Spatiotemporally controlled, aptamers-mediated growth factor release locally manipulates microvasculature formation within engineered tissues

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ABSTRACT

Spatiotemporally controlled growth factor (GF) delivery is crucial for achieving functional vasculature within engineered tissues. However, conventional GF delivery systems show inability to recapitulate the dynamic and heterogeneous nature of developing tissue’s biochemical microenvironment. Herein, an aptamer-based programmable GF delivery platform is described that harnesses dynamic affinity interactions for facilitating spatiotemporal control over vascular endothelial GF (VEGF165) bioavailability within gelatin methacryloyl matrices. The platform showcases localized VEGF165 sequestration from the culture medium (offering spatial-control) and leverages aptamer-complementary sequence (CS) hybridization for triggering VEGF165 release (offering temporal-control), without non-specific leakage. Furthermore, extensive 3D co-culture studies (human umbilical vein-derived endothelial cells & mesenchymal stromal cells), in bi-phasic hydrogel systems revealed its fundamentally novel capability to selectively guide cell responses and manipulate lumen-like microvascular networks via spatiotemporally controlling VEGF165 bioavailability within 3D microenvironment. This platform utilizes CS as an external biochemical trigger for guiding vascular morphogenesis which is suitable for creating dynamically controlled engineered tissues.

1. Introduction

A long-sought goal of tissue engineering is to successfully bioengineer artificial tissues that could repair or regenerate damaged tissues after implantation in patients. However, establishing hierarchically organized, perfusable and mature vascular networks within engineered tissues is fundamental for their survival post-implantation. [1]. In native tissues, various processes are synergistically responsible for achieving vascularization, including vasculogenesis (de novo vessel formation), angiogenic sprouting, intussusception, vascular remodeling and maturation. Similar to other tissue formation processes, vascularization is also spatiotemporally controlled via various interactions of the cells with the extracellular matrix (ECM) which undergoes constant biochemical modifications. [2]. The complex, dynamic and independent nature of these interactions in native tissues provides great challenges in tissue engineering, but also highlights the need to design dynamic biomaterials that could spatiotemporally control various biochemical cues for better mimicking the in vivo microenvironment.

Spatiotemporally controlled biochemical cues (growth factor, GF) within biomaterials provide unique opportunities for governing defined cell behaviors such as adhesion, proliferation, migration, and differentiation. [3–10]. Recent research has demonstrated a variety of bio-inspired designs that manifest the ability to temporally control GF delivery by being transiently activated with external stimuli such as pH, enzymes, light, abiotic avidin affinity, cell-selective traction forces, or a logic-based integrated combination of multiple stimuli. [11–21]. Although these approaches have their own merits and show efficient functionality in vitro, they have limited in vivo applicability. For instance, light-dependent approaches exhibit significant light scattering and lower absorption at higher tissue depths. [22]. Indeed, diffusion-dependent approaches such as pH leverages on temporal control within deep tissue, but shows suboptimal spatial selectivity. Moreover, synthetic approaches like abiotic-affinity designs, logic-based multiple stimulus combination, cell-selective traction force and matrix
metalloproteinase based-triggers overcome the issues of spatial selectivity, but often impose challenges including complex system designs, low cost-effectiveness, or rely on protein modifications of the GF. Approaches enabling spatial control over GF delivery via covalent immobilization, covalent tethering with stimuli-sensitive linkers (such as pH, enzymes, light), or protein modification, also have their own limitations. For example, covalent immobilization inhibits GF internalization that could inadvertently interfere with the biological processes. [23]. Besides, it is challenging to maintain conformations and/or biological activity after conjugation (or during the process) within polymer matrices. [23].

To address the aforementioned limitations and enable high specificity along with controlled therapeutic release, we sought to develop a versatile aptamer-based GF delivery platform that (i) is highly selective for rapid GF sequestration, (ii) enables on-demand triggered GF release via complementary sequence (CS) hybridization, (iii) decouples GF loading and hydrogel formation (avoiding harsh conditions or GF modifications), thus promising maximum bioactivity, (iv) is suitable for exploring 3D cell encapsulation (specifically, for tissue engineering applications), and (v) is capable of guiding cell behaviors such as survival, attachment, migration and proliferation, by spatiotemporally controlling GF bioavailability in 3D. In our rational approach, we set out to harness the unique capabilities of aptamers which are small, single-stranded oligonucleotides selected from synthetic RNA/DNA libraries. [24]. Depending on sequence, oligonucleotides form unique 3D conformations that enable selective binding to target biomolecules with high affinity and specificity, thus minimizing risk of non-specific release. [24]. While oligonucleotides are naturally susceptible to DNases degradation in vivo, approaches such as aptamer PEGylation and RNA-modified oligonucleotides have significantly reduced DNase-mediated degradation in vivo. [25]. Consequently, the clinically approved vascular endothelial growth factor (VEGF)-binding RNA-based aptamer, Pegaptanib sodium, which is used for macular degeneration treatment (Macugen, Eyetech/Pfizer), utilizes both PEGylation and oligonucleotide modification to achieve an in vivo half-life of 10 (±4) days. [25]. Moreover, aptamers can be designed for higher affinity to CS than the target GF resulting in aptamer-CS binding via intermolecular hybridization, thus enabling on-demand GF release using CS as an external trigger. [24]. It has been shown that biomaterials (or polymeric particles) functionalized with aptamers specific for different GFs (for example, VEGF and platelet-derived growth factor BB (PDGF-BB) or basic fibroblast growth factor) [26], [28]–[28] can sustain sequential release of multiple GFs via aptamer-CS hybridization.

Even though aptamer based biomaterials have been reported in literature, so far their bioactivity has been mainly studied through in vitro experiments. [29]. In this work, we developed a versatile aptamer-based GF delivery platform that is (i) highly selective for rapid GF sequestration, (ii) enables on-demand triggered GF release via complementary sequence (CS) hybridization, (iii) decouples GF loading and hydrogel formation (avoiding harsh conditions or GF modifications), thus promising maximum bioactivity, (iv) is suitable for exploring 3D cell encapsulation (specifically, for tissue engineering applications), and (v) is capable of guiding cell behaviors such as survival, attachment, migration and proliferation, by spatiotemporally controlling GF bioavailability in 3D. In our rational approach, we set out to harness the unique capabilities of aptamers which are small, single-stranded oligonucleotides selected from synthetic RNA/DNA libraries. [24]. Depending on sequence, oligonucleotides form unique 3D conformations that enable selective binding to target biomolecules with high affinity and specificity, thus minimizing risk of non-specific release. [24]. While oligonucleotides are naturally susceptible to DNases degradation in vivo, approaches such as aptamer PEGylation and RNA-modified oligonucleotides have significantly reduced DNase-mediated degradation in vivo. [25]. Consequently, the clinically approved vascular endothelial growth factor (VEGF)-binding RNA-based aptamer, Pegaptanib sodium, which is used for macular degeneration treatment (Macugen, Eyetech/Pfizer), utilizes both PEGylation and oligonucleotide modification to achieve an in vivo half-life of 10 (±4) days. [25]. Moreover, aptamers can be designed for higher affinity to CS than the target GF resulting in aptamer-CS binding via intermolecular hybridization, thus enabling on-demand GF release using CS as an external trigger. [24]. It has been shown that biomaterials (or polymeric particles) functionalized with aptamers specific for different GFs (for example, VEGF and platelet-derived growth factor BB (PDGF-BB) or basic fibroblast growth factor) [26], [28]–[28] can sustain sequential release of multiple GFs via aptamer-CS hybridization.

2. Materials and methods

2.1. Materials

Type A 300 bloom porcine skin gelatin (G1890-500G, Sigma Aldrich), Methacrylic anhydride (MA, 276685-500 ML, Sigma Aldrich), Dulbecco’s phosphate buffered saline (DPBS, D8537-500 ML, Sigma Aldrich), Fisherbrand™ regenereated cellulose dialysis tubing (12–14 kDa, 21-152-14, Fisher Scientific), bovine serum albumin (BSA, A9418, Sigma Aldrich), deuterium oxide (151890, Sigma-Aldrich), 2-hydroxy-4-(2-hydroxyethoxy)-2-methylpropionophenone (Irgacure 2959, 410896, Sigma Aldrich), VEGF specific control aptamer (47-nt, DNA, IDT), 5'-acryde modified aptamer (47-nt, DNA, IDT), complementary sequence (Comp. Seq., 46-nt, DNA, IDT), 5’Alexa Fluor 488 modified complementary sequence (Fluooro-Comp. Seq., 46-nt, DNA, IDT), nuclease free water (11-04-02-01, IDT), Fluoro-Max dyed blue aqueous fluorescent particles (2 μm diameter) (B0200, Thermo Scientific), human VEGF ELISA kit (RAB0507-1 KT, Sigma Aldrich), glutaraldehyde solution (340855, Sigma Aldrich), human umbilical vein endothelial cells (HUVECs, C2519A, Lonza), human mesenchymal stromal cells (MSC, PT-2501, Lonza), α –MEM medium (+nucleosides, 22571-020, Gibco), fetal bovine serum (FBS, 17552, Sigma), GlutaMax™ supplement (35050061, Gibco), penicillin-streptomycin (pen/strep, 15140-122, Gibco),漆-tarobic acid (A9860, Sigma Aldrich), trypsin-EDTA 0.25% (+phenol red, 25200072, Gibco), vascular endothelial growth factor 165 human (VEGF, H9166, Sigma Aldrich), Gibco™ FGABasic AA 1–155 recombinant human protein (bFGF, PHG0264, Fisher Scientific), endothelial cell basal medium 2 (EGM 2, C-22211, PromoCell), endothelial cell growth medium 2 supplement pack (EGM 2, C-39211, PromoCell), PDMS silicone elastomer (2401673921, Sylgard), live/dead cell double staining kit (04511, Sigma Aldrich), formaldehyde solution (FA775-25 ML, Sigma Aldrich), Triton X-100 (49409, Sigma Aldrich), Invitrogen™ AlexFluor™ 647 Phalloidin (A22827, Fisher Scientific), Invitrogen™ hoechst 33342 (H1399, Fisher Scientific), monoclonal anti-actin, α-smooth muscle antibody (mouse, A2547, Sigma Aldrich), anti-von Willebrand factor antibody (rabbit, F3520, Sigma Aldrich), human VE-Cadherin antibody (mouse, MAB9381, R&D systems), IgG (H + L) cross-adsorbed goat anti-rabbit, Alexa Fluor® 488, Invitrogen™ (A11008, Fisher Scientific), IgG (H + L) cross-adsorbed goat anti-mouse, Alexa Fluor® 488, Invitrogen™ (A11001, Fisher Scientific), IgG (H + L) highly cross-adsorbed goat anti-mouse, Alexa Fluor® 594, Invitrogen™ (A11032, Fisher Scientific), Alexa Fluor® 647 Anti-VEGFA antibody (EP1176Y) (ab206887, abcam), Corning® 96 well black polystyrene microplate (CLS3603, Sigma), Corning® Costar® ultra-low attachment well plates were used for all cell culture experiments (CLS3473, CLS3474, Sigma).

2.2. Prediction of secondary structures

The secondary structures of DNA based aptamers and their 3D conformations were generated using NUPACK. The presented structure with the lowest free energy at 37 °C was presumed to be the dominant structure (Fig. S1).

2.3. Synthesis and characterization of gelatin methacryloyl

Gelatin methacryloyl (GelMA) was synthesized as described previously elsewhere with medium degree of methacryloyl substitution (~60%). [29]. Briefly, gelatin was dissolved at 10% (w/v) into DPBS at 60 °C and stirred until fully dissolved. Subsequently, 1.25% (v/v) of MA was added at a rate of 0.5 mL/min to the gelatin solution at 50 °C for 1 h with continuous stirring. Afterwards, the reaction was stop with additional warm (40 °C) DPBS leading to 5% dilution. The solution was then dialyzed against distilled water using 12–14 kDa cutoff dialysis tubing at 40 °C for 1 week to remove residual salts and methacrylic acid. The solution was freeze-dried for 1 week and stored at -20 °C until further use. The degree of functionalization was quantified by H1 NMR. [29]. The H1 NMR spectra were collected at 35 °C in deuterium oxide at a frequency of 400 MHz using NMR spectrometer. The obtained spectral data was analyzed using MestReNova software (Fig. S2).
2.4. Synthesis of aptamer-functionalized hydrogels

The freeze-dried GelMA macromer was mixed with Irgacure 2959 as a photoinitiator in DPBS at 60 °C until fully dissolved. Afterwards, VEGF165 specific aptamers (control aptamer or acrydite aptamer, see Table S1) reconstituted in nuclease free water, was added into this solution with 5% (w/v) GelMA, 0.5% (w/v) Irgacure 2959 and 2.5 nmoles of aptamers in every 50 μl of pre-polymer solution. This pre-polymer solution was then used to prepare hydrogel samples for different experiments. Additionally, plain GelMA hydrogels were synthesized using same pre-polymer solution but without aptamer. For photo-crosslinking, requisite amount of pre-polymer solution was added into a PDMS mold, covered with a coverslip and exposed to 0.23 mW/cm² UV light (360–480 nm) (UV-KUB 2, Kloe, France) for 2 min. Samples were detached from the coverslips, transferred to their respective well plates using sterilized spatulas and used for different experiments.

2.5. Evaluation of aptamer retention within the hydrogels

To evaluate aptamer incorporation and their retention within the hydrogels, the aptamer-functionalized (acrydite-aptamer or control-aptamer) hydrogels and GelMA hydrogels were prepared by adding 25 μl of pre-polymer solution into PDMS mold (2 mm thickness and 4 mm diameter) following 2mins of UV crosslinking. Post photo-crosslinking, the hydrogels were transferred into clear-bottom black polystyrene 96-well plate with one hydrogel per well and were incubated. For photo-crosslinking, requisite amount of pre-polymer solution was added into a PDMS mold, covered with a coverslip and exposed to 0.23 mW/cm² UV light (360–480 nm) (UV-KUB 2, Kloe, France) for 2 min. Samples were detached from the coverslips, transferred to their respective well plates using sterilized spatulas and used for different experiments.

2.6. Analysis of VEGF165 sequestration and its triggered release

To conduct VEGF165 sequestration and triggered release experiments, the aptamer-functionalized hydrogels (2.5nmoles of acrydite-aptamer or control-aptamer) were prepared by adding 50 μl of pre-polymer solution into a PDMS mold (2 mm thickness and 6 mm diameter) following 2mins of UV crosslinking. Additionally, plain GelMA hydrogels without aptamers were prepared. Once crosslinked, the hydrogels were transferred to ultra-low attachment 24-well plate for 24hrs in 100 μl DPBS. A mole ratio of 1:1 for aptamer to CS was maintained. After 24hrs, the supernatant was discarded and the hydrogels were washed once with DPBS to remove unbound CS. To this end, 100 μl of fresh DPBS was added into the hydrogel and imaged using fluorescence microscope (EVOS M7000, Thermo Fisher Scientific) as day1 for the experiment. The experiment was carried out for 10days where after every 24hrs, the hydrogel’s supernatant was replaced with fresh 100 μl DPBS. The hydrogels were imaged at 60% intensity & 120 ms exposure time; 2x objective. The experiment was performed with three experimental replicates for each group. To quantify the fluorescence intensity, ImageJ (NIH, USA) software was used.

2.7. Scanning electron microscopy

For scanning electron microscopy (SEM) analysis, aptamer-functionalized hydrogels with different (acrydite- or control-) aptamer amounts (0.25nmoles, 2.5nmoles & 25nmoles) were prepared as previously discussed. The GelMA hydrogel without any aptamer was prepared as control. For preparing the hydrogel, 500 μl of pre-polymer solution was added into 24-well plate followed by 2mins of UV crosslinking. The hydrogel discs of about ~4 mm thickness and 15 mm diameter were obtained. The hydrogels were washed with DPBS followed by fixing in 2.5% glutaraldehyde for 24hr at 4 °C. Afterwards, the hydrogels were flash-frozened in liquid nitrogen and freeze-dried for 4days. The freeze-dried hydrogel discs were broken in liquid nitrogen to observe the cross-section, gold-sputtered (Sputter Coater 108 Auto, Cressington Scientific Instruments) and imaged using SEM (JSM-IT100, JEOL). Considering the large hydrogel disc size, three different regions from the same sample were imaged for pore size measurement within each group using ImageJ software.

2.8. Rheological analysis

The viscoelastic properties analysis of the aptamer-functionalized hydrogels with different (acrydite- or control-) aptamer amounts (0.25nmoles, 2.5nmoles & 25nmoles) were performed using parallel plate geometry (PP8, 8 mm) on a stress-controlled rheometer (Physica MCR301, Anton Paar). The parallel plate and bottom rheometer plate were treated with suitable sandpaper to prevent slipping. The GelMA hydrogel was prepared as control. Post UV crosslinking, hydrogel samples (8 mm diameter and 1 mm thickness) were placed onto the rheometer plate at ambient room temperature (20 °C) and the parallel plate was lowered to the desired gap height of 1 mm. A customized 3D-printed solvent trap was used to prevent evaporation during the rheological measurement. The amplitude sweep measurement at a constant 1 rad/s angular frequency (ω) was carried out over the range of 0.1% till 100% strain (γ) to examine viscoelastic properties of hydrogels. The storage modulus (G’) and loss modulus (G”) were determined as the strain (%) changes during oscillatory shear. The elastic modulus of hydrogel was determined from the linear region of amplitude sweep data.

2.9. Cell culture

The human umbilical vein-derived endothelial cells (HUVECs,
Lona) and human mesenchymal stromal cells (MSCs, Lonza) were cultured as per the standard protocol. In brief, HUVECs were cultured in EGM-2 medium (EGM-2 Basal medium + EGM-2 Supplements) with 1% (v/v) Pen/Strep. However, for MSCs, α-MEM medium supplemented with 10% (v/v) FBS, 2 mM l-glutamine, 0.2 mM ascorbic acid, 1% (v/v) Pen/Strep and 1 ng/ml BFGF was used. Both cell types were cultured in a humidified atmosphere with 5% CO₂ at 37 °C and passaged till 80% confluence.

2.10. 3D Co-culture in aptamer-functionalized hydrogels

For co-culture experiments, 1:1 ratio of HUVECs to MSCs with a total seeding density of 2.5 × 10⁶ cells ml⁻¹ was used throughout this study. Similarly, MSCs medium (α-MEM, 10% FBS, 2 mM l-glutamine, 0.2 mM ascorbic acid & 1% Pen/Strep) and HUVECs medium (EGM-2 Basal medium+ α-MEM+ Pen/Strep) were used. The cell suspension was then centrifuged at 300g for 3mins at 4 °C to get the cell pellet. The supernatant was removed and pre-polymer solution (5% GelMA+0.5% Irgacure 2959) was directly added and mixed gently by pipetting. 50 μl from this pre-polymer solution was added and dispensed into ultra-low attachment 96-well plate, followed by 2mins of UV crosslinking. The cells were trypsinized, counted (using trypan blue) and re-suspended in media (5% GelMA and 5% CS) were added.

The samples were then washed with DPBS, IgG (H + L) highly cross-adsorbed goat anti-mouse, Alexa Fluor® 594 (1:1000 in DPBS) and IgG (H + L) cross-adsorbed goat anti-rabbit, Alexa Fluor® 488 (1:1000 in DPBS) solutions were added for 2hrs in dark. Subsequently, the samples were washed with DPBS and incubated with Hoechst 33342 (1:2000 in DPBS) for 1hr at room temperature, followed by 10mins incubation with Hoechst 33342 (1:2000 in DPBS). For immunostainings, post blocking solution, the samples were washed with DPBS and incubated for overnight at 4 °C with monoclonal anti-actin, α-smooth muscle antibody (mouse, 1:300 in DPBS) and anti-β-actin antibody (rabbit, 1:200 in DPBS). The samples were then washed with DPBS, IgG (H + L) cross-adsorbed goat anti-mouse, Alexa Fluor® 594 (1:1000 in DPBS) and IgG (H + L) cross-adsorbed goat anti-rabbit, Alexa Fluor® 488 (1:1000 in DPBS) solutions were added for 2hrs in dark. Subsequently, the samples were washed with DPBS and incubated with Hoechst 33342 (1:2000) for 10mins in dark. For tight-junctions specific proteins staining, after blocking solution & DPBS washing, the samples were incubated for overnight at 4 °C with human VE-Cadherin antibody (mouse, 1:200 in DPBS). The samples were washed with DPBS and incubated with IgG (H + L) cross-adsorbed goat anti-mouse, Alexa Fluor® 488 (1:1000 in DPBS) for 2hrs in dark. After 1hr, Phalloidin 647 (1:40 in DPBS) was added to the secondary antibody staining solution and allowed to incubate for 1hr in dark. Thereafter, the samples were washed and incubated with Hoechst 33342 (1:2000) was added for 10mins in dark. Once stained, samples were washed with DPBS, 500 μl DPBS was added and stored at 4 °C until imaged.

2.11. Bi-phasic cell-laden hydrogels

To fabricate bi-phasic cell-laden hydrogels, autoclaved PDMS molds (3 mm thickness & 15 mm diameter) with a removable insert blocking half of the mold were used. To start with, HUVECs (P3) and MSCs (P4) were trypsinized, counted (using trypan blue) and re-suspended in 1:1 ratio with total seeding density of 2.5 × 10⁶ cells ml⁻¹. To prepare two different pre-polymer solutions, the cell suspension with same seeding density was centrifuged at 300g for 3mins at 4 °C in two separate 15 ml falcon tubes. For pre-polymer solution 1, supernatant was removed and pre-polymer solution (5% GelMA+0.5% Irgacure 2959 + 2.5mnoles acrydite- or control-aptamer in DPBS) was added and mixed gently by pipetting. Similarly, for pre-polymer solution 2, supernatant was removed and 5% GelMA+0.5% Irgacure 2959 in DPBS solution was added. Afterwards, PDMS molds with an insert blocking half of the area were placed in sterile cell culture petri dish (90 mm diameter) and 200 μl of pre-polymer solution 1 was added. The mold was covered with coverslip and exposed for UV crosslinking (1min, 0.23 mW/cm² intensity). After the crosslinking of one side, the PDMS mold insert and coverslip were carefully removed and pre-polymer solution 2 was added in the rest of the mold followed by UV crosslinking of whole sample (1min, 0.23 mW/cm² intensity). Thereafter, the coverslip and PDMS molds were removed and bi-phasic cell-laden hydrogels were transferred to ultra-low attachment 24-well plate using sterilized spatula.

Herein, for VEGF₁₆₅ loading, 1 ml of co-culture medium with VEGF₁₆₅ (10 ng) was added into each sample and incubated at 37 °C for 1hr. Subsequently, the medium was replaced with fresh 1 ml co-culture medium. The medium was changed once every day throughout the experiment. To evaluate the effect of triggered VEGF₁₆₅ release on bi-phasic cell-laden hydrogels, 2.5mnoles CS (1:1 mol ratio of aptamer to CS) into 1 ml co-culture medium was added for acrydite- and control-aptamer based bi-phasic hydrogels on day4 for 24hrs. These samples were analyzed at different time-points. For vessel properties quantification, each sample was categorized into four regions; near aptamer- and near GelMA-side being in immediate vicinity of the interface (< 600 μm; interface at 0 μm), whereas far aptamer- and far GelMA-side were at the far end from the interface (between 2000 to 4000 μm from interface).

2.12. Immunostainings

Prior to staining, all of the samples were fixed using 4% formaldehyde solution in DPBS for 30mins, washed with DPBS and permeabilized using 0.1% Triton X-100 in DPBS. Afterwards, the samples were washed with DPBS and blocking solution of 1% FBS in DPBS was added for 45mins. For actin cytoskeleton staining of bi-phasic cell-laden hydrogels, the samples were incubated with Phalloidin-Atto 488 (1:50 in DPBS) or Alexa Fluor-647 Phalloidin (1:40 in DPBS) for 1hr at room temperature, followed by 10mins incubation with Hoechst 33342 (1:2000 in DPBS). For immunostainings, post blocking solution, the samples were washed with DPBS and incubated for overnight at 4 °C with monoclonal anti-actin, α-smooth muscle antibody (mouse, 1:300 in DPBS) and anti-von Willebrand factor antibody (rabbit, 1:200 in DPBS). The samples were then washed with DPBS, IgG (H + L) highly cross-adsorbed goat anti-mouse, Alexa Fluor® 594 (1:1000 in DPBS) and IgG (H + L) cross-adsorbed goat anti-rabbit, Alexa Fluor® 488 (1:1000 in DPBS) solutions were added for 2hrs in dark. Subsequently, the samples were washed with DPBS and incubated with Hoechst 33342 (1:2000) for 10mins in dark. For tight-junctions specific proteins staining, after blocking solution & DPBS washing, the samples were incubated for overnight at 4 °C with human VE-Cadherin antibody (mouse, 1:200 in DPBS). The samples were washed with DPBS and incubated with IgG (H + L) cross-adsorbed goat anti-mouse, Alexa Fluor® 488 (1:1000 in DPBS) for 2hrs in dark. After 1hr, Phalloidin 647 (1:40 in DPBS) was added to the secondary antibody staining solution and allowed to incubate for 1hr in dark. Thereafter, the samples were washed and incubated with Hoechst 33342 (1:2000) was added for 10mins in dark. Once stained, samples were washed with DPBS, 500 μl DPBS was added and stored at 4 °C until imaged.

2.13. Statistical analysis

Statistical analysis was performed using GraphPad Prism 7 software. Two-way analysis of variance (ANOVA) with Tukey’s multiple comparisons test was used to analyze the data. Significance was set at p < 0.05. The data is represented as mean ± SD.

3. Results and discussion

3.1. Acrydite-modified aptamers enhance aptamer retention and CS recognition within aptamer-functionalized hydrogels

To develop the desired versatile platform, we synthesized photo-crosslinked aptamer-functionalized gelatin methacryloyl (GeMA) hydrogels via free-radical polymerization initiated with UV light
exposure (Fig. 1A). The oligonucleotide sequence specific for VEGF<sub>165</sub> [30,31], was utilized in both unmodified form (hereafter referred as “control-aptamer”) and 5′-acrydite modified form (hereafter referred as “acrydite-aptamer”) (Fig. S1 and Table S1). Upon initiating the free-radical polymerization, the unsaturated bonds present in acrydite groups enable covalent aptamer incorporation within the polymer network. [32,33]. The methacrylation degree and monomer concentration are critical parameters governing the polymer crosslinking density, that in-turn affects the hydrogel porosity, its pore size and mechanical properties. [34,35]. Specifically, GelMA hydrogels with high methacrylation degree have often showed low porosity and high elastic modulus. [34]. As high porosity supports mass transport to cells and suitable pore size allows cell ingrowth, both of which are favorable conditions for 3D culture [34], GelMA with a medium methacrylation degree (~60%) was utilized for the fabrication of aptamer-functionalized hydrogels throughout the study (Fig. S2).

To demonstrate the versatility of the platform, we first set out to verify aptamer retention and their ability for CS recognition within the hydrogels by using 5′-Alexa Fluor 488 fluorophore labelled CS (Fluoro-CS) at physiological temperature (i.e., 37 °C) (Fig. 1B and C). The acrydite-aptamer-functionalized hydrogels exhibited higher fluorescence throughout the study, confirming stable aptamer incorporation into the hydrogel network (Fig. 1C). The control-aptamer-functionalized hydrogels showed lower fluorescence after the initial 24hrs, followed by gradual decrease of the fluorescent signal over time. Without acrydite modification, the control-aptamers tend to physically entrap within polymer network during polymerization, which enables gradual aptamer diffusion out of the polymer network. As expected, the plain GelMA hydrogels (without aptamer) showed no detectable fluorescence after 48hrs (Fig. 1C). Furthermore, to evaluate the presence of the acrydite-aptamer and Fluoro-CS diffusion throughout the acrydite-aptamers hydrogel thickness, bi-phasic hydrogels (having one side with acrydite-aptamer and one side without aptamer; 3 mm thickness & 15 mm diameter) were imaged using confocal microscopy after 24hrs incubation with Fluoro-CS at 37 °C. Three-dimensional projections of confocal z-stacks confirms the CS-Fluoro penetration (z = 240 μm) through the hydrogel thickness while a clear interface with GelMA region (in blue color) is maintained (Fig. S3 & Video S1). Combined, these observations suggest that Fluoro-CS was able to penetrate within the (acrydite- & control-) aptamer-functionalized hydrogels at macroscopic scale within a timeframe of 24hr incubation (irrespective of aptamer being covalently-bound or physically entrapped) and displayed high molecular recognition via aptamer-Fluoro-CS hybridization at physiological temperature.

![Fig. 1.](image_url)

Fig. 1. The concept of aptamer-functionalized hydrogels and aptamer retention within gelatin methacryloyl (GelMA) hydrogels. (A) Schematic representation of photo-crosslinked GelMA with 5′-acrydite-modified VEGF<sub>165</sub> specific aptamer sequence, in the presence of photoinitiator (Irgacure 2959) and UV light. (B) Schematics explaining the concept of 5′-acrydite-modified aptamers being covalently linked into GelMA network, whereas, due to the absence of 5′-acrydite group (red circle), the control-aptamers are freely diffusible within the polymer network. Due to molecular recognition, upon addition of 5′-fluorescently labelled complementary sequence (Fluoro-CS) in the 5′-acrydite-aptamer-functionalized hydrogels (Acrydite Apmatmers) and control-aptamer-functionalized hydrogels (Control Apmatmers), the Fluoro-CS-aptamer hybridization occurs. (C) The fluorescence microscopic images of the GelMA, “Control Apmater” and “Acrydite Apmater”-functionalized hydrogels after 24hrs incubation with Fluoro-CS at 37 °C. #D10 indicates the brightfield images of hydrogels on day10. The fluorescence intensity corresponding to the presence of Fluoro-CS indicates the presence of aptamers within the hydrogels at various time-points. The red dotted line in GelMA hydrogels from day2 onwards shows the hydrogel border. The mean fluorescence of all samples over different time-points have been shown in the graph. The experiment was performed with n = 3 experimental replicates and values are represented as mean ± SD.
3.2. Aptamers crosslinked within hydrogel matrix mediates maximum VEGF<sub>165</sub> sequestration and programmable release via CS hybridization

Having verified the aptamer retention and its ability for CS hybridization within aptamer-functionalized hydrogels, we next set out to validate their functionality for VEGF<sub>165</sub> sequestration and programmable, triggered VEGF<sub>165</sub> release via aptamer-CS hybridization. Firstly, for verifying the rapid VEGF<sub>165</sub> sequestration ability and subsequent GF release over time, the hydrogels were incubated with VEGF<sub>165</sub> (10 ng) containing loading medium for 1 hr (Fig. 2A and B). The results indicated higher VEGF<sub>165</sub> sequestration within aptamer-functionalized hydrogels even after 1 hr incubation (50% and 46% for control-aptamer and acrydite-aptamer, respectively) compared to GelMA hydrogel (33%) (Fig. S4). To further visualize the differential sequestration of VEGF<sub>165</sub> within hydrogels, bi-phasic hydrogels (having one side with acrydite-aptamer and one side without aptamer; 3 mm thickness & 15 mm diameter) loaded with VEGF<sub>165</sub> for 1 hr were immunostained using anti-VEGF antibody. The confocal microscopic images showed increased levels of VEGF<sub>165</sub> sequestration (in red color) within acrydite-aptamer region compared to the GelMA region (in blue color) (Fig. S5). Gelatin-based hydrogels display inherent GF sequestration properties because of their electrostatic interactions with oppositely charged GFs like VEGF, which can explain the relatively high sequestration for GelMA hydrogels. As expected, the increased sequestration in aptamer-functionalized hydrogels is likely due to the binding affinity of GF with aptamer. The insignificant difference in VEGF<sub>165</sub> sequestration among aptamer-functionalized hydrogels confirm the similar GF sequestration ability of the used aptamer, irrespective of being covalently crosslinked or freely diffusible within the polymeric network. However, as shown in Fig. 1C, the freely diffusible behavior of control-aptamer becomes prevalent within the 24 hrs of incubation that could result into the initial burst release of VEGF<sub>165</sub>. Notably, the present GF loading route decouples the loading process from the hydrogel fabrication process, thus providing a clear advantage over conventional GF loading strategies, where free-radicals generated during hydrogel preparation could potentially harm the GF bioactivity. In a tissue engineering context, the present approach provides great flexibility over GF loading at different time-points after hydrogel fabrication, which is adaptable according to the needs of developing engineered tissue.

To explore the dynamic nature of this platform, we next examined the triggered GF release behavior from aptamer-functionalized hydrogels by supplementing CS (1:1 aptamer to CS mole ratio) into the releasing medium at different time-points (Fig. 2C). As expected, the control-aptamer-functionalized and GelMA hydrogels showed high burst release of VEGF<sub>165</sub> on day1 (10.99 pg and 18.89 pg, respectively) and near zero release from day2 onwards. In contrast, acrydite-aptamer-functionalized hydrogels exhibited minimal initial release on day1 (3.30 pg), confirming significantly reduced initial burst release and efficient VEGF<sub>165</sub> sequestration within the hydrogel. Upon the first round of triggered release with CS addition on day4, the acrydite-aptamer-functionalized hydrogels showed high VEGF<sub>165</sub> release within 24 hr (20.81 pg), compared to GelMA and control-aptamer-functionalized hydrogels (2.62 pg and 0.32 pg, respectively). For testing the platform’s functionality at later time-point, the second triggered release with CS addition on day9 was attempted with all samples. Interestingly, the acrydite-aptamer- and control-aptamer-functionalized hydrogels that were already triggered on day4 showed no additional release. The acrydite-aptamer-functionalized hydrogel triggered only on day9 showed high VEGF<sub>165</sub> release on day10 (25 pg) (Fig. 2C). This indicates that CS used in present study was able to regulate the binding functionality of aptamers and induce rapid dissociation of aptamer-protein complexes to trigger VEGF<sub>165</sub> release within acrydite-aptamer samples. The used aptamer to CS mole ratio (1:1) showed a maximum release of 25 pg at once, followed by no subsequent release upon second CS triggering indicating its saturation. However, it is speculated that the triggered VEGF<sub>165</sub> release can be modulated by varying triggering time and CS mole ratio. This proof-of-concept data shows the rapid sequestration (within 1 hr) and on-demand VEGF<sub>165</sub> releasing capability of acrydite-aptamer-functionalized hydrogels. The cumulative VEGF<sub>165</sub> release data confirms overall higher release from acrydite-aptamer-functionalized and GelMA hydrogels (25–29 pg), than control-aptamer-functionalized hydrogels (12–14 pg) (Fig. 2C). Considering control-aptamers not being covalently crosslinked within the polymeric network, as confirmed by retention data (Fig. 1C), a portion of VEGF<sub>165</sub> molecules diffusing into the releasing medium is likely to be bounded with control-aptamers via affinity interaction. Upon investigating the sensitivity of the used ELISA assay for detecting VEGF<sub>165</sub> molecules complemented with control-aptamers, lower signals (with overall difference of 28%) were detected for VEGF<sub>165</sub> incubated with control-aptamers compared to VEGF<sub>165</sub> in DBPS (Fig. S6). It is anticipated that this disparity in ELISA sensitivity towards VEGF<sub>165</sub> molecules in the presence of control-aptamers, could be a possible reason for the overall lower VEGF<sub>165</sub> release values from control-aptamer-functionalized hydrogels. It should further be noted that a cumulative release of approximately 30 pg constitutes only a small fraction of the 10 ng that was loaded per sample, from which up to half was sequestered into the sample as determined by analysis of the remaining VEGF<sub>165</sub> in the loading fluid. Even though this could indicate that the major fraction of loaded VEGF<sub>165</sub> remains in the samples after 10 days of release, this is unlikely given the low level of staining for VEGF<sub>165</sub> in the biphasic system in Fig. S5. If the binding characteristics of unmodified GelMA would be strong enough to retain the major fraction of VEGF<sub>165</sub> over the course of 10 days, the differential effect of binding to the aptamer would be small, resulting in only a small or undetectable difference in the staining. It remains unclear what has caused this discrepancy, but as the VEGF<sub>165</sub> sequestration and release experiments were performed separately, variations among individual samples used in the experiments cannot be ruled out. It is speculated that by modulating aptamer’s binding affinity and mole ratio, an improved GF retention capacity can be achieved. Aptamer-functionalized superporous hydrogels with high binding affinity have been previously shown to retain up to 90% of the GF (PDGF-BB) after 24 hr incubation with washing solution, compared to the non-functionalized (14%), low binding affinity (28%) and medium binding affinity aptamer-functionalized hydrogels (63%), respectively. [33]. Similarly, higher GF retention was reported with increased mole ratio of aptamer to GF such as 53% (1:1 mol ratio), 90% (1:0.1 mol ratio) and 96% (10:1 mol ratio), respectively. [33]. In the previously reported study, small volumes of highly concentrated GF solution were directly loaded onto partially dehydrated superporous hydrogels which were then incubated with the GF solution for 24 hr. [33]. The present study on the other hand employs GF sequestration from a large volume of medium supplemented with a more physiological concentration of GF during 1 hr of incubation. Even though the previously employed approach is likely to result in more efficient GF loading of the aptamer functionalized hydrogel, the presented approach is better compatible with patterned hydrogels or potential applications where the GF is sequestered in vivo. Taken together, these results confirmed acrydite-aptamer functionalized hydrogel’s potential for rapid (within 1 hr loading) and localized (confined within aptamer region) VEGF<sub>165</sub> sequestration from the culture medium as well as their capability for triggering VEGF<sub>165</sub> release via aptamer-CS hybridization.

3.3. Aptamer incorporation influences physicochemical properties of aptamer-functionalized hydrogels

To understand the effect of aptamer incorporation in GelMA hydrogels on their overall physicochemical properties, aptamer-functionalized hydrogels with varying aptamer amounts were assessed (Fig. 3). The scanning electron micrographs (SEM) data of acrydite-aptamer-functionalized hydrogels showed significant aptamer amount dependent pore properties. For example, low aptamer (0.25nmoles) resulted in an increase of the average pore area compared to GelMA
Fig. 2. Temporally controlled VEGF$_{165}$ release from the aptamer-functionalized hydrogels. (A) Schematic representation of the aptamer-functionalized hydrogel fabrication, its ability to sequester VEGF$_{165}$ from the release media via affinity interactions and on-demand triggered release of VEGF$_{165}$ in presence of complementary sequences (CS). (B) The complete timeline of the experiment. The aptamer amount used was 2.5 nmoles (25 μM) and initial VEGF$_{165}$ loading amount was 10 ng. (C) The daily (top) and cumulative (bottom) VEGF$_{165}$ release from control-aptamer- and acrydite-aptamer-functionalized hydrogels over 10 days in presence or absence of CS. To trigger the first round of VEGF$_{165}$ release on day5, the CS was added to specified hydrogels on day4 for 24hrs (as indicated in figure). Furthermore, to trigger the second round of VEGF$_{165}$ release, the CS was added to all aptamer hydrogel samples on day9, except for GelMA hydrogels. The mole ratio of CS to aptamers used was 1:1. The graphs are represented as mean with individual data points. The experiment was performed with three experimental replicates, n = 3.
samples (13,320 μm² compared to 6079 μm² respectively). An increase in the concentration of acrydite-aptamer resulted in denser polymeric crosslinking, which is evident by increased wall thickness and decreased pore size for 2.5nmoles samples (~13 μm and 1,462 μm²) and collapsing pores in 25nmoles samples accompanying higher wall thickness and larger remaining pores (~35 μm and 15,551 μm²). Control-aptamer-functionalized hydrogels on the other hand showed similar pore properties as GelMA hydrogels for all tested aptamer amounts (for example,

Fig. 3. Pore properties of aptamer-functionalized hydrogels with various aptamer amounts. SEM images of the cross-section of acrydite-aptamer- and control-aptamer-functionalized hydrogels with different aptamer amounts, namely 0.25 nmoles, 2.5 nmoles and 25 nmoles aptamers per sample. The GelMA hydrogels without aptamers were also used. Scale bar is 100 μm and magnification is 200x. Pore area and pore wall thickness of different hydrogel samples quantified by ImageJ using the representative SEM images. The data is represented as mean ± S.D. The statistical significance among pore area and pore wall thickness within each sample type was calculated using two-way ANOVA with Tukey’s multiple comparisons test where **** means p < 0.0001.

Fig. 4. Rheological analysis of aptamer-functionalized hydrogels with various aptamer amounts, namely 0.25 nmoles, 2.5 nmoles and 25 nmoles aptamers per sample. (A) control-aptamer-functionalized and (B) acrydite-aptamer-functionalized hydrogels, along with GelMA hydrogels. The storage modulus (filled symbols) and loss modulus (empty symbols) of the respective hydrogels at fixed angular frequency of 1 rad/s are shown.
an average pore area within 6000–6,900 μm² range). It is noteworthy that pore size determined from SEM images of lyophilized hydrogels, does not represent the hydrogel ultrastructure in a hydrated state that is experienced by encapsulated cells. Instead, the observed porosity is a result of the freezing-lyophilization process and represents the relative density of the hydrated hydrogels for comparison.

In correlation with SEM data, the rheology results revealed higher storage modulus for acrydite-aptamer-functionalized hydrogels (~450Pa) compared to GelMA hydrogel (404Pa) (Fig. 4). This effect was independent of the aptamer amount present within the hydrogel. The control-aptamer-functionalized hydrogels exhibited a slightly lower storage modulus for the lowest tested aptamer amount (~379Pa for 0.25nmoles). The storage modulus increased upon further increasing the aptamer amounts (~430Pa for 2.5nmoles; ~509Pa for 25nmoles). While the exact reason for these changes in mechanical properties of the hydrogels is unclear, one possibility may be linked to the hydrophilic nature of aptamers resulting in additional swelling of hydrogels when present in higher amounts, and thus contributing towards a higher storage modulus. This behavior could be anticipated for control-aptamers that are physically entrapped within hydrogel network, whereas covalently crosslinked acrydite-aptamers in the amounts used in this study are less likely to be available for participating in swelling, mainly because of the steric hindrances caused due to the close proximity of covalently immobilized acrydite-aptamers with GelMA polymeric network.

3.4. Bioactivity of acrydite-aptamer functionalized cell-laden hydrogels post-VEGF₁₆₅ sequestration and its triggered release via CS hybridization

ECM is a highly dynamic microenvironment consisting of multi-functional components (for example, proteoglycans and glycoproteins) that are capable of promoting cell adhesion as well as controlling GF presentation to cells by sequestering soluble GFs and releasing them via enzymatic degradation. Both soluble and ECM-bound GFs facilitate the spatiotemporal regulation of cellular responses. While many strategies have been developed to recapitulate the critical complexities of native ECM, the use of CS for triggering GF release, along with unique properties of aptamers and GelMA, provide this platform with the ability to precisely tune GF bioavailability within hydrogel microenvironment to probe and direct encapsulated cell responses in 4 dimension (4D). To this end, we next demonstrated the bioactivity of the platform throughout the process of VEGF₁₆₅ loading and triggered CS mediated release; co-cultures of human umbilical vein endothelial cells (HUVECs) and human mesenchymal stromal cells (MSCs) were used as model cell

![Cell viability of the 3D co-cultured aptamer-functionalized hydrogels with or without VEGF₁₆₅ loading. Live/Dead stained fluorescent microscopic images of 3D cultured hydrogels with control-/acrydite-aptamer-functionalized hydrogels in the presence or absence of CS addition on day4, on different time points. The HUVECs and MSCs were co-cultured in 1:1 ratio. For VEGF₁₆₅ loading, the hydrogels were incubated with co-culture medium (without VEGF₁₆₅ supplement) added with known VEGF₁₆₅ amount (10 ng), for 1hr at 37 °C. Plain GelMA hydrogels, with or without 1hr VEGF₁₆₅ loading were also studied. The scale bar is 200 μm. Cell viability % quantification was performed using ImageJ software. The quantification was performed with three experimental replicates, n = 3. The statistical significance was calculated using two-way ANOVA with Bonferroni’s multiple comparisons test where ***p = 0.0005, ****p < 0.0001 and ns stands for not significant.](image-url)
systems and were subjected to 3D cell encapsulation for fabricating cell-laden aptamer-functionalized hydrogels. These proof-of-concept experiments were designed to incubate cell-laden samples with VEGF_{165} in co-culture medium for 1hr to enable rapid GF sequestration, followed by washing and medium replacement. Additionally, VEGF_{165} release was triggered by CS addition into the culture medium for 24hrs on day4 of culture. To ensure that the observed cell behavior is mainly due to the bioavailability of externally loaded GF, no VEGF supplements were added into the co-culture medium after the initial GF loading step. Regardless of CS addition, high cell viability (>90%) in acrydite-aptamer-functionalized and GelMA hydrogels on day1 and day5 were observed (Fig. 5). However, in control-aptamer-functionalized hydrogels relatively lower cell viability (~70%) on day1 was observed that significantly increased (~90%) by day5, irrespective of CS addition. The initial lower viability in control-aptamer samples remains unclear, but could be related to the cells preference for matrix bound GFs as opposed to soluble ones. [39].

3.5. Spatially patterned aptamers within bi-phasic hydrogels selectively guide cell behavior within 3D microenvironment

Following the successful bioactivity confirmation of cell-laden acrydite-aptamer-functionalized hydrogels, we next aimed to demonstrate their fundamentally unique capabilities compared to the existing GF releasing strategies. We hypothesized that by spatially controlling local GF bioavailability using this platform, we could selectively guide cell behavior in 3D microenvironment. To test this, we designed bi-phasic cell-laden hydrogels with (control- or acrydite-) aptamer-functionalized hydrogels on one-side and plain GelMA on the other side via two-step photocrosslinking. These bi-phasic hydrogels were subsequently loaded with VEGF_{165} (10 ng) for 1hr. Microscopy data of F-actin stained samples confirmed the ability of the encapsulated cells to sense the localized presence of acrydite-aptamer-bound-VEGF_{165} within the hydrogels, resulting in increased cell adherence and spreading within the aptamer regions on day3 (Fig. 6). For quantification, the images with F-actin staining were processed via thresholding, binary and watershed segmentation tools in ImageJ software. The results indicate higher average cell area (2569 μm^2) and aspect ratios in acrydite-aptamer side than the GelMA side (542 μm^2). In control-aptamer samples, both sides displayed similar cell responses (average cell area ~3133 μm^2 & 2220 μm^2; aspect ratio ~2.8 & 2.9). This limited effect in the control-aptamer samples is likely to be attributed to the freely diffusible nature of control-aptamer-bound-VEGF_{165} molecules within the hydrogel matrix, correlating with our previous aptamer retention (Fig. 1C) and VEGF quantification data (Fig. 2C). Due to the bi-phasic nature of the samples, with both sides being exposed to the same pool of culture medium, it is likely that control-aptamer-bound-VEGF_{165} molecules diffuse out of the aptamer hydrogel region, thus being accessible to cells encapsulated within both regions. On the other hand, as acrydite-aptamers are covalently immobilized, this confines the acrydite-aptamer-bound-VEGF_{165} bioavailability within one region. When comparing the aptamer containing regions of control-aptamer samples with acrydite-aptamer samples, higher cell aspect ratios are seen in the acrydite-aptamer samples. Even though the exact mechanisms behind this are unclear, the increased cellular response within acrydite-aptamer regions could be attributed to the ability of matrix-bound VEGF to elicit more pronounced cell responses compared to soluble VEGF. [39]. It has previously been shown that matrix-bound VEGF induces its receptor (VEGFR2)-β1-integrin complex formation and increases the β1-integrin targeting to focal adhesions. [39].

![Fig. 6. Bi-phasic cell-laden acrydite-aptamer-functionalized hydrogels generate selectively guided cell responses on day3. The maximum projection confocal images of the HUVECs and MSCs, co-cultured within VEGF_{165} loaded (A) control-aptamer- and (B) acrydite-aptamer-functionalized bi-phasic hydrogels. For the ease of understanding, each bi-phasic hydrogel was categorized into two regions; aptamer side and GelMA side where the white dashed line indicates the interface. The cells were stained with cytoskeletal F-actin Phalloidin (green) and Hoechst (blue). The scale bar is 100 μm. Quantification of the individual cell area plotted against the distance from the interface within the (A) control-aptamer- and (B) acrydite-aptamer-functionalized bi-phasic hydrogels. The quantification was performed using ImageJ software, where values are represented in scatter plot with individual data points along with overall mean ± S.D. The calculations were performed with three technical replicates, n = 3. The statistical significance between aptamer and GelMA sides were calculated using unpaired two-tailed t-test with Welch’s correction where **p = 0.0004, ***p < 0.0001 and ns stands for not significant.](image-url)
3.6. Spatiotemporally controlled VEGF₁₆₅ bioavailability within acrydite-aptamer bi-phasic hydrogels manipulates microvascular network organization in 3D microenvironment

VEGF plays a vital role in early regulation of angiogenesis. Spatiotemporally controlling its bioavailability within engineered matrices (hydrogel) could be a valuable tool to direct the tubular morphogenesis of endothelial cells for achieving mature and stable vascular networks. Having established the effect of aptamer-functionalized bi-phasic hydrogels on cellular adherence and spreading, we next investigated its capacity for translating this effect with temporal control in guiding vascular morphogenesis. To do so, the bi-phasic samples were triggered with CS for VEGF₁₆₅ release at different time-points and the effect of this on microvascular network organization within biphasic samples was studied. The microscopy data confirms positive expression of von Willebrand factor (vWF, endothelial cell specific marker) and α-smooth muscle actin (α-SMA, smooth muscle cells specific marker) (Figs. 7 and 8). In agreement with previous research, the interaction between endothelial cells and MSCs in co-cultures directs MSCs differentiation towards the smooth muscle cell lineage, indicating vascular network stabilization and maturation. [40]. Furthermore, the microscopic images in orthogonal view revealed the presence of circular/elliptical lumen-like structures within vWF + networks wrapped by α-SMA + cells, mimicking the early tubular morphogenesis of endothelial cells in native tissues (Figs. 7B and 8B & S7 and Video S2). VE-cadherin, which is an endothelial intercellular junctional protein, plays a substantial role in endothelial cell tubular morphogenesis, vascular stability and maturation. The positive expression of VE-cadherin in the developing vascular networks is an early marker for vessel stabilization and maturation. As expected, VE-cadherin expression increased over time (Figs. 7C and 8C).

Quantification of vascular organization revealed that triggered VEGF₁₆₅ release (via CS addition) could spatiotemporally regulate vessel properties within acrydite-aptamer and GelMA regions (Fig. 7A,D&E) (Fig. S8, video S3). Upon comparing the network parameters among acrydite-aptamer samples treated with/without CS, the temporal control over vascular network organization triggered via CS addition becomes apparent. For instance, samples without CS addition showed higher vessel density in near aptamer region (region close to the interface on aptamer side; 4.99%) compared to near GelMA (region close to the interface on GelMA side; 2.21%) (p = 0.0421). Herein, the vessel density is defined as the percentage of area occupied by vWF + vessels within the total area (i.e., %vessels/total area). However, after 24hr CS treatment, this difference became insignificant (Fig. 7D). Even though the vessel density became indistinguishable across the interface by day10 (independent of CS addition), pronounced vessel density was observed specifically in near aptamer region for samples treated with CS (15.94%) compared to samples without CS (4.57%) (p < 0.001) (Fig. 7E). Similar trends were observed in near GelMA regions with higher vessel density in CS treated samples (12.30%) compared to samples that were not treated with CS (2.61%) (p = 0.004) by day10 (Fig. 7E). Evaluation of other vessel parameters such as branching density (i.e., number of vessel

**Fig. 7.** Spatiotemporally controlled vascular network organization within acrydite-aptamer-functionalized bi-phasic cell-laden hydrogels. (A) Immunostained maximum projection confocal images showing von Willebrand factor (vWF, endothelial cell specific marker) and α-smooth muscle actin (α-SMA, smooth muscle cells specific marker) (red) as a marker for MSCs differentiation to smooth muscle cells and vWF (green) as a marker for endothelial cells. (B) Acrydite aptamer bi-phasic samples treated with CS on day 4. (C) Vessel density quantification in acrydite-aptamer samples treated with/without CS, the temporal control over vascular network organization triggered via CS addition becomes apparent. For instance, samples without CS addition showed higher vessel density in near aptamer region (region close to the interface on aptamer side; 4.99%) compared to samples without CS (4.57%) (p < 0.001) (Fig. 7E). Similar trends were observed in near GelMA regions with higher vessel density in CS treated samples (12.30%) compared to samples that were not treated with CS (2.61%) (p = 0.004) by day10 (Fig. 7E). Evaluation of other vessel parameters such as branching density (i.e., number of vessel

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81
control-aptamer-bound-VEGF and average vessel length (mean length of all the vessels in image) corroborated this trend. On day5, lower average vessel length in near aptamer regions were observed for samples without CS treatment (0.0513 mm) compared to samples with CS (0.035 mm), but by day10, samples with CS treatment showed a pronounced increase in average vessel length (0.211 mm) compared to samples without CS (0.058 mm) (Fig. 7D&E). As expected, in control-aptamer samples significantly increased vessel network formation was observed when comparing CS treated samples with samples without CS on day5 and day10 (Fig. S9 & video S4). However, no significant differences were observed across the treated samples with samples without CS on day5 and day10 (Fig. S9). However, upon CS addition no significant differences were observed. The overall increased vessel properties in CS-treated samples (control-aptamer + CS) at different time-points (day5, 10), confirms the higher bioactivity of the released GF via CS hybridization and spatially controlled vascular organization in acrydite-aptamer samples across the interface by day5, shows the platform’s potential for regulating vascular morphogenesis in vitro by spatiotemporally controlling GF bioavailability.

The establishment of optimally organized vascular networks is currently a major hurdle for clinical translation of engineered tissues at large-scale. While many strategies have previously shown success with pre-vascularized scaffolds or cell patterning, they often overlook to account for vascular remodeling, thus hampering the functionality of these networks. Given the mobile nature of encapsulated endothelial cells within scaffolds, the initially developed vascular networks tend to remodel after implantation. At this stage, using platforms that provide

Fig. 8. Spatiotemporally controlled vascular network organization within control-aptamer-functionalized bi-phasic cell-laden hydrogels. (A) Immunostained maximum projection confocal images showing von Willebrand factor (vWF) expression (green) as a marker for endothelial cells and α-smooth muscle actin (α-SMA) (red) as a marker for MSCs differentiation to mural cells, within the HUVECs and MSCs co-cultured, bi-phasic hydrogels (with or without CS treatment on day4). The representative photographic image of the actual bi-phasic hydrogel, showing two distinct sides (one with aptamer and other one without) with a distinct interface. Considering its big size, each hydrogel was categorized into four regions; near aptamer- and far GelMA-side being in immediate vicinity of the interface, whereas far aptamer- and far GelMA-side were at the far end from the interface. Scale bar is 200 μm. Orthogonal views of the confocal z-stacks showing (B) vWF + α-SMA stained samples & (C) F-actin Phalloidin + VE-Cadherin stained samples, at higher magnification, showing the developing vascular networks. At the cross-section of the developing vessel, a round lumen like vascular structure could be observed in orthogonal view. The scale bar is 50 μm. Quantification of vWF + stained vessel network in these samples using Angiotool Software on day5 (D) and day10 (E). The values are represented as mean ± SD, along with individual data points. The calculations were performed with three technical replicates, n = 3. The statistical significance was calculated using two-way ANOVA with tukey’s post-hoc test were *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 and ns means not significant.
additional cues to guide the vascular remodeling processes, can enable the sustainment of well-organized vascular networks for longer durations. The cell culture data in this study confirm the platform’s ability to not only form lumen-like microvascular networks within a hydrogel matrix, but also to enable spatiotemporally controlled network organization. These unique properties could possibly help to better control vascular network remodeling processes in vivo.

In addition to demonstrating their ability for selectively directing cell responses by controlling GF bioavailability, this study also establishes that (i) the acrylde-aptamer-functionalized hydrogels were stable, exhibiting minimum VEGF leakage and (ii) the hydrogels display high bioactivity of sequestered VEGF in situ, for at least ten days in culture. Although GelMA degradation could be a limiting factor affecting GF release in studies involving longer culture periods, this could be addressed by modulating polymer concentration, methacyration degree and crosslinking density. [41]. GelMA being the polymeric backbone, makes this a versatile bioactive platform. GelMA hydrogels possess cell adhesion properties, owing to RGD motifs from gelatin and allow for in vitro enzymatic degradation, indicated by the presence of matrix metalloproteinase (MMP)-sensitive motifs suitable for ECM remodeling. [41]. Moreover, GelMA is compatible with various biofabrication techniques (such as, 3D bioprinting, photo-patterning, etc.), that ensures adaptability of the platform for creating tissue-specific 3D structures and patterns with controlled cellular behavior in 4D. While in this study spatial patterning is limited to a relatively large biphasic disk, 3D bioprinting and photopatterning will also result in more defined spatial patterns of aptamer availability, enabling a more precise investigation of the spatial control over vascular organization. Furthermore, different cellular responses were observed within aptamer-functionalized regions compared to GelMA regions for at least ten days of culture (Fig. 7), displaying the long-term stability of GFs in these systems. This indicates the aptamers ability to stabilize GFs against various environmental stresses [42] and thereby contributing towards their long-term bioavailability. Similar behavior is observed in nature where upon ECM binding, GFs become stabilized and protected. [43].

Given the simplicity of designing aptamer sequences to capture and release various GFs, this platform is well adaptable for any GF of choice. In complex processes such as angiogenesis, multiple GFs are required at different stages of development. [44]. Specifically, VEGF is responsible for initiating the angiogenesis process and formation of early blood vessels, whereas presence of PDGF-BB ensures their stabilization and maturation. [44]. Owing to aptamers high binding specificities, it is possible to incorporate multiple aptamers within the same hydrogel with different release kinetics for multiple GFs. In outlook, our future studies will focus on achieving mature perfusable vascular networks in vitro, by combining both GFs that initiate and stabilize vasculature in a single hydrogel system. On the longer term, these results, combined with other developments in aptamer technology, provide significant potential for in vivo and clinical applications. The platform’s ability to induce temporally controlled cell responses by modulating GF bioavailability, provides significant advantages over the currently available strategies for enhancing tissue regeneration within the actively remodeling microenvironment of an implanted engineered tissue. Prior research has revealed the in vivo efficacy of intravenous delivery of drugs conjugated with oligonucleotide sequence for targeted refilling of implanted drug depot and mitigate tumors. [45]. Thus, we speculate the possibility to achieve temporally controlled targeted GF release via intravenous delivery of the external trigger (CS) post-implantation. Alternatively, a local injection of CS could also be used for triggered GF delivery. Notably, the aptamers used in present study were not modified to increase their nuclease resistance. However, various modifications such as PEGylation, locked nucleic acids (LNA), phosphorothioate modification, Spiegelmers (mirror-image L-oligonucleotide aptamers) and q-quadraplex aptamers have been previously reported to enhance their resistance for nuclease-based degradation which is important for long-term in vivo application. [46].

Supplementary data related to this article can be found at https://doi.org/10.1016/j.bioactmat.2021.10.024.

4. Conclusion

In conclusion, we demonstrate an aptamer-based dynamic platform for spatiotemporally controlling the bioavailability of an angiogenic GF by exploiting affinity interactions, within a biofabrication friendly polymeric matrix. The platform showcases various outstanding features namely, (i) sustained aptamer incorporation with polymeric matrix via chemical modification, (ii) rapid and localized GF sequestration from culture medium, (iii) triggered GF release via CS hybridization without any non-specific leakage until ten days, (iv) bioactivity for 3D co-cultures, (v) the ability to selectively guide cell responses in 3D, as well as (vi) spatiotemporally regulating tubular morphogenesis of endothelial cells by controlling aptamer-GF bioavailability within bi-phasic hydrogels, similar to native ECM. Moreover, the novel ability to locally manipulate microvascular network formation within bi-phasic hydrogels that are incubated in a single pool of culture medium, opens up new avenues for in situ manipulation of cell behavior within aptamer-patterned 3D scaffolds. Altogether, these data provide sufficient proof for harnessing aptamers as biochemical triggers to locally manipulate cell response in 4D, opening up the avenues for creating clinically relevant dynamic biomaterials to be explored for regenerative medicine and tissue engineering applications.

Author contributions

D.R. and J.R. conceived the project. D.R. designed the study, performed experiments/analysis and prepared the manuscript; A.K. conducted rheological experiments; N.S. & I.I. contributed in ELISA data analysis; B.K. and J.R. provided financial and administrative support to the project; J.R. contributed to the designing of experiments and manuscript writing/correction. All authors discussed the results and commented on the manuscript.

CRediT authorship contribution statement

Deepti Rana: Conceptualization, Methodology, Investigation, Formal analysis, Writing – original draft, Writing – review & editing. Ajoy Kandar: Formal analysis, Investigation. Ilyas Inci: Formal analysis, Investigation. Ilyas Inci: Resources, Formal analysis. Bart Koopman: Supervision, Project administration. Jeroen Rouwkema: Conceptualization, Writing – review & editing, Supervision, Project administration, Funding acquisition.

Declaration of competing interest

The authors declare no competing interests.

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Appendix A. Supplementary data

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