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Highly efficient intracellular transduction in three-dimensional gradients for programming cell fate

Hoda M. Eltaher¹, ², Jing Yang¹, Kevin M. Shakesheff*¹ & James E. Dixon*¹

¹Wolfson Centre for Stem Cells, Tissue Engineering, and Modelling (STEM), Centre of Biomolecular Sciences, School of Pharmacy; University of Nottingham, Nottingham, NG7 2RD, UK

²Department of Pharmaceutics, Faculty of Pharmacy; Alexandria University, Alexandria, 21521, Egypt

*Correspondence should be addressed to Kevin M. Shakesheff or James E. Dixon

kevin.shakesheff@nottingham.ac.uk

james.dixon@nottingham.ac.uk

Wolfson Centre for Stem Cells, Tissue Engineering, and Modelling (STEM)

Centre for Biomolecular Sciences

University of Nottingham

University Park

Nottingham NG7 2RD, UK

Tel: +44 (0) 115 9515 104/121
Abstract

Fundamental behaviour such as cell fate, growth and death are mediated through the control of key genetic transcriptional regulators. These regulators are activated or repressed by the integration of multiple signalling molecules in spatio-temporal gradients. Engineering these gradients is complex but considered key in controlling tissue formation in regenerative medicine approaches. Direct programming of cells using exogenously delivered transcription factors can by-pass growth factor complexity but there is still a requirement to deliver such activity spatio-temporally. We previously developed a technology termed GAG-binding enhanced transduction (GET) to efficiently deliver a variety of cargoes intracellularly using GAG-binding domains to promote cell targeting, and cell penetrating peptides (CPPs) to allow cell entry. Herein we demonstrate that GET can be used in a three dimensional (3D) hydrogel matrix to produce gradients of intracellular transduction of mammalian cells. Using a compartmentalized diffusion model with a source-gel-sink (So-G-Si) assembly, we created gradients of reporter proteins (mRFP1-tagged) and a transcription factor (TF, myogenic master regulator MyoD) and showed that GET can be used to deliver molecules into cells spatio-temporally by monitoring intracellular transduction and gene expression programming as a function of location and time. The ability to spatio-temporally control the intracellular delivery of functional proteins will allow the establishment of gradients of cell programming in hydrogels and approaches to direct cellular behaviour for many regenerative medicine applications.

**Keywords:** Intracellular transduction; Gradient; Diffusion, Hydrogel, GAG-binding enhanced transduction (GET), CPP, Myogenesis.
1. Introduction

Authentic regeneration of adult tissue requires exquisite control of the biological microenvironment and the appropriate cellular plasticity in order to recapitulate the developmental processes from which the tissue was originally formed. Key cellular processes like migration, proliferation, adhesion and differentiation occur in response to chemical cues present within the microenvironment such as interaction with extracellular matrix (ECM) and stimulation with morphogens or growth factors [1, 2]. These macromolecules trigger intracellular signal transduction pathways, with complex cross-talk between the cascades and eventual programming of cell behaviour changes at the level of gene expression. Such networks provide important positional information that control cell behaviour in a concentration dependent manner inside the tissue [2-4]. In this context, recreating gradients of such molecules in 3D scaffolds has been used extensively to study such biological responses in vitro and by exploiting bioengineering strategies, this knowledge may help permit the development of complex tissues for regenerative applications.

Many approaches have been employed to produce scaffolds with biochemical gradients to control cell behaviour [1, 5]. These methods include internal microfluidic channels [6-8], controlled protein release using drug delivery systems [9-12], gradual immobilization of signalling molecules throughout the scaffold geometry [13-15], simple molecular diffusion throughout the scaffold [16, 17] or combinations of these methods [18-21]. The simplest and most representative method is to employ diffusion to generate gradients using a source (So) and a sink (Si) separating a cell-laden hydrogel (gel, G) structure (So-G-Si). This method, despite being time consuming, has been successfully used to study cellular responses to soluble chemical gradients and has been chosen in several studies (over convection-based
methods which employ microfluidics) as they may introduce extra cellular shear stress and asymmetric mass transport unlike simple diffusion systems [22, 23].

Within a tissue, 3D concentration gradients are thought to exist for every soluble component, such as those consumed or produced by the cells from basic nutrients to signalling molecules. Gradients form due to competition between diffusion of these molecules and convection by cell consumption or release [24]. Local signal concentration and duration affect the cellular response directly; meaning cells have alternate interpretation of signals in different zones of tissue depending on this dynamic [4]. The ultimate realization of signalling is the interpretation by the cell to activate key regulatory genes, which in turn, instruct cell behaviour. We and others have shown that by delivering specific transcriptional master regulators either genetically (via viruses [25]) or recombinantly (using purified proteins, [26]) powerful cell fate changes can be programmed, effectively by-passing this highly-complicated interaction of extracellular signalling.

Various methodologies have been shown to deliver recombinant proteins intracellularly and elicit changes in cell behaviour [27-29], and direct differentiation of somatic [30, 31] or stem cells [32] which in essence could replace complex signalling network reconstruction. Cell penetrating peptides (CPPs) can be tethered to the cargo protein of interest [33-35] and these trigger endocytosis-mediated uptake into cells [27-29]. Even though CPPs significantly increase cellular uptake, their activity requires them in vast extracellular excess to drive uptake, with the quantities required to elicit any cell behaviour change in the micromolar scale. To overcome this inefficiency in delivery, we recently described a new technology named glycosaminoglycan GAG enhanced transduction (GET) [26]. We demonstrated enhanced intracellular transduction upon coupling CPPs with a GAG-binding peptide
(the composite peptide termed GET peptides). Functional quantities of many cargos including fluorescent proteins, transcription factors and enzymes were successfully delivered using GET with significant increase in functional delivery compared to conventional CPP-mediated delivery and using sub-micromolar doses.

Here, we demonstrate that the efficient cellular uptake of proteins using the GET technology may be employed in a 3D hydrogel system (Gelatin-Agarose; GA). Using a diffusion-based model (So-G-Si), we have created 3D cell–laden hydrogels with dynamic spatio-temporal gradients of GET proteins to study cell uptake and the potential of GET to be used to program cell behaviour in 3D.

2. Materials and methods

2.1. Hydrogel fabrication in compartmentalized diffusion chamber

Gelatin-Agarose (GA) hydrogels contained Type IX, ultra-low gelling temperature agarose (1% w/v, Sigma-Aldrich, UK), bovine skin Gelatin (1% w/v, Sigma-Aldrich, UK), and foetal calf serum (FCS, 10% w/v) in Dulbecco’s Modified Eagle’s Medium (DMEM; Gibco, UK). Prior to final formulation Gelatin and Agarose (2% w/v each) were heated to 60°C for 30 minutes to dissolve the powder, sterilized by exposure to UV lamp for 20 minutes and the pre-warmed (37°C) FCS and DMEM added to the correct concentration. Molten Gelatin-Agarose (GA) hydrogel was kept at 37°C in humidified incubator for usage. 500 µL hydrogel was cast in custom made moulds to create three compartmentalized assembly of source-gel-sink. The assembly was modified from X. Cao and MS. Shoichet [16] such that acetal resin mould (ARM) and portable polytetrafluoroethylene (PTFE) fitted blocks were used to create the middle hydrogel compartment separating two compartments of equal dimensions (source and sink compartments) (Figure S1C). ARMs were oxygen etched rendering the surface hydrophilic and improving hydrogel attachment prior to diffusion studies [36].
The oxygen etching was carried out for 15 minutes in a T-shaped borosilicate chamber closed with stainless steel endplates that were sealed with Viton O-rings at a power of 20 W, under a working pressure of 300 m Torr. Prior to hydrogel casting, the ARM and PTFE blocks were sterilized in industrialized methylated spirit (IMS; 70% v/v) for 30 minutes, washed twice with sterile PBS, air dried in microbiological safety cabinet and then exposed to UV light for 20 minutes. This resulted in scaffolds with the dimension of 15×5×6.6 mm (Length × Width × Height) (Figure S1D).

For cellular scaffolds, NIH 3T3 mouse fibroblasts cells were suspended in the prepared gels at 2x10⁶ /ml at 37°C. Hydrogels were allowed to cast in moulds for 30 minutes at 4°C then the PTFE blocks were removed creating the sink and source compartments.

2.2. Diffusion studies and GET proteins

Source compartments were loaded with 50 µg/ml of test recombinant protein in 500 µL growth medium unless otherwise stated. Non-transducing monomeric red fluorescent protein (mRFP1; mR), CPP (eight arginine, 8R) -tagged mRFP1 (mR-8R), heparin binding domain; HBD (P21) –tagged mRFP1 (P21-mR), or the complete Glycosaminoglycan-binding enhanced transduction (GET is CPP and HBD) –tagged mRFP1 or MyoD protein (P21-mR-8R and P21-mR-MyoD-8R, respectively) were expressed and purified from E.coli using GST affinity purification as previously described by Dixon et al. [26]. Sink compartments were loaded with 500 µL growth medium. Diffusion experiments were carried out at 37°C under 5% CO₂ in humidified incubator.

2.3. Diffusion gradient evaluation

At pre-determined time points post-diffusion, cellular and acellular scaffolds were directly frozen in OCT mounting media (VWR, USA) at -80°C which prevented
further diffusion. These were then serially sectioned perpendicular to the direction of protein diffusion using Leica CM 1100 cryostat at -20ºC into 20 µm thick slices. Each slice was vortexed in warm in PBS (350µl) and the protein concentration within was then determined by fluorometric analysis (excitation: 584 nm; emission: 620 nm) using a TECAN infinite 200PRO multimode reader. Each time point diffusion study was carried out in triplicate, and analysis to detect protein concentration per slice was performed in triplicate for each hydrogel. Diffusion gradient profiles across the width of the hydrogel scaffolds were constructed by plotting protein concentrations per 20 µm width slice at different time points as a function of distance in the scaffold away from source.

Alternatively, cellular scaffolds at different time points were exposed to a selective washing protocol which removed all fluorescence that was not associated with or internal to cells (as determined by optimisation using acellular scaffold). Cells within hydrogel were fixed by 3.7% (w/v) para-formaldehyde (PFA; Sigma-Aldrich, UK) for 3 hours at 4 ºC, followed by 2 hours in sucrose 15% w/v (Sigma-Aldrich, UK) in PBS at 4ºC and finally overnight in sucrose 30% w/v at 4ºC. Scaffolds were then placed in OCT mounting media, frozen in liquid nitrogen cooled isopentane bath then stored at -80ºC. Scaffolds were subsequently sectioned parallel to the direction of protein diffusion using Leica CM 1100 cryostat at -20 ºC into 20 µm thick slices. Retrieved gel slices were then viewed under fluorescence image capturing using Nikon stereomicroscope SMZ 1500 for entire scaffold imaging.

2.4. Intracellular transduction analyses

Cellular scaffolds at determined time points were manually sliced perpendicular to the direction of the protein diffusion into five parallel sections each 1 mm in diameter using sterile disposable blades. Each hydrogel slice was degraded by
enzymatic digestion following a protocol modified from Ran Yu et al. [37]. Briefly, hydrogel slices were digested by addition of 10 µL 10X agarase reaction buffer (New England Biolabs, USA) to each slice. The mixture was heated at 50 ºC for 5 minutes and left to cool to 44ºC for 25 minutes. 2 µL β-agarase I enzyme (New England Biolabs, USA) were added to each sample and the mixtures were maintained for 1 hour at 44ºC. Hoechst 33342 at 2.5 µg/mL (Cell Signaling, USA) was added to the liquefied gel slices and stained for 30 minutes at room temperature. Samples were then analysed using a MoFlo™ DP (DAKO) Flow Cytometer using the green (561 nm) and violet (405 nm) lasers to detect staining for red fluorescent proteins and Hoechst 33342 respectively (20,000 events per sample). Mean red fluorescence per cell was used to compare transduction activity with background from unlabelled cells being subtracted and values taken as ratios to the control monolayer of NIH mouse fibroblasts (1x10^6 cells) exposed to an equivalent concentration of recombinant for 24 hours. Data shown was collected from three independent experiments.

2.5. Protein size and molecular weight

Protein size (hydrodynamic radius) of the variable test proteins was determined using the Zetasizer Nano ZS (Malvern instrument) applying the principle of dynamic light scattering. Measurements were done on protein solutions in HEPES buffer pH 7.2 at 37ºC. For determination of protein molecular weights, sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) using Novex® 10% Tris-Glycine mini protein gel (Invitrogen) was applied following manufacturer’s instructions. Gels were stained with Coomassie blue (Invitrogen) and de-stained before imaging.

2.6. Hydrogel porosity
The mesh size of GA hydrogels were estimated using transmission electron microscopy. Sucrose-dehydrated hydrogel sample was sectioned down to 80 nm thick slices at -60 °C and 0.9 mm/s cutting speed using a freshly prepared glass knife in the RMC Powertome with CR-X cryo-attachment. Sections were placed on holey carbon support films in copper grids (HC300-Cu, Electron Microscopy Sciences, UK), then transferred to the JEOL 2100F transmission electron microscope operating at an accelerating voltage of 200 kV. Imaging was performed using the High Angle Annular Dark Field STEM mode. Randomly selected pores in the transmission electron micrographs were used to determine the average pore size using ImageJ software.

2.7. Real time-qPCR

Cellular scaffolds (Human embryonic kidney cells; HEK 293, 2×10^6 cells/mL), exposed to culture medium or GET-MyoD (P21-mR-MyoD-8R) (60 µg/mL) for 3 days, were serially sliced into five parallel sections each 1 mm in diameter and snap frozen at -80 °C. RNA was extracted using a Qiagen RNeasy mini kit (Qiagen, UK) and quantified using a Nanodrop spectrophotometer (ND-100 Thermo Scientific). This was followed by reverse transcription into cDNA using SuperScript™ III Reverse Transcriptase enzyme (Invitrogen, UK) according to manufacturer's instructions. Quantification of the gene expression was performed using rT-qPCR analysis (ΔΔC₅) using Taqman assays for human Actin-β, MyoD1, Myogenin, Myf5 and Desmin (Applied Biosystems, UK) as detailed Dixon et al.[26].

2.8. Statistical analysis

Statistical comparisons were carried out using the GraphPad Prism version 6. The statistical significance was determined using two-way ANOVA followed by Dunnett’s multiple comparisons test for viability experiments. Results were
considered significant if \( p<0.01 \). Spearman’s regression was used to calculate the rank correlation of the gradients produced. Further information on these analyses can be found in supplemental methods.

3. Results

3.1. Engineering of a hydrogel diffusion method

We engineered a moulding system (acetal resin mould; ARM) in which we could cast a hydrogel (gel, G) sample between a source (So) and sink (Si) area (creating a compartmentalised chamber of So-G-Si) (Figure S1). This system, by the use of plasma oxygen etching prior to casting the hydrogel, reproducibly demonstrated no leakage (using Fluorescein) between source and sink (by-passing the construct) for a variety of hydrogels (collagen, alginate, agarose, fibrin) (Figure S2). This was further confirmed by conducting opposing diffusion studies of TRITC- and FITC-labelled bovine serum albumin (BSA) from different compartments neighbouring the hydrogel (Figure 1 and S2). Mirror image diffusion profiles of TRITC- and FITC-BSA were established confirming the potential of the compartmental diffusion model/quantification protocol in developing and evaluation of chemical gradients irrelevant of the fluorescent tag employed. We used cryosectioning to analyse hydrogels (which required freezing samples without sucrose embedding) to avoid dilution of the soluble fluorescent signals and disturbance of the diffusion profile. Our technique created large mesh sizes in the samples (Figure 1B-D; produced by ice crystals) but faithfully retained the diffusion profile (Figure 1E) [38].

We chose to use agarose hydrogel due to its rapid casting and the ability to simply vary its diffusion kinetics by altering hydrogel concentration. The agarose was modified with gelatin (Gelatin-Agarose; GA) to aid cell adhesion to the matrix. This
modification behaved similarly to agarose-only matrices and allowed the cells to retain viability over a 3-day experiment (Figure S3). We thought it important to assess the GA hydrogel porosity by assessing mesh size (with sucrose embedding). Transmission electron microscopy (TEM) demonstrated the GA hydrogel possesses pores with an average size of 0.198 ± 0.079 µm (Figure S4A). The pore size of frozen-hydrated GA samples was, furthermore, confirmed using focused ion beam scanning electron microscope (FIB-SEM) coupled with TEM [39]. This nanoporous hydrogel therefore dictates that encapsulated cells have restricted movement, allowing the gradients of proteins to be determined without changes confounding analyses over time such as cell migration.

To further define our system, we sectioned cell-laden GA hydrogels in all three dimensions and determined the number of cells in each section (compared to the expected numbers if uniformly distributed). We were able to confirm that cells were evenly distributed throughout the GA hydrogel (Figure S5) and therefore we had created homogenously cell-laden hydrogel system for diffusion analyses.

**3.2. Using GET-tagged proteins to assess the effect of intracellular transduction on protein gradients**

In our previous studies [26], we characterized the dynamics of GET-mediated transduction and compared the efficiency of intracellular transduction to non-transducing controls (mRFP1 only), less efficient transducing proteins (tagged with a CPP; mRFP-8R), and proteins with limited transduction but efficient cell interaction (P21-mR). We aimed to use all these variants within the gradient system which would allow us to determine whether cell binding verses transduction had the most significant influence of gradient profile. To allow accurate data interpretation, we
Initially assessed if differences in our test proteins’ hydrodynamic radii in 10 mM HEPES buffer (Dynamic Light Scattering analyses; DLS; Figure S4B and S6) and molecular weight (SDS gel electrophoresis; Figure S4C) were likely to affect diffusion. DLS assessment revealed similar sizes for all test proteins (mR, mR-8R, P21-mR and P21-mR-8R) ranging from 2.36 to 5.17 nm hydrodynamic radii (Figure S4B and S6). All proteins ran close to their theoretical molecular weights (all around ~28 kDa; SDS-PAGE assessment) calculated based on their amino acid sequences (Figure S4C) [40]. We therefore conclude that simple protein size was unlikely to be a major factor influencing diffusion behaviour.

Initially we tested the diffusion gradient profiles at different time points (Figure 2 & 3) within our GA hydrogel system with mRFP1 recombinant protein (50µg/ml source concentration, ~1.8µM, Figure 2A). GA scaffolds (Figure 2B), showed a steep fluorescent concentration gradient across the length of the hydrogel after 6 hours. Further time led to the gradient being less pronounced (at 24 hours) with an increase in mRFP1 fluorescence within slices distant from the source and reciprocal decrease in areas close to the source, at which time sink concentration also significantly increased (Figure S7). At 72 hours, the gradient is lost with equal mRFP1 fluorescence throughout the hydrogel (irrespective to distance away from source) suggesting equilibrium over the scaffold width. We used an exponential decay model (72h data) to model the relationship between time and fluorescence (Figure 2B, right graph) which resulted in a $R^2 = 0.405$ suggesting a relative scatter of fluorescence signal was around an average value rather than a gradient, statistically demonstrating equilibrium. Further emphasis on the statistical dependence between the distance in scaffolds and the corresponding protein concentration is denoted by Spearman’s correlation rank coefficient ($r_s$). A low value ($r_s = -0.53$ at $p < 0.01$) was
calculated indicating relatively poor correlation between variables, further demonstrating equilibrium. Importantly we show that our system can rapidly create a diffused gradient of mRFP1 protein within 24 hours and that this reaches equilibrium within the hydrogel by 72 hours.

### 3.2.1. Cells do not prevent equilibration of non-transducing proteins

We next assessed if diffusion within constructs was affected by the presence of mammalian cells (NIH 3T3 cells) aiming to determine the rate of gradient formation and equilibration. Testing different cell densities in scaffolds (0.1-2x10^6 cells/ml) gave similar diffusion profiles to acellular scaffolds with comparable trends at respective time points (Figure 2C). The coefficient of determination (for exponential decay fitted data; Figure 1C, right graph, R^2) and Spearman’s correlation rank coefficient (r_s) for cellular scaffolds was comparable to acellular experiments with 72 hours post-initiation being that both achieved complete equilibrium (cellular and acellular being R^2 = 0.503 and 0.405, and r_s = -0.6374 and -0.5297, respectively; Table S1). These data demonstrate that early gradients (up to 24 hours) are dissipated and an equilibrium state is achieved regardless of presence or absence of cells within 72 hours. It is important that mRFP1 protein diffusion is statistically unaffected by the presence of cells in our system as we have previously shown it cannot efficiently interact with or transduce into cells [26].

### 3.2.2. GET-proteins transduce cells efficiently in 3D hydrogels

We hypothesized that the diffusion profile of a protein that efficiently interacts with or transduces into cells would be significantly affected in cellular versus acellular scaffolds. To test this, we assessed diffusion of GET-tagged mRFP1 protein (P21-mR-8R, Figure 3A) which transduces cells rapidly (~1 hour) and has been shown previously to concentrate intracellularly depending on extracellular concentration
mRFP1 (non-transducing) and P21-mR-8R (transducing) diffusion was very similar through both acellular and cellular hydrogels over 6 and 24 hours (Figure 3B & 3C). However, unlike as demonstrated for mRFP1, the well-established gradient of P21-mR-8R is retained in the cellular scaffold at 72 hours. This gradient was a function of distance from source with further zones being lower in protein concentration compared to those neighbouring the source. Importantly this effect was not seen for acellular scaffold which equilibrated with the same efficiency as mRFP1 with or without cells. Analyses of the data from P21-mR-8R diffused cellular scaffolds (Figure 3C) showed they maintain a steep gradient at 72 hours \( R^2=0.8811 \) and \( r_s=-1.0000 \) relative to equilibrium seen in acellular hydrogels \( R^2=0.5091 \) and \( r_s=-0.6573 \) (Figure 3B and Table S1). Determination of the diffusion coefficients of both test proteins across GA hydrogels (Figure S8) was calculated using the model permeation method previously described in Cao and Shoichet [16] (using Equation 1). The diffusion coefficients of mRFP1 & P21-mR-8R were determined to be \( 7.15 \times 10^{-7} \) & \( 7.53 \times 10^{-7} \) cm\(^2\)/s respectively.

3.3. GET-proteins transduce cells in a gradient

We wanted to assess whether the gradient we observed for transducing P21-mR-8R in cellular scaffolds was due to intracellular transduction and retention in cells or possible slowing of the diffusion kinetics. From our previous work, we have shown that GET-tagged mRFP transduces cells orders of magnitude more than untagged or CPP-tagged proteins [26]. This efficiency of transduction would likely deplete freely soluble protein from the system and significantly change the gradient kinetics. In order to achieve this, we developed a washing protocol which was employed following hydrogel/cell fixation to remove all of the freely diffusible proteins within the hydrogel slice analysed. Any fluorescence retained would therefore be from
transduced cells which could be observed microscopically (Figure 4). The protocol was set up such that cells remained preserved and intact, while all signal from the gel component was completely removed (optimised on acellular hydrogels using mRFP1). Fluorescence microscopy was used to compare the effect of the transducing P21-mR-8R to the non-transducing mRFP1 in cellular hydrogels, which demonstrated that P21-mR-8R scaffolds possessed strongly red fluorescently-labelled cells compared to unlabelled cells for mRFP1 diffused hydrogels (Figure 4). We assessed a time course of diffusion using these analyses and were able to demonstrate that width of the hydrogel bands through which the transduced cells could be observed (following P21-mR-8R diffusion) was increased as a function of time and closely mirrored the initial protein gradient established in hydrogels (Figure 3).

Equilibrium was observed across the width of cellular and acellular hydrogel scaffolds exposed to mRFP1 as demonstrated by low value of coefficient of determination of the exponential phase decay model. This statistic acts as an indication of free diffusion of the reporter protein without remarkable cellular uptake within 72 hours. A similar behaviour was observed for acellular scaffolds exposed to P21-mR-8R; however, the presence of cells maintained a steep gradient (exponential decay) demonstrated by higher goodness of fit $r^2 = 0.881$. This data emphasizes enhanced cellular uptake of P21-mR-8R shifts the state of equilibrium (demonstrated by evenness in concentrations of mRFP1 across the hydrogel width) to an exponential decay curve.

3.4. Intracellular transduction of GET proteins is directly proportional to early concentration gradient
To confirm the intracellular transduction of P21-mR-8R, intact cells were retrieved from the hydrogel and their fluorescent intensities analysed by flow cytometry with the mean cell intensity assessed as function distance from the source. We optimized the recovery procedure to ensure cells remained intact (recovered from liquefied 1 mm thick hydrogel slices) and were counter-labelled with nuclear stain Hoechst 33342. It was found essential to counterstain cells to exclude non-digested hydrogel debris as events in the data analysis. Only cells that were positively stained for mRFP1 and Hoechst 33342 were used for the evaluation and we assessed the transduction of P21-mR-8R versus mRFP1, and singly tagged proteins P21-mR and mR-8R (Figure 5A).

Cells retrieved from hydrogels through which mRFP1 or control culture medium diffused showed equal fluorescence intensities all over the scaffold width at different time points (Figure 5B & 5C). Similarly, minimal cell fluorescence uptake/labelling were observed following diffusion of the cell binding P21-mR. As hypothesized, cells in scaffolds treated with P21-mR-8R, showed gradual decrease in fluorescence intensity as a function of distance at 24 and 72 hours (Figure 5B & 5C). These cells possess significantly higher fluorescence values compared to cells within control culture medium and mRFP1 treated samples at each slice position (Figure S9). Furthermore, the gradual intracellular transduction mediated by P21-mR-8R in samples was significantly higher than samples with CPP-tagged mRFP (mR-8R) for both 24 hours and 72 hours. Interestingly, both CPP- and GET-tagged proteins maintained a similar spatial trend, suggesting that the intracellular sequestering of the mRFP1 signal was responsible for maintenance of the gradient.

We next assessed slices microscopically; liquefied hydrogel suspensions from slices were analysed for control media, and mRFP1 or P21-mR-8R (at 24 and 72
hours; Figure 6A and 6B, respectively) were assessed for background signal and cell-associated fluorescence. We defined cell-associated fluorescence as Hoechst 33342-positive foci (highlighting the presence of cells) which we used to visualize the nucleus of cells to exclude any red fluorescence not associated with cells. Cells in the liquefied hydrogel slices from control culture media diffused did not show any red fluorescence signal, which also demonstrated no background fluorescence from the hydrogel or the culture medium. Liquefied scaffold slices from mRFP1 treatments showed negligible red fluorescence associated with cells. However, they did show gradual decrease in the fluorescence intensity emerging from the hydrogel as a function of distance (24h, Figure 6A) and relatively less steep to equilibrated fluorescence intensity change over time (72 hours, Figure 6B).

For P21-mR-8R treated scaffolds, early in culture, significant cell-associated and background red fluorescence was observed with progressive decrease of the overall signal relative to the distance from the source (24 hours, Figure 6A). Further culturing yielded stronger cell-associated fluorescence, which was proportional to distance from source (72 hours, Figure 6B). Interestingly the background fluorescence which represents freely-diffusible fluorescence was progressively reduced over culture time, concomitant with a reciprocal cell-associated increase in fluorescence. To confirm this qualitative assessment, we used semi-quantitative fluorescent image analysis (Figure S10), and plotted mean red fluorescence of the images for mRFP1 and P21-mR-8R (at 24 and 72 hours; Figure S10A & B, respectively) as a function of distance (4 images/distance). As determined by fluorometry and our qualitative assessment both proteins have matching the trends. This confirms that at least over the time analysed (up to 72 hours) that distance of fluorescence from source is not affected by whether the protein can transduce cells.
Even though the mean fluorescence appeared similar for mRFP1 and P21-mR-8R proteins in these analyses, it was clear that the cell-association of the fluorescence distinguished their diffusion profiles. To express this, we analysed variation (standard deviation of pixel intensity) across the same images; as cell-associated fluorescence appeared as punctate signal in the cells, more cell association creates an increased signal variation in a sparsely populated scaffold. The fluorescence signal for the mRFP1 scaffold had low variance compared to significantly higher variation for P21-mR-8R images (Figure S10B). These data further confirm that though early in this diffusion model for cellular scaffolds that analyses that do not distinguish between cell-association yield essentially the same diffusion trend; however, it is clear that cell-transduction can change diffusible protein availability by sequestering the signal inside cells.

3.5. Induction of early myogenic differentiation using GET-MyoD and a 3D concentration gradient

To demonstrate that the GET technology (combined with the compartmental diffusion model) was capable for efficient gradient delivery of functional proteins, the transcription factor MyoD (the myogenic differentiation master-regulator) was GET-tagged [26] and allowed to diffuse across cell laden GA (GET-MyoD, P21-mR-MyoD-8R; Figure 7A). We initially assessed the diffusion profile in acellular and cellular GA hydrogels (Figure S11) and the diffusion co-efficient of GET-MyoD (Figure S12) which confirmed enhanced cellular transduction and a similar diffusion profile to P21-mR-8R. Next we aimed to show the gradient produced leads to a corresponding induction of early myogenic differentiation (in terms of myogenic gene expression) as a function of distance away from source protein or control culture medium by RT-
qPCR. Initially we showed that HEK293T cells cultured in conventional monolayers respond dose-dependently to GET-MyoD and initiate endogenous myogenic gene expression in 72 hours (Figure 7B). Comparing monolayer- versus GA hydrogel-cultures directly exposed to GET-MyoD, we showed relatively inhibited myogenic induction in GA hydrogels but that GET-MyoD did indeed significantly induce myogenic gene expression in this microenvironment (Figure 7B and C). Importantly, without stimulating cells with GET-MyoD, there was no effect on background myogenic- or house-keeping gene expression (β-actin) when culturing cells in GA scaffolds. Using our 3D diffusion system, we found a pronounced increase of the endogenous expression of MYOD and other myogenic markers (MYOGENIN, DESMIN and MYF5) in GA slices retrieved from scaffolds exposed to GET-MyoD protein for 72 hours (Figure 7D). This was inversely proportional to the distance from the source and no induction was observed in scaffolds exposed to control culture medium. Interestingly, induction of myogenesis in GA gradients (at similar concentrations taken at specific points), was lower than the corresponding monolayer or uniformly-stimulated GA hydrogels (Figure 7C and 7D). Importantly, as our gradient system is dynamic (with a progressive increase in protein over time as a function of distance from the source), cells would not be exposed to the same duration and concentration of GET-MyoD in the gradient system. Here, we therefore show that transcription factors delivered by GET can be supplied in a gradient, elicit different gene expression programming efficiencies and that the duration and concentration of GET-MyoD stimulation is distinct from non-gradient systems.

4. Discussion

Here we describe the development of a methodology to create reproducible protein gradients in hydrogels (using a compartmentalized diffusion system). We
created assays to assess the diffusion of recombinant proteins both qualitatively and quantitatively using microscopy, fluorometry and cytometry. Using this system, we assessed the effect of cellular transduction on gradient establishment and retention over time. We employed GET-tagged proteins that represent a novel and efficient transduction approach and compared them to a conventional cell penetrating peptide (CPP)-tagged and control non-transducing proteins. GET-proteins efficiently transduce cells ‘locking in’ the diffusion profile of the protein by immobilising it intracellulary.

The diffusion model employed has been modified from one previously described in Cao and Shoichet [16]. We put significant effort to ensure that a complete seal between the hydrogel matrix and the base of the mould was achieved which is a significant technical flaw in most diffusion models. The diffusion chamber (ARM, acetal resin mould) was designed to create a well with portable PTFE blocks creating source and sink chambers that the hydrogel could be cast between. This is distinct from that of Cao and Shoichet in which they used glued components to create the source, sample, sink arrangement [16]. The acetal polymer used here is hydrophobic, therefore the base of the mould underneath the hydrogel compartment was oxygen plasma etched to render the surface more hydrophilic and achieve a seamless contact between the hydrogel and the mould base [36]. The surface modification and diffusion chamber design was necessary to guarantee that the diffusion occurred only through the hydrogel with minimal leakage from the source to sink compartment underneath the hydrogel scaffold (Figure 1 and S1-2). Importantly this was confirmed by monitoring the protein concentration in the source and sink compartments over time which demonstrated that time versus sink and source fluorescence were directly and inversely proportional, respectively, over 72 hours.
Other systems that aimed to assess gradients within hydrogels in the presence of cells mostly employ hydrogels cast over a cell monolayer [16, 41]. We aimed that cellular components were evenly dispersed in the whole 3D matrix (Figure S5) rather than plating cells on the base of the mould with the hydrogel being cast above or cells seeded on top of the hydrogel construct [16, 42]. This was needed in the current study to demonstrate the effect of cells on the diffusion of proteins and to expose the cells to conditions mimicking the natural microenvironment that accommodates the cells.

The diffusion profiles of mRFP1 across cellular and acellular hydrogel scaffolds were similar. We demonstrated that the fluorescent signal detected was that retrieved from freely diffusible proteins and of minimal contribution by the cells. This was further confirmed by applying successive washing steps to cellular hydrogel scaffolds. The washing protocol completely removed any freely bound proteins in the hydrogel matrix and only fluorescent signals from transduced cells were visible. Cellular scaffolds through which mRFP1 diffused did not show any fluorescent signal associated with the cells (Figure 5 & 6) suggesting that the gradient observed (Figure 2A & B) was mainly in the hydrogel rather than incorporated inside the cells. This confirms that the key factors regulating diffusion of mRFP1 through GA hydrogel were dependant on the protein-hydrogel interaction and not on the presence of cells similar to that described by Lieleg and Ribbeck [43]. mRFP1 molecular weight was estimated experimentally (by SDS-PAGE) to be ~28kDa (theoretically 26.6 kDa and isoelectric point (pI) of 5.66 [40]). This implies that the protein carries a net negative charge in GA hydrogel scaffold which is neutral physiological pH. The ultra-low melting temperature agarose is composed of frequently hydroxyethylated galactose residues so that the agarose fibres are almost
neutral with low electroendosmosis value \( \leq 0.05 \) and low sulphate content \( \leq 0.10\% \). Gelatin type B is a denatured collagen with RGD-like sequences that support cell proliferation and attachment \([44]\) and it is known to have a pI of 4.7-5.2 \([45]\) suggesting that the overall charge of the hydrogel is slightly negative. These attributes would suggest that there is minimal electrostatic interaction between mRFP1 and the GA hydrogel.

The pore size of the GA system was calculated to be \(0.198 \pm 0.079\ \mu m\) and this matches with the reported average pore size of 0.1-0.3 \(\mu m\) for different concentrations and types of agarose (Figure S4A) \([46]\). This pore size cut off is far larger than the hydrodynamic radii of test proteins (Figure S5B) and therefore our system supports diffusion. Interestingly P21-mR-8R, which has a similar molecular weight to the other tests (Figure S4C) but a higher pI of 9.59 \([40]\) is consistently slower diffusing than the other test proteins (Figure 3 and 5). It could be possible that positively charged P21-mR-8R has an electrostatic interaction with the negatively charged hydrogel resulting in relatively slower diffusion. However, the practical estimation of the diffusion coefficients of mRFP1 and P21-mR-8R was more or less similar \(7.15 \times 10^{-7}\) versus \(7.53 \times 10^{-7}\) \(cm^2/s\) respectively (Figure S8, Equation 1) suggesting these differences are small and overall the test proteins possess similar diffusion behaviours in our system.

In addition to protein–hydrogel interaction, the cellular uptake of P21-mR-8R was demonstrated to effectively deplete the freely diffusible protein in the hydrogel and showed that the cells themselves can act as sink, which explains the diffusion behaviour in cellular scaffolds confirming our previous findings \([26]\). This cellular sequestering of P21-mR-8R was seen microscopically (following the washing
protocol) with fluorescent signal only detectable inside cells (confirmed by flow cytometry) and not in the hydrogel matrix.

Only background fluorescence was retrieved from scaffolds exposed to mRFP1 due to its poor cell interaction and transduction activity [26]. Minimal but detectable cellular fluorescence was observed for both CPP-only (mRFP-8R) and HBD-only (P21-mRFP) tests. However, the gradient was retained at significantly lower level for mRFP-8R (compared to that of P21-mR-8R); suggesting that cell transduction was a key factor in retaining the gradient over time (Figure 5). P21-mRFP does transduce cells at low levels but this activity was found insufficient to sequester the amounts of protein intracellularly to retain the gradient. It is likely that cells close to the source (and over time due to diffusion) quickly become saturated with test protein; if the protein is not effectively transduced inside of cells it will still be readily diffusible and the gradient rapidly equilibrates.

Having demonstrated the efficient delivery of GET proteins in cell-laden hydrogels, functional cargo; GET-MyoD (P21-mR-MyoD-8R) was gradually delivered to responsive cells (HEK293T cells) using the compartmental diffusion chamber and the subsequent RT-qPCR analysis assessed myogenic differentiation. Using our 3D system, cells retrieved from all slices of GA scaffolds exposed to GET-MyoD exhibited enhanced myogenic gene expression as compared to the control scaffold demonstrating differences in gene programming (Figure 7D). The profile of gene expression from different myogenic loci was distinct but was evident as gradient of induction. Myogenic programming was lower in the gradient than in statically cultured GA hydrogels where GET-MyoD was homogeneously distributed (Figure 7B and C). This could be due to the slower diffusion of P21-mR-MyoD-8R across GA (D = 4.2×10^{-7} \, \text{cm}^2/\text{s}) due to size (65.3kDa) or the time and duration of GET-MyoD
exposure in gradient being different than if homogeneously delivered in static cultures. Furthermore, this result could be attributed to the altered gene expression in later differentiation stage that might be induced by higher doses of GET-MyoD. This is plausible as threshold dose and duration of transcription factor activity will be different for activation of particular downstream genes. Importantly, MyoD is a myogenic regulatory factor that is highly expressed in early myogenesis and its activity declines in the later phases of differentiation [47] which is consistent with our observations. Importantly, we demonstrate significant induction of zonal myogenic differentiation after 72 hours in response to GET-MyoD gradient in absence of myogenic medium.

Our study complements other’s work aiming at engineering complex gradients in matrices [48]. However, the method employed for our study was unique as it generated a soluble transcription factor gradient which was simply achieved by molecular diffusion. This is in comparison to studies using non-covalent immobilization, such as Runx2 retroviruses (on poly(L-lysine)-coated collagen constructs as described in Phillips et al. [42]) which are far more complex and not based on simple diffusion kinetics. Furthermore, the introduction of highly efficient GET transduction system overcomes the limitations associated with efficiency and toxicity of viral vectors [27], and programs cells without genetic modification. Moreover, our system investigates the effect of gradient transcription factor delivery in a genuine 3D matrix with cells encapsulated within the material rather than seeded on top or underneath the scaffolds [16, 42].

Alternative hydrogel systems with improved cellular viabilities and proliferation characteristics can be employed to recreate soft and vascularized tissue structures beyond that of GA. Also the continuous change of the cellular environment in terms
of cell number, degradation of the hydrogel system combined with the secretion of extracellular matrix would need to be studied to fully understand the factors contributing to concentration gradient manipulation. Finally, careful choice of a clinically relevant cell type and analyses with multiple differentiation markers corresponding to the continuous change of gene profiling with time together with immunohistochemistry will need to be employed to completely study cellular responses towards the generated GET-protein gradients.

5. Conclusions

The compartmental diffusion model presented here succeeded in the production of a 3D gradient of GET proteins which transduce into cells. The higher efficiency of transduction using the synergistic combination of GAG binding proteins together with cell penetrating peptides offers a promise of enhanced delivery of bioactive molecules. We show that the GET system’s enhanced transduction activity can be exploited to deliver functional concentrations of macromolecules including transcription factors inside cells, effectively using the cells themselves as sinks of the delivered molecule. With the ability to control the intracellular delivery of functional cargoes spatio-temporally, we believe that this approach will allow creation of gradients of multiple biologically active proteins to direct cellular behaviour. Ultimately understanding how to control the fate and programming of cells is key for creating successful regenerative medicine systems in three-dimensions.

6. Acknowledgments

The authors acknowledge Bryan Morris (Medical Engineering Unit; M.E.U, University of Nottingham) for diffusion chamber manufacture. Special thanks to Dr David Onion (Flow cytometry facility, University of Nottingham) for the fruitful discussions. We
acknowledge the expertise of Nichola Weston and Dr Michael Fay (Nottingham Nanotechnology and Nanoscience Centre; NNNC, University of Nottingham) for TEM expertise. We acknowledge the financial support from the Egyptian Ministry of Higher Education for PhD scholarship granted to H.M.M.A.E. The research leading to these results has received funding from the European Research Council under the European Community’s Seventh Framework Programme (FP7/2007-2013)/ERC grant agreement 227845. J.E.D and K.M.S acknowledge the support of the Medical Research Council, the Engineering and Physical Sciences Research Council, and the Biotechnology and Biological Sciences Research Council UK Regenerative Medicine Platform Hub “Acellular Approaches for Therapeutic Delivery” (MR/K026682/1).


8. Competing Financial Interests. The authors declare no competing financial interests.

9. References


10. Figure Legends

A

B

C

D

E

TRITC BSA concentration per slice (mg/mL)

0.5

0.45

0.4

0.35

0.3

0.25

0.2

0.15

0.1

0.05

0

So        Distance away from source (μm)        Si

Day 1

Day 3
Figure 1. Testing of compartmentalized diffusion using the engineered chamber and TRITC- and FITC-fluorescently labelled BSA. A) Photograph of setup at 72 hours. Fluorescence microscopy of 1% Agarose (w/v) hydrogel scaffold (15×5×6.6 mm; Length × Width × Height) treated with TRITC- & FITC-BSA diffusing from different sides of the scaffold (TRITC, red in source (So) and FITC, green in sink (Si)). B-D) Fluorescent images of 20 µm thick 1% agarose hydrogel slices sectioned parallel to the direction of the diffusion at day 1 (using Nikon stereomicroscope SMZ 1500). Scale bar is 1mm. E) Pattern of BSA fluorescence of opposing diffusion of TRITC- (left, So) and FITC- (right, Si) labelled BSA across the 1% Agarose (w/v) hydrogel. Each time point was evaluated from triplicate scaffolds through which 0.5 mg/mL TRITC- & FITC-BSA diffused from either side of the hydrogel, respectively.
Figure 2. Non-transducing mRFP1 diffuses to equilibrium in acellular or cellular scaffolds. A) Schematic of mRFP1 recombinant protein used in diffusion studies. Diffusion gradients of mRFP1 (50 µg/mL) across B) acellular and C) cellular (NIH 3T3 mouse fibroblasts, 2x10⁶/mL) GA hydrogels at 6, 24 and 72 hours (n=3).
Figure 3. Efficiently transducing P21-mR-8R maintains a gradient profile in cellular scaffolds. A) Schematic of P21-mRFP1-8R (P21-mR-8R) recombinant protein used in diffusion studies. Diffusion gradients of P21-mR-8R (50 µg/mL) across B) acellular and C) cellular (NIH 3T3 mouse fibroblasts, 2x10^6/mL) GA hydrogels at 6, 24 and 72 hours (n=3).
Figure 4. Temporal assessment of gradients of transduction using fluorescence microscopy. Fluorescent images of cross sections of the cellular hydrogels sliced parallel to the direction of the protein flow: non-transducing mRFP1 (left) and cell penetrating P21-mR-8R (right). Red dots represent transduced cells. Dashed white lines represent the border of the hydrogel scaffold next to the protein source (So) & red arrows represent the width of scaffold through which red transduced cells are dispersed, Sink (Si) (scale bar=0.5 mm).
Figure 5. Intracellular transduction gradient profiles following control, non-transducing (mRFP1), cell binding (P21-mR), CPP-tagged (mR-8R) and enhanced transducing GET (P21-mR-8R) protein diffusion through cellular hydrogels. A) Schematic of recombinant proteins used in diffusion studies. GA hydrogels (NIH 3T3 mouse fibroblasts, 2x10^6/mL) with test proteins were assessed post-sectioning and liquefaction for cell transduction using flow cytometric analysis. Data were normalized to the mean fluorescence intensity per monolayer of cells (1x10^5) exposed to mRFP1 (50 µg/mL) for 24 hours. Data are represented as the mean (n=3) ± standard deviation. Asterisks represent statistical difference between novel enhanced transduction protein P21-mR-8R and cell penetrating peptide mR-8R (p<0.01). While ¥ signs represent statistical difference between CPP-tagged mR-8R and enhanced transduction peptide P21-mR-8R, and the non-transducing reporter protein mRFP1 (p<0.01) using Dunnett’s multiple comparison test.
<table>
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<th>P21-mR-8R</th>
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- **A** 24h
- **B** 72h
Figure 6. Gradients of fluorescence intensity in cells as a function of distance from Source. GA hydrogels were exposed to non-transducing mRFP1 and enhanced transducing P21-mR-8R. Gels were sliced into 1mm sections and liquefied. Representative fluorescence images of digested hydrogel slices retrieved from cellular scaffolds (NIH 3T3 mouse fibroblasts, 2x10⁶/mL) exposed to media, mRFP1 or P21-mR-8R respectively after A) 24 hours or B) 72 hours. Hoechst 33342; blue (left) and red fluorescent protein; red (right), scale bar = 100 µm.
Figure 7. Relative myogenic gene expression as a function of distance away from source using GET-MyoD gradients. A) Schematic of GET-MyoD recombinant proteins used in diffusion studies. B) Relative gene expression of P21-mR-MyoD-8R dose-treatments of HEK293T cells (200,000) over 72h cultured as a monolayer. C) GET-MyoD dosing of HEK293T cells (2x10^6 /mL) over 72h cultured in GA hydrogel. D) GET-MyoD gradients in GA hydrogels. 60 µg/mL P21-mR-MyoD-8R diffused across cellular HEK293T (2x10^6 /mL) GA scaffolds for 72h. Myogenic differentiation was evaluated based on the fold increase in MYOD, MYOGENIN, DESMIN and MYF5 gene expression compared to control cells recovered from 5 different slices of the scaffold. β-actin was used as the house keeping gene control using real time quantitative PCR. Data is expressed as mean (n=5) ± SE. ** p<0.01, * p<0.05.
Supplementary Materials

11. Supplementary Methods

11.1. Temporal monitoring of source/sink protein concentrations

The source and sink compartments of the compartmentalized diffusion model described above were continuously monitored for protein concentrations. This was done to ensure the perfect seal of the middle hydrogel compartment to the base of the mould. No leakage was observed in the proposed model. This was proven by the relatively high proportion of source to sink protein concentrations at day one that started to decrease as more proteins diffused through the hydrogel out to the sink compartments on day three as shown in Figure S1.

11.2. Determination of proteins diffusion coefficients and equilibrium

The diffusion coefficients of mRFP1 and P21-mR-8R across GA hydrogels were determined using the model permeation method. The method was extensively described in Cao and Shoichet [16]. Diffusion studies of mRFP1 and P21-mR-8R in the compartmentalized diffusion model were conducted in acellular hydrogel scaffolds. At different time points, aliquots of the sink compartments (triplicate for each time point) were analysed for the protein concentrations from which a cumulative mass of protein versus time curve was established. The data points on the curve, after achievement of steady state concentrations (24 hours), were fitted using linear regression. The equation developed was then solved for the diffusion coefficients of the different proteins according to

\[ m_i = P_i C_{di} L D_i \]

where \( m_i \) is the mass of proteins received in the sink compartment; \( P_i \) is the permeability of protein in the hydrogel, \( C_{di} \) is the initial concentration of the source compartment, \( L \) is the thickness of the hydrogel membrane and \( D_i \) is the diffusion coefficient of proteins in the hydrogel system. From which, the diffusion coefficients of mRFP1 and P21-mR-8R can be calculated to be 7.15x10^{-7} & 7.53x10^{-7} cm^2/s respectively.

The membrane permeation model assumes that if:

(i) the source side is held at a constant concentration
(ii) the sink is held at zero concentration
(iii) the hydrogel scaffold is initially at zero concentration

The total amount of the diffusing solute that passed through the membrane varies with time according to the following relation:

\[ m_i = P_iC_{di}L(t - \frac{L^2}{6D_i}) \]

where \( m_i \) is the mass of proteins received in the sink compartment; \( P_i \) is the permeability of protein in the hydrogel, \( C_{di} \) is the initial concentration of the source compartment, \( L \) is the thickness of the hydrogel membrane and \( D_i \) is the diffusion coefficient of proteins in the hydrogel system. A plot of \( m \) versus time from this equation is a straight line (once steady state is established) with a time-axis intercept of \( \frac{L^2}{6D_i} \), from which \( D_i \) is calculated.

To describe the statistical significance between the measured concentrations of diffused proteins in hydrogel slices and the distance within the scaffold, we used the non-parametric measure of Spearman’s Rank correlation coefficient. This measure is best suited if the relation between the two variables is not linear. The diffusion model across the diffusion chamber with the experimental conditions applied is best fit using exponential curve decay kinetics (data not shown). Using GraphPad Prism (version 6), we applied correlation testing of 72 hours mRFP1 and P21-mR-8R diffusion data in cellular and acellular scaffolds. Setting the significance level (alpha=0.01) and determining the probability values for each set of data, we demonstrated that a perfect negative correlation was detected in presence of cells for the transducing P21-mR-8R \( (r_s=-1, P<0.0001) \) suggesting the cellular uptake altered the relatively weak non-significant correlation detected in acellular counterpart \( (r_s=-0.6573, P=0.0238) \). In the case of non-transducing protein; mRFP1, the correlation was not significant at the selected significance level in both conditions; cellular and acellular due to minimal cellular uptake and relative free diffusion at the test time point (72 h).
11.3. Expression and Purification of Recombinant Proteins

We obtained cDNA for *mRFP1* (*mR*) as a kind gift from Prof. R. Y. Tsien (University of California, USA) [26]. *8R, P21* and *MYOD* cDNAs were synthesized *de novo* (Eurofins MWG Operon). We cloned cDNAs into the pGEX6-P1 expression vector (Novagen) to create in-frame fusions and expressed proteins in BL21 (DE3) pLysS *Escherichia coli* (Novagen). Exponentially growing LB cultures (OD\textsubscript{600} = 0.4) shaken at 220rpm at 37°C were induced using 1 mM IPTG for 24 hours at 25°C. We lysed and sonicated (7 amplitudes, 1 minute, 5 times) bacterial pellets in 1X STE extraction buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 1 mM EDTA containing 1mM DTT, 0.2 mg/ml lysozyme, and 1X protease inhibitor cocktail). Insoluble protein was retrieved using the Rapid GST inclusion body solubilisation and renaturation kit (AKR-110; Cell Biolabs, Inc., San Diego, CA). We purified recombinant proteins by affinity chromatography using Glutathione-Sepharose resin (GE Healthcare). We removed GST-tags and eluted from resin by PreScission\textsuperscript{TM} Protease cleavage (GE healthcare) in 1X cleavage buffer (50 mM Tris-HCl pH 7.0, 150 mM NaCl, 1 mM EDTA and 1 mM DTT). We determined protein concentration using a BCA-based protein assay (BioRad) with absorbance measured at 595nm using recombinant mR protein as a standard. Integrity and full-length protein expression was confirmed by SDS-PAGE. We determined the fluorescence of recombinant proteins (excitation: 584 nm; emission: 607 nm) with all preparations <10% intensity difference between samples (fluorescence/µg). Standards and samples were analysed using the TECAN infinite 200PRO multimode reader.
2. Supplementary Figure Legends

Figure S1. Compartmentalized diffusion chamber and Acetal resin mould (ARM) design. A) Schematic view of acetal resin mould (ARM) with PTFE spacer fitted inside between which the hydrogel is cast. B) Side schematic view of the ARM design showing mould dimensions. C) Top schematic view of the ARM showing sink (Si), source (So) and gel (G) compartments with their relevant dimensions. D) Top view of compartmentalized ARM diffusion chamber showing hydrogel membranes of
15×5×6.6 mm (Length × Width × Height) dimensions cast in the middle of the chambers.

Figure S2. Faithful diffusion of Fluorescein using the ARM compartmentalized diffusion chamber. A) Representative images of diffusion of 32.15 µg/mL Fluorescein in 1% w/v agarose hydrogel scaffolds of 15×5×6.6 mm
(Length \times Width \times Height) dimensions with Fluorescein diffusing from the Source (So) compartment through the hydrogel scaffold at 0, 2, 6, 16 and 24 hours. B) The corresponding concentration profiles assessed as $C/C_0$ as a function of distance away from the source ($n=3$).

**Figure S3. Assessment of cellular health in different GA hydrogel formulations.**

Different GA hydrogels (containing NIH 3T3 mouse fibroblasts; $2\times10^6$/mL) were set
up (% is w/v) and A. viability assessed over 3 days using Live/Dead cytotoxicity assay. B. Metabolic activity assessed over 3 days using Presto® blue assay. Data was normalised to a monolayer control. (n=3, bars are standard deviation). ** p<0.01 for day 2, * p<0.05 for day 1.

**Figure S4. Determination of the characteristic features of GA hydrogels and the test proteins.** A) transmission electron micrograph image of ultrathin section (80 nm thick) gelatin/agarose hydrogel showing the nanoporous structure. Labels represent randomly selected pores in the TEM image for measurement of the, average pore diameter using ImageJ software, from which the average pore diameter is calculated to be 0.198±0.079 µm. B) table showing the practical estimation of the hydrodynamic radius (nm) of the test proteins in 10 mM HEPES buffer pH7.2 at 37°C using Zetasizer Nano ZS, their theoretical molecular weights and isoelectric points (* using the isoelectric point calculator). C) Estimation of the practical molecular weight of the test protein using SDS-PAGE.
Figure S5. Distribution analyses for cells in three dimensions. A) GA hydrogels (containing NIH 3T3 mouse fibroblasts; 2x10^6/mL) were set up with the standard dimensions shown. Cellular GA scaffolds were sectioned against the B) x-axis (5 sections of 1mm), C) y-axis (6 sections of 1mm) or D) z-axis (7 sections of 2mm). Cells were extracted and counted. Numbers are expressed as a % of cells numbers expected from the gel volume sectioned if distributed uniformly (n=3, bars are standard deviation).
Figure S6. Determination of the hydrodynamic radius of test proteins. The practical estimation of the hydrodynamic radius (nm) of the test proteins in 10 mM HEPES buffer pH7.2 at 37°C using Zetasizer Nano ZS showing the traces obtained. Traces for: A) mRFP1, B) P21-mR-8R, C) P21-mR, D) mR-8R, E) P21-mR-MyoD-8R, F) control protein FITC-BSA, G) Standard divinyl nezene polymer spheres for calibration, are shown.
Figure S7. Monitoring of protein concentrations in the source and sink compartments. GA hydrogels (containing NIH 3T3 mouse fibroblasts; 2x10^6/mL) were set up and source and sink protein concentrations analysed at 24 and 72 hours by fluorometry. (n=3, bars are standard deviation)
Figure S8. Calculation of the diffusion coefficient of mRFP1 and P21-mR-8R in acellular hydrogels using membrane permeation method. Inserts represent linear regression of the same set of data after 24 h, from which the diffusion coefficients of mRFP1 and P21-mR-8R in GA are determined to be $7.15 \times 10^{-7}$ & $7.53 \times 10^{-7}$ cm$^2$/s respectively.
Figure S9. Flow cytometry showing gradients of fluorescence intensity in cells as a function of distance from Source. GA hydrogels were exposed to non-transducing mRFP1 and enhanced transducing P21-mR-8R. Gels were sliced into 1mm sections and liquefied. Representative cytometry dot-plots of digested hydrogel slices retrieved from cellular scaffolds (NIH 3T3 mouse fibroblasts, 2x10^6/mL) exposed to media, mRFP1 or P21-mR-8R respectively after A) 24 hours or B) 72 hours.
Figure S10. Image analyses of pixel mean and standard deviation for mRFP1 and P21-mR-8R in cellular GA scaffolds. (n=3, bars are standard deviation).
Figure S11. Efficiently transducing GET-MyoD maintains a gradient profile in cellular scaffolds. A) Schematic of P21-mR-MyoD-8R (GET-MyoD) recombinant protein used in diffusion studies. Diffusion gradients of GET-MyoD (60 µg/mL) across B) acellular and C) cellular (NIH 3T3 mouse fibroblasts, 2x10^6/mL) GA hydrogels at 6, 24, 72 and 96 hours (n=3). An exponential decay model to assess the gradient is detailed for the 72 and 96 hour time-points.
Figure S12. Calculation of the diffusion coefficient of GET-MyoD (P21-mR-MyoD-8R) in acellular hydrogels using membrane permeation method. Insert represents linear regression of the same set of data after 42 h, from which the diffusion coefficient of P21-mR-MyoD-8R in GA is determined to be $4.2 \times 10^{-7} \text{ cm}^2/\text{s}$. 

$y = 0.2504x - 6.915$

$R^2 = 0.9667$
Table S1. Calculation of Spearman’s Rank correlation coefficient for acellular and cellular GA hydrogels for mRFP1 or P21-mR-8R gradients at 72 hours.

Details of the calculation are described in supplemental methods.

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<th>Cellular</th>
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Equation 1. Determination of the diffusion coefficient of protein macromolecules in GA hydrogels using model permeation method [16].
Graphical Abstract.