dedicated set of wells on days 1, 3, 10, and 23, following the protocol of the manufacturer. Shortly, the cell culture medium in the wells was exchanged for fresh medium containing 10% (v/v) of alamarBlue® reagent, followed by incubation at 37°C for color development. In view of high cell activity on day 23, an incubation time of only 1.5 h was used for these samples, whereas for the earlier days 4 h was used. The medium containing the reagent was subsequently removed for measuring fluorescence (ex 560 nm, em 590 nm; FLx800 reader, Bio-Tek Instruments), and all values were normalized to activity per hour. To compare cell activity on different scaffolds, the average fluorescence of the medium was calculated (n = 3).

Cell viability and spreading
To study cell viability, and to visualize changes over time in cell distribution and morphology (spreading), a LIVE/DEAD Viability/Cytotoxicity Kit (L3224, Thermo Fisher Scientific) was used on days 1 and 3, and only the calcine-AM compound from the kit on days 10, and 23. Fluorescent staining of F-actin and nuclei was performed on samples from days 1, 3, and 23 for further confirmation of cell distribution and spreading.

For the LIVE/DEAD assay, three samples of each gel type (n = 3) were washed twice with PBS and afterwards incubated in 0.5 mL of PBS with 2 mM calcein-AM and, optionally, 4 mM ethidium homodimer, in the dark, for 30 min at 37°C. Next, scaffolds were washed twice with PBS and cells were visualized using a fluorescence microscope (Axio Imager Microscope Z1; Carl Zeiss Micro Imaging GmbH). Overlay images were prepared, with dead cells appearing in red (ex 517 nm, em 617 nm) and live cells in green (ex 488 nm, em 517 nm).

Samples for fluorescent staining of F-actin and nuclei were prepared as follows. First, the cell culture medium was removed from the cell culture well-plate. Subsequently, the scaffolds in the wells were washed gently with PBS, fixed for 1.5–4 h with 0.7 mL per well of 3.3% PFA (paraformaldehyde) in PBS, and then kept in 70% (v/v) ethanol until the day of imaging (for a maximum of 5 days). Prior to imaging the gels, ethanol was removed, scaffolds were washed twice in PBS, and cells were permeabilized for 20–30 min with 0.8 mL per well of 1% Triton X-100 in PBS containing 1% FBS. Scaffolds were then incubated for 2 h with Alexa Fluor® 568 conjugated phalloidin (Thermo Fisher Scientific) for F-actin staining and with DAPI (Thermo Fisher Scientific) for visualization of nuclei. Both reagents were diluted in PBS containing 1% FBS (1:200 and 1:2,500, respectively), and 0.5 mL of this mixture was added per well. After washing in PBS for 10 min, scaffolds were examined using an Axio Imager Microscope Z1 (Carl Zeiss Micro Imaging GmbH).

Statistical analysis
Statistical analysis was performed using SPSS Statistics software. Statistical differences were estimated using one-way analysis of variance followed by an LSD post-hoc test. Differences were considered significant at p < 0.05, if not stated differently. Results were reported as the mean ± standard deviation.

RESULTS
Polymer design
We previously reported the molecular design of the fibril-forming protein C248C2.35,36 It features an S48H (silk-inspired) middle block consisting of 48 repeats of the octapeptide GAGAGGH. Whereas at low pH the proteins are soluble owing to repulsion by the positively charged histidines, at neutral pH the charges are neutralized36 and the S48H domains form β-rolls or β-sheets.39–41 These, in turn, form the fibril core through stacking, mediated by hydrophobic interactions. The S46H block is flanked on both sides by hydrophilic C2 (random coil) blocks, each consisting of two 99 amino acid-long unfolded domains in tandem, which form a corona surrounding the core and provide colloidal stability [Fig. 1(A)]. At higher protein concentrations (above several g/L), gels are formed, most likely as a result of the entanglement of long fibrils,42 and possibly the occurrence of weak physical crosslinks.36

Here, we designed two functionalized variants of C248S2H C2 with N-terminal cell-binding extensions (B domains, for short). One variant, BBDN C248S2H C2, contains integrin-binding RGD motifs,13 and the other, BRRS C248S2H C2 contains
proteoglycan-binding KSR motifs.\(^{17}\) [Fig. 1(B)]. For the design, the following considerations were taken into account: (1) To potentially increase cell interaction, the design should preferably feature several copies of these motifs. (2) The length of the extensions should be limited, so as to minimize possible effects on the well-characterized fibril-forming properties of the \(C_2S_{48}^{SH}C_2\) core component. (3) The repeated cell-binding motifs should be separated by suitable flexible linkers to allow sufficient conformational freedom.

In \(B^{RGD}C_2S_{48}^{SH}C_2\), a flexible Gly/Ser-rich sequence was chosen to separate the RGD motifs because this type of spacer is widely used as a constituent of engineered proteins produced in various hosts including \(P.\) \(pastoris.\)\(^{44,45}\) A spacer length of 12 residues in the form of three GSGG repeats was used. Two RGD motifs were incorporated, where the specific motif used was GRGDSP from fibronectin.\(^{33}\) The N-terminal extension relative to \(C_2S_{48}^{SH}C_2\) thus became: GRGDSPGGSGGSGGSGGRGDSP, which adds ~3% to the length of the protein sequence. This sequence is N-terminally preceded by the cloning-derived sequence YVEF, as is the case in \(C_2S_{48}^{SH}C_2.\)

In the design of \(B^{KRSR}C_2S_{48}^{SH}C_2\), poly-glycine stretches were used as spacers between the KRSR motifs, in view of a study that reports osteoblast adhesion onto glass surfaces with C-terminally immobilized KRSR, KRSRG\(_3\), or KRSRG\(_6\) peptides.\(^{17}\) However, as the authors found similar adhesion efficiency for all peptides tested, spacer length does not appear to be critical. Rather, because a tetravalent poly-lysine peptide core with four attached KRSR peptides performed significantly better, the authors suggest peptide density may be more important. We, therefore, incorporated three KRSRG\(_3\) motifs, while maintaining the same small length of the N-terminal extension, namely 24 residues, as used in \(B^{RGD}C_2S_{48}^{SH}C_2.\) Another critical design consideration was to prevent proteolysis of Lys-Arg in the KRSR motifs by the \(P.\) \(pastoris\) homolog of Kex2 protease. This serine endoprotease was first identified in \(S.\) \(cerevisiae\) as the protease required for maturation of the -mating factor pheromone, via cleavage at the C-terminal side of Lys-Arg.\(^{46}\) As we have shown previously,\(^{36,37}\) \(P.\) \(pastoris\) Kex2 can also digest particular mono-arginyl sites, as governed to a large extent by the residues at positions -2 to -4 relative to the scissile bond. Under the assumption that \(P.\) \(pastoris\) and \(S.\) \(cerevisiae\) Kex2 have similar substrate specificity, the extent of putative cleavage at the C-terminal side of the KRSR sequences is likely low, in view of the moderately unfavorable Ser at position -2 relative to the scissile bond.\(^{49}\) As for potential digestion of the canonical Lys-Arg motif in KRSR, we reasoned it might be possible to minimize this by placing highly unfavorable residues at positions -4 and -3. We thus arrived at the following sequence design for the N-terminal \(B^{KRSR}\) extension: GPKRSGGGPKRSGGGPKRSGGGGPK (again N-terminally preceded by the cloning-derived sequence YVEF, and C-terminally followed by the \(C_2S_{48}^{SH}C_2\) sequence starting with Gly). For all three Lys-Arg motifs in this sequence, position -4 relative to the scissile bond is occupied by Gly, and position -3 by Pro. Both amino acids rank as highly unfavorable in these positions, at least for \(S.\) \(cerevisiae\) Kex2.\(^{49}\)

**Protein synthesis and characterization**

The polymers were produced as secreted proteins in methanol fed-batch bioreactor cultures of \(P.\) \(pastoris,\) and purified as previously described.\(^{37}\) Based on the determined amino acid composition, the purity of \(B^{RGD}C_2S_{48}^{SH}C_2\) was estimated at 99.8%, and that of \(B^{KRSR}C_2S_{48}^{SH}C_2\) at 99.3%. The gravimetrically determined yields of the purified, desalted, and lyophilized material, expressed per volume of cell-free fermentation broth, were 3.2 g/L for \(B^{RGD}C_2S_{48}^{SH}C_2\) and 3.5 g/L for \(B^{KRSR}C_2S_{48}^{SH}C_2.\)

The functionalized proteins and unmodified \(C_2S_{48}^{SH}C_2\) were analyzed by SDS-PAGE (Fig. 2). The main bands of all three proteins migrated in SDS-PAGE at an apparent molecular weight of ~105 kDa, although the calculated masses are ~66 to ~68 kDa. As described previously for \(C_2S_{48}^{SH}C_2,\)\(^{36,37}\) this discrepancy is due to the high content of hydrophilic and small amino acids in this protein. The faint upper band and
low mass smear visible in all proteins are probably SDS-PAGE artifacts, as has previously been shown for $C_2S_{48}H_{4}C_2$. N-terminal sequencing of $B^{RGG}C_2S_{48}H_{4}C_2$ showed the expected YVEFGRGD sequence without any secondary reactions, confirming that the protein was N-terminally intact. Likewise, N-terminal sequencing of $B^{KRSSR}C_2S_{48}H_{4}C_2$ revealed mainly the expected YVEFGPKRSR sequence, which shows that most of the N-terminal $B^{KRSSR}$ extension was N-terminally intact. However, besides spurious background signals, a minor secondary species, most likely corresponding to SRGG, was also observed in this sample. This sequence may have resulted from a limited extent of Kex2 cleavage at the C-terminal side of the dibasic GPKR sequences. The molecular mass of the proteins was verified by MALDI-TOF (Fig. 3). For both $B^{RGG}C_2S_{48}H_{4}C_2$ and $B^{KRSSR}C_2S_{48}H_{4}C_2$, the observed masses of the main peaks (67,932 and 68,412 Da, respectively) correspond, within experimental error, to the calculated masses of the intact proteins (68,061 and 68,535 Da, respectively). The MALDI-TOF spectra furthermore show that $B^{RGG}C_2S_{48}H_{4}C_2$ was fully monodisperse, and $B^{KRSSR}C_2S_{48}H_{4}C_2$ nearly so. The slight shoulder to the left of the main peak in the spectrum of $B^{KRSSR}C_2S_{48}H_{4}C_2$ (indicated by an arrow in Fig. 3) contains three peaks whose masses correspond to that of the protein after N-terminal truncation following either the first, second, or third GPKR motif. This result is in agreement with the minor SRGG species observed in N-terminal sequencing. Additionally, three minor peaks are present, with masses that could correspond to N-terminal truncation following either the first, second, or third KRSSR motif. Apparently, Kex2 cleavage occurred at the C-terminal side of the dibasic GPKR and monobaric KRSSR motifs, but only at a very low level. The shoulder to the right side of the main peak of both proteins (Fig. 3) corresponds to Glu-Ala extended species in a small portion of the molecules, as observed previously for $C_2S_{48}H_{4}C_2$.37 and results from incomplete processing of the $\alpha$- mating factor propeptide.38

Material characterization

AFM and rheology were performed to compare the fibril formation and mechanical properties of $B^{RGG}C_2S_{48}H_{4}C_2$ and $B^{KRSSR}C_2S_{48}H_{4}C_2$ with those previously established for $C_2S_{48}H_{4}C_2$.36,37 For AFM (Fig. 4), dilute solutions in 10 mM HCl were prepared of the unmodified $C_2S_{48}H_{4}C_2$ protein, and of a 1:1 mixture of the functionalized proteins $B^{RGG}C_2S_{48}H_{4}C_2$ and $B^{KRSSR}C_2S_{48}H_{4}C_2$. After raising the pH to a physiological level of 7.4 and 24 h of incubation, both samples showed self-assembled fibrils in AFM with a length in the micrometer range, as described previously for $C_2S_{48}H_{4}C_2$.36,37 This indicates that the fibril forming capacity was not affected by functionalization of the molecules.

A rheological study (Fig. 5) revealed that gel formation is as well relatively unaffected by the addition of functional domains to $C_2S_{48}H_{4}C_2$. After adjusting the pH to a physiological level of 7.4, 2% (w/v) solutions of $C_2S_{48}H_{4}C_2$, $B^{RGG}C_2S_{48}H_{4}C_2$, and $B^{KRSSR}C_2S_{48}H_{4}C_2$ all developed storage moduli in the range of 500–1,500 Pa. The observed differences
in the final storage modulus and in the kinetics of gel formation can be attributed to batch-to-batch variations. Also, minor differences in pH adjustment could play a role, in view of the known influence of pH on the storage modulus of C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>.<sup>36</sup> All proteins achieved a storage modulus above 200 Pa already after 1–2 h, and reached a plateau after 5 h. Gelation, defined as the crossover point of storage modulus and loss modulus, occurred within minutes after adjusting pH (see Supporting Information Figure 1). As has been shown for C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>,<sup>36</sup> also BRGDC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> and BKRSRC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> retained the ability to quickly recover after breaking, reaching storage moduli of 58–83% of the initial values.

**Cell culture**

Scaffolds for two-dimensional cell culture studies were obtained by preparing 2% (w/v) gels with six different protein compositions, as listed in Table 1. This design allowed to determine the influence of the incorporated functional domains on cell behavior. By mixing BRGDC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> and BKRSRC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>, hydrogels are obtained that possess both RGD and KRSR motifs. By mixing C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> with either BRGDC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> or BKRSRC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>, a reduced content of these cell-adhesive domains is obtained.

As we aim at developing materials for future use as bone cell scaffolds, and investigate the influence of KRSR as bone cell-specific adhesive motifs, MG-63 osteoblastic cells were used in the cell culture studies. The influence of the active domains on early cell response was studied by evaluating cell adhesion (Fig. 6). As compared to the unmodified 0R/0K gel, the number of adherent cells was significantly higher on the functionalized gels, except for 50K. The simultaneous presence of both RGD and KRSR did not have a synergistic effect on cell adhesion.

The largest variation in the number of adherent MG-63 cells was found in unmodified 0R/0K. Apparently, the distribution of cells on the surface of the scaffold was not uniform. In accordance with our previous study, where rat bone marrow stem cells (MSC) were grown on nonfunctionalized C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>,<sup>37</sup> this suggests that scaffolds containing C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> without adhesive domains are not particularly favorable for cell attachment. In contrast, the smallest variation was found for the 100R scaffold, which indicates a consistent positive effect from the RGD domains.

The effect of addition of RGD and KRSR domains on cell metabolic activity, as a measure of viability and possible proliferation, was tested with the alamarBlue<sup>®</sup> assay (Fig. 7). In this assay, the oxidized reagent resazurin is added to the culture medium, which after entering viable cells is reduced to resorufin by mitochondrial enzymes. This reduction can be observed as a change in color.<sup>50</sup>

Already after one day of cell culture, significant differences in cell activity were observed. Cells on the 50R, 100R, and 50R/50K scaffolds (i.e., scaffolds containing RGD domains) were metabolically more active in comparison with unmodified material. On day 23, 50R

![FIGURE 4](image4.png) AFM of 0.1% (w/v) protein solutions after 24 h of incubation at pH 7.4 and room temperature: A: C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>, and B: C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>/BRGDC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub> mixed in a 1:1 ratio.

![FIGURE 5](image5.png) Rheometry of 2% gels of C<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>, BRGDC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>, and BKRSRC<sub>2</sub>S<sub>H</sub>48C<sub>2</sub>. Development of storage moduli as a function of time during gel formation, and recovery after strain-induced breakage of the gel at ~15 h.

![FIGURE 6](image6.png) Number of cells adherent to the scaffold per mm<sup>2</sup> after 4 h of incubation. * p < 0.05, compared to 0R/0K; ** p < 0.07 compared to 0R/0K; *** p < 0.05 compared to 50K; (n = 3).

| TABLE I. Protein Composition of 2% (w/v) Cell Culture Scaffolds |
|---------------------------------------------------------------|
| **Scaffold** | **Relative Composition** |
|---------------|-------------------------|
| 0R/0K         | 100% – –                |
| 50R           | 50% 50% –               |
| 100R          | – 100% –                |
| 50K           | 50% – 50%               |
| 100K          | – – 100%                |
| 50R/50K       | – 50% 50%               |

### *Nanofibrillar Hydrogel Scaffolds*

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showed significantly more impact on cell activity than 50K and 100K, respectively, suggesting a stronger influence of RGD than of KRSR domains. Nonetheless, a synergistic effect of the scaffold modified with both functionalities (KRSR and RGD) became apparent from day 1 onward. With significantly higher metabolic activity of cells on 50R/50K compared to most of the other gel variants at each time point, there is a clear tendency for superior metabolic responses to our material when it contains both functionalities.

Cell survival was assessed after 1 and 3 days with a LIVE/DEAD assay. As visible in Figure 8, for all scaffold types, only a few individual cells were identified as dead (red-stained nuclei), and the rest remained alive (green cells). For $C_2S_{48}C_2$, this result is in agreement with previous findings for scaffolds prepared from that protein. However, modification with RGD and/or KRSR motifs did not negatively affect the cytocompatibility of the gels. The LIVE/DEAD images (Fig. 8) also revealed changes over time in terms of cell morphology and cell distribution. Already on day 1, cells seeded on the scaffolds containing RGD motifs (50R, 100R, 50R/50K) started to show a more stretched morphology. On day 3, some cells with elongated shape were observed also on the gels exclusively containing KRSR domains (50K, 100K), although to a lesser extent than on scaffolds containing RGD. On day 3, the scaffold containing both RGD and KRSR motifs (50R/50K) yielded the highest degree of confluency and thus seemed most profitable for the cells. As of day 10, these samples showed a fully confluent layer. Also the 100K scaffold highly favored cell spreading and formation of a confluent layer on days 10 and 23. Interestingly, 50K scaffolds instead led to formation of clusters, and did not show cell spreading until day 23. A similar overall trend as apparent in Figure 8 was observed with fluorescent phalloidin/DAPI staining for the visualization of actin filaments and nuclei (Supporting Information Figure 2).

**DISCUSSION**

In previous work, we have shown that the $C_2S_{48}C_2$ protein-based polymer can form self-assembling, noncytotoxic fibrillar gels allowing to maintain viable cells for at least 21 days. However, the cells did not reveal a spread morphology over time, and did not proliferate to form a confluent layer. To further develop our material toward use as a scaffold in bone cell culture, we created in this study two variants of the protein with a genetic design that includes either integrin-binding RGD motifs, or proteoglycan-binding bone cell-specific KRSR motifs.

The polymers were efficiently produced as secreted proteins in P. pastoris at up to 3.5 g/L of cell-free broth. The possibility to obtain multiple grams of these polymers from benchtop bioreactors readily allows their use in preclinical research. The $B^{RGD}C_2S_{48}C_2$ polymer produced was completely monodisperse. Also $B^{KRSR}C_2S_{48}C_2$ was found to be nearly intact, apart from a very minor level of proteolytic cleavage in the KRSR motif C-terminal of Lys-Arg, and probably also C-terminal of Ser-Arg. As described in Results, proteolytic processing of these motifs in the N-terminal random-coil $C_2$ block by the endogenous Kex2 protease of P. pastoris had been anticipated. The fact that this protease is responsible for processing at Lys-Arg of the $\alpha$-mating factor prepro peptide, typically used to drive high-level secretion of heterologous proteins in this host, only attests to the enzyme’s processivity and/or abundance. In our sequence design, we therefore attempted to particularly minimize cleavage to the C-terminal side of the canonical Lys-Arg motif in KRSR by placing expectedly unfavorable residues at position $-4$ and $-3$, relative to the scissile bond. Because only a negligible level of degradation was observed, this approach was apparently remarkably effective. To our knowledge, this represents the first reported case where Kex2 cleavage of a dibasic Lys-Arg motif in a recombinant protein was largely prevented by the careful design of the preceding sequence.
context. Importantly, the essentially monodisperse nature of the polymers adds to the level of control gained over the properties of the scaffolds produced from them.

By mixing nonfunctionalized $C_2S_{48}C_2$ with $B^{RGG}_C S_{48}S_{48}$ and/or $B^{KRSR}_C S_{48}S_{48}C_2$ in a chosen ratio (Table I), tailored fibrillar scaffolds for cell culture were obtained with different concentrations of the functional groups, independent of the total polymer concentration, and thus, in principle independent of properties such as nominal pore size and storage modulus. The storage moduli of gels prepared from either of the functionalized proteins were indeed in the same range as that of $C_2S_{48}C_2$, also in accordance with the observation that functionalized and nonfunctionalized proteins showed similar fibril-forming properties in AFM. Apparently, the functional domains have little or no effect on the physico-chemical characteristics of the proteins. This suggests

FIGURE 8. Cell survival, morphology, and distribution as monitored by the LIVE/DEAD assay. Viable cells show green fluorescence, whereas dead cells appear red (tested on days 1 and 3 only). Scale bar = 200 μm.
that also other domains could be incorporated in the future, allowing convenient tuning of multiple functionalities by mixing. $B_{\text{RGD}}^2C_{29\text{H}4\text{C}_2}$ and $B_{\text{KRSR}}^2C_{29\text{H}4\text{C}_2}$ gels also retained the self-healing characteristics of unmodified $C_{29\text{H}4\text{C}_2}$. This may be a useful property for potential biomedical use, allowing recovery of the gels if inadvertently broken during in situ application or afterwards. While the mixing approach allows to instantaneously prepare hybrid scaffolds containing both RGD and KRSR domains, we do not claim that hybrid fibrils are certainly formed. Stochastically, however, this will most likely be the case, as each fibril consists of thousands of polymer molecules. To the cells it is probably not relevant whether both domains are present in the same or separate fibrils, as each cell is in contact with countless nanofibrils. In either case, there will likely be a certain optimal adhesive domain content, which can be reproduced at will by mixing the proteins in the appropriate ratio.

We previously observed that proliferation and mineralization of primary rat bone MSC were relatively low on non-functionalized $C_{29\text{H}4\text{C}_2}$ in comparison with a collagen control, and anticipated that the cell response may be improved by functionalization of the polymer with cell-adhesive motifs. Indeed, cell adhesion, metabolic activity, and spreading studied here using MG-63 osteoblastic cells were all enhanced on the scaffolds containing either of the functional domains (KRSR or RGD). RGD domains had a slightly more pronounced influence than KRSR. Based on similar observations, Rezania et al. suggested that the principle binding domain is RGD, and that heparin-binding sites are secondary.

A clear synergistic effect on cell metabolic activity and spreading was apparent from day 1 onwards when RGD and KRSR domains were simultaneously present in the scaffolds. Incorporating both integrin-binding and heparin-binding domains may create conditions more similar to the environment of the multifunctional native ECM and may thus be more profitable for the cells. Coughman and Woods suggested that integrins and proteoglycans strongly cooperate during regulation of cell adhesion and spreading.

Interestingly, although cell activity and spreading in our study clearly benefited from the presence of both functional domains, no synergistic effect was seen for adhesion. Possibly, proteoglycan-binding motifs influence the cell response only after initial adhesion and particularly play a role in mediating cell spreading. Similarly, cell adhesion, activity, and spreading in our study did not appear to have a simple causal relation to the degree of confluency. As mentioned above, RGD had a slightly more pronounced influence than KRSR on cell adhesion, activity, and spreading. However, the highest degree of confluency on days 10 and 23 was seen on scaffolds 100K and 50R/50K.

CONCLUSIONS

We described the design and production of functionalized variants of the genetically engineered polymer $C_{29\text{H}4\text{C}_2}$ and evaluated the performance of osteoblastic cells on self-assembled fibrillar hydrogels formed by these proteins. Functionalization involved incorporation of either integrin-binding (RGD) or bone cell-specific heparin-binding (KRSR) sequences into the genetic design. Both proteins, efficiently secreted by *Pichia pastoris* at g/L levels, were essentially monodisperse, and formed self-supporting gels under physiological conditions. Control over the overall concentration of functional groups, independently of the total polymer concentration, was readily achieved by mixing functionalized and nonfunctionalized proteins in various ratios. Functionalization promoted attachment, metabolic activity, and spreading of osteoblastic cells. The combined cell culture studies suggest that the scaffold containing both bioactive domains was most profitable. Overall, these results hold promise for further development of the material toward bone tissue engineering applications.

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