



Cite this: *Biomater. Sci.*, 2024, **12**, 2885

Developing an *in vitro* model of haematoma for study of intracerebral haemorrhage†

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Intracerebral haemorrhage (ICH) is a devastating neurovascular attack with limited treatment options. Alternative, pre-clinical modelling approaches are required to identify and trial therapeutic drug compounds. In this study we have used alginate hydrogels to model blood insult *in vitro*. Human whole blood was mixed with alginate and encapsulated into hydrogel beads. Beads were then incorporated in a second layer of alginate containing hyaluronic acid/chitosan nanoparticles to mimic the mechanical properties of brain tissue and create a model haematoma. Beads and model haematomas were characterised to profile size, volume, mechanical properties, release capacity and storage stability over time. Beads and model haematomas stimulate a pro-inflammatory phenotype in human monocytic and macrophage-like cells, however have no pathogenic effect on brain endothelial and neuronal cell survival or function. In conclusion, we have developed an effective strategy to model ICH *in vitro*, to investigate the human immune response to blood insult.

Received 9th January 2024,
Accepted 17th April 2024

DOI: 10.1039/d4bm00039k

rs.c.li/biomaterials-science

Introduction

Intracerebral haemorrhage (ICH) occurs when blood vessels in the brain spontaneously rupture and release an insult of blood into the brain parenchyma.¹ This devastating neurological damage causes widespread cell death and inflammation resulting in disability and mortality, and there are no specific therapies for patients. Primary injury associated with the rupture of blood into the brain causes swelling, increased intracranial pressure, oedema and compression of surrounding brain structures. After the bleeding stops the haematoma condenses, red blood cells lyse, and there is a release of toxic blood compounds and breakdown products into the brain tissue.² Best clinical practice includes surgically removing the haematoma to reduce the mass effect for better clinical outcomes³ but this

is only possible in ~10% of patients.⁴ There has been little investigation into how the extravasated blood damages cells when left to degrade and reabsorb.

Commonly, modelling intracerebral haemorrhage pre-clinically relies on the use of animals to recreate the complex, multi-system disease and investigate behavioural outcomes.⁵ Despite improving our understanding of disease mechanisms, pre-clinical animal studies have been limited in translation of outcomes to the clinic.⁶ As such, there is an urgent need for novel, complementary models⁷ to address specific questions about blood toxicity and the localised human immune response.

Human-relevant data has historically come from clinical observations, patient peripheral blood samples, or analysis of post-mortem tissue. Such approaches limit the information available to specific time frames and holds little relevance for understanding the immediate pathology caused by extravasated blood. Traditionally, *in vitro* models of ICH recreate injury by adding haemoglobin,⁸ hemin,^{8,9} glutamate¹⁰ or hydrogen peroxide to damage cells and the advantages and limitations have been reviewed elsewhere.¹¹ This approach does not recreate the full complement of factors in the blood that contribute to primary cell damage and stimulate secondary inflammatory injury.¹² Neurotoxic factors from blood include thrombin, haemoglobin, free iron, fibrinogen, complement, leukocytes, platelets, oxyhaemoglobin, methemoglobin and hemin.¹³ Adding whole blood to cell culture introduces practical problems due to opacity and reduction of available nutrients and oxygen in the media with increasing blood doses. To overcome these problems, in this study we have used

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4bm00039k>



alginate-based hydrogels to encapsulate human blood to control injury in a cell culture system.

Alginate is a polysaccharide derived from brown algae that has been widely used in biomedical applications and tissue engineering due to structural similarities with extracellular matrices (ECM).¹⁴ Alginate is composed of different sequences of β -D-mannuronic acid (M) and α -L-guluronic acid (G) monomers determined by the species of source algae used. Alginate hydrogels are also used as drug delivery systems, enabling the release of both small chemicals and biomolecules in a controlled manner dependant on crosslinking methods.¹⁵ Alginate hydrogels form easily with the addition of divalent cations, such as Ca^{2+} , upon which G monomers are bound into 'egg box' structure.¹⁶ The G/M ratio is often used to determine gelation, crosslinking density and the resulting mechanical properties of the formed hydrogels. In this study, we investigated 4 types of alginate and different concentrations of CaCl_2 for crosslinking, to encapsulate whole blood and characterise breakdown product release for ICH modelling *in vitro*. This is the first study, to our knowledge, that investigates whether alginate-based hydrogels can provide controllable release of blood breakdown compounds *in vitro* that mimic the cellular toxicity following an ICH in a culture system.

To further model characteristics of the brain tissue, alginate hydrogels containing hyaluronic acid (HA) were characterised to identify mechanical properties that mimic the cerebral ECM. HA was selected as a crucial component of the brain ECM, providing a backbone structure for HA-binding lecticans.¹⁷ In addition to providing biomechanical integrity, HA alters tissue hydration and plays a key role in synaptic plasticity,¹⁸ stroke recovery¹⁹ and inflammation.²⁰ We incorporated HA into the model haematoma to promote hydration of the gel, and to provide structure that would mimic the soft, damaged, oedemic brain tissue following a haemorrhage that surrounds a haematoma.²¹ In polymeric form, HA in hydrogel absorbs water and weakens the network, making the hydrogel unstable within 24–48 hours in cell culture.²² In this study we have fabricated nanoparticles from HA and chitosan (CS), to stabilise the hydrogel whilst maintaining HA in the network as an essential component of the ECM. HA forms a corona around the nanoparticle^{23,24} and ensures hydrogel stability by reducing hydration.

This methodology is highly adaptable and offers an entirely human, *in vitro* platform for interrogating the mechanistic pathology of ICH, and screening neuroprotective compounds that may have translational benefit for the clinic.

Results

Differences in alginate composition affect inflammation in culture

To investigate the effect of different alginate compositions on inflammation, beads generated from 10 mg mL^{-1} alginate hydrogel (AB) were firstly formed using 4 different alginate types varying in G/M ratio and molecular weight (MW).

Different alginate-only beads were cultured for 6 hours with THP-1 cells to determine inflammatory stimulation (Fig. 1A). A panel of primers were used to detect differential gene expression from the HEPES buffered saline (HBS) control indicative of proinflammatory phenotypes. Phosphate buffered solutions cannot be used as phosphate anions degrade alginate gels by sequestering Ca^{2+} ions from the matrix. Reverse transcriptase-qPCR analysis of gene transcription was performed to determine whether alginates stimulated a pro-inflammatory response in THP-1 cells. Tumour necrosis factor (TNF) is a pro-inflammatory cytokine involved in regulating macrophage function and cell survival. Interleukin-1 β (IL-1 β) is a potent pro-inflammatory cytokine that is stimulated in response to injury and drives downstream transcription of other pro-inflammatory molecules. Data showed that A3 was proinflammatory and A1 produced a variable 2-fold increase of TNF transcription however this was not significant from the HBS control. A2 did not produce an inflammatory response however, was excluded from further experiments because of practical difficulty with filter sterilisation due to high viscosity. A4 consistently showed the least inflammatory activation (Fig. 1A) Interleukin-6, arginase 1, interleukin-10, nitric oxide synthase-2 and CC motif chemokine ligand 18 were all undetected confirming the absence of an anti-inflammatory phenotype.

To confirm whether the G : M ratio is as different as reported in A1 and A4, proton nuclear magnetic resonance ($^1\text{H-NMR}$) analysis was performed. The ratio is known to influence physicochemical and mechanical properties of alginate hydrogels.²⁵ $^1\text{H-NMR}$ shows ratio G : M is 2.1 in A1 and 1.3 in A4 (Fig. 1B). The high M content of A4 is desirable to make soft hydrogels with properties that mimic brain tissue ($\sim 1 \text{ kPa}$).²⁶

Synthesising a hyaluronic acid-alginate layer

In physiological conditions, the haematoma is surrounded by damaged tissue and oedema is observed by CT scan,²⁷ likely to contain disrupted ECM and cell matter. This provides a physical layer between the healthy tissue and contact with the blood in the haematoma. In order to generate a model of haematoma that mimics human disease physiology, we sought to introduce hyaluronic acid (HA) to alginate hydrogels to recreate the layer of water-dense damaged brain ECM between the haematoma and cells in culture. Hyaluronic acid has been previously reported to be unstable in water after 24–48 hours due to water absorbance.²² To mitigate instability in water, hyaluronic acid/chitosan (HA/CS) nanoparticles were prepared and mixed with A4 for hydrogel formulation.²⁴ HA/CS nanoparticles were made using two MW of hyaluronic acid: 35 kDa (HA_{35}) and 180 kDa (HA_{180}). HA_{35}/CS and $\text{HA}_{180}/\text{CS}$ nanoparticles were consistent in Z-average size ($\sim 210 \text{ nm}$) and polydispersity index ($\text{PDI} < 0.25$) (Fig. 2A). Of note, HA_{35}/CS nanoparticles showed more uniformity ($\text{PDI} = 0.10$) than $\text{HA}_{180}/\text{CS}$ ($\text{PDI} = 0.23$). Alginate HA/CS gels were prepared by mixing 10 mg mL^{-1} A4 solution with 1 mg mL^{-1} HA/CS nanoparticles solution at 1:1 volume ratio, and hydrogels obtained after incubation with 75 mM CaCl_2 (aq.) for 10 min at RT. CaCl_2 concentration was optimised to generate alginate hydrogel



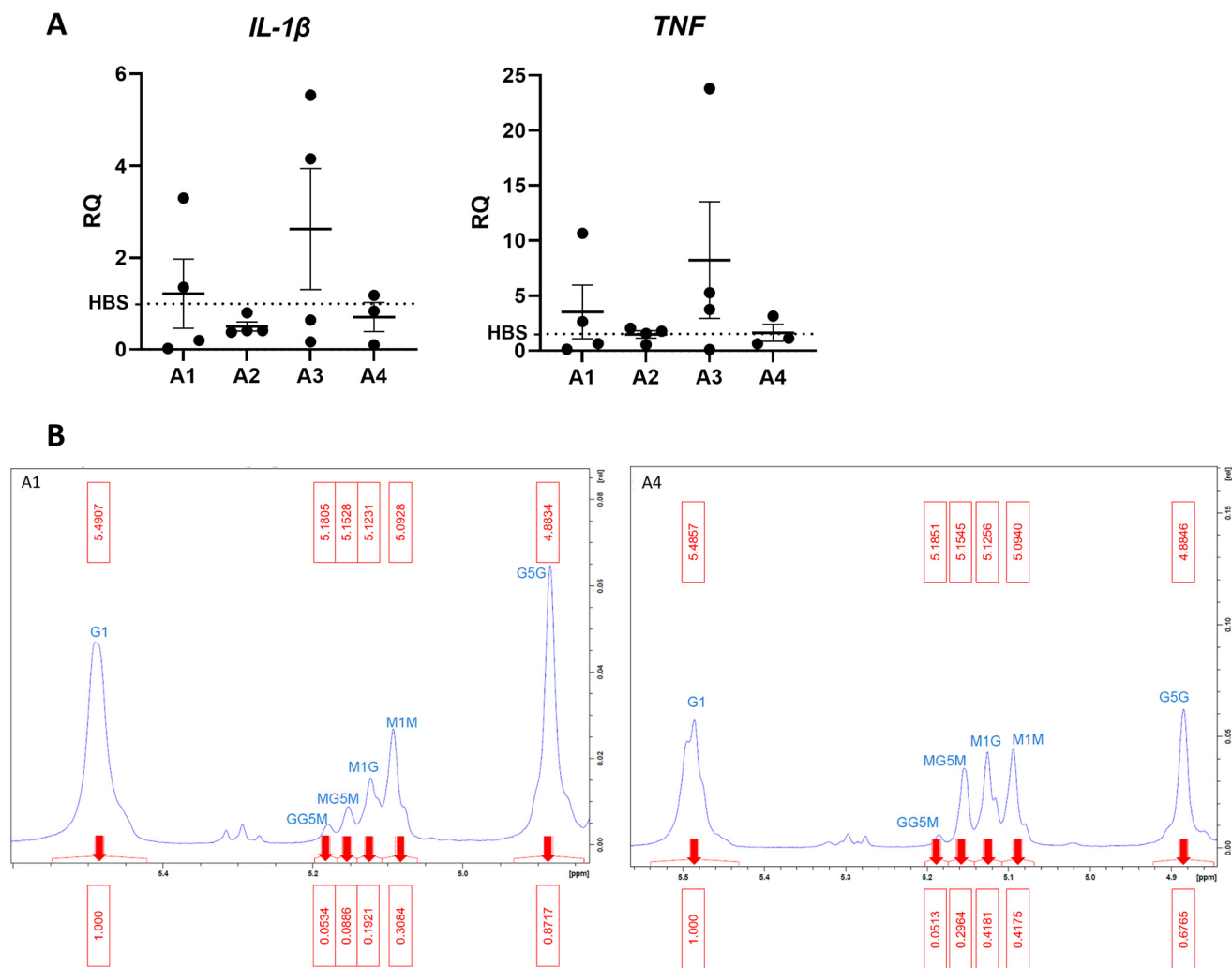


Fig. 1 Investigating the differences between alginate formulations (A) qPCR detecting human IL-1 β and TNF expression in monocytic THP-1 cells as a marker for inflammatory stimulation for 4 different alginates ($n = 4$). (B) Annotated ^1H -NMR results show the GM ratio in each selected alginates A1 and A4. RQ: relative quantification.

containing HA/CS nanoparticles that had a stiffness close to 1.1 kPa, the reported stiffness of grey matter.²⁶ Rheological analysis of the elastic modulus (G') and viscous modulus (G'') was used to determine material stiffness. Data revealed that A4-HA₃₅/CS hydrogel has an average elastic modulus of 1.12 kPa \pm 0.25 s.d (Fig. 2B) compared to A4-HA₁₈₀/CS hydrogel which is 1.51 kPa \pm 0.27 s.d (Fig. 2C). Therefore, A4-HA₃₅/CS hydrogels were selected for the stiffness most closely fit with grey matter and used for the rest of the experiments. Data from further experiments with A4-HA₁₈₀/CS hydrogels is available in ESI Fig. 1.†

Alginate blood beads are consistent in fabrication, but release blood in a short time period

Beads generated from 10 mg mL⁻¹ alginate hydrogel mixed with whole blood to form blood beads (BB) showed no difference in size or volume from alginate-only beads (AB) (Fig. 3A–D). On average, beads are 250 μm in diameter and have a

volume of 0.007 mm³. Constant values were observed with different ratios of blood:alginate for bead preparation however protein release was accelerated at higher ratios. Blood beads were made 50:50 v/v blood and 10 mg mL⁻¹ alginate for all further experiments as beads had good stability and blood content. Blood beads showed evidence of degradation and protein release into the buffer after 24 hours at 4 °C (Fig. 3E). This varied between preparation repeats however did not reach levels of significance. After 3 weeks stored at 4 °C, blood beads had released ~15% (Fig. 3F) of the hemin levels that were detected in the dissolved blood beads, implying that although blood beads are initially leaking protein from breakdown products, hemin can still release after 3 weeks to initiate a pathogenic response.

Generating a model haematoma adds stability to beads

Alginate-only beads and blood beads were incorporated into A4 HA₃₅/CS hydrogel, final concentration of 5 mg mL⁻¹ ALG4





Fig. 2 Hyaluronic acid/chitosan nanoparticles incorporated into alginate cell forms a layer with a similar stiffness to brain tissue. (A) Dynamic light scattering measurements of HA₃₅/CS and HA₁₈₀/CS nanoparticles, *n* = 4 independent repeats. (B) Shear moduli for A4 HA₃₅/CS hydrogel crosslinked using 75 mM CaCl₂ solution (aq.) (*n* = 3 independent repeats) (C) Shear moduli of A4 HA₁₈₀/CS hydrogel crosslinked using 75 mM CaCl₂ (*n* = 3 independent repeats). PDI: polydispersity index, LVR: linear viscoelastic region, G': elastic modulus, G'': viscosity modulus.

and 0.5 mg mL⁻¹ HA₃₅/CS nanoparticles, prior to extrusion and gelation in a 75 mM CaCl₂ solution (aq.) at RT to form model haematomas (controls with alginate-only beads = MHA, and with blood beads = MHB) (Fig. 4A and B). Model haematomas are on average 2 mm in diameter with a volume of 15 mm³ (Fig. 4C) and on average contain the same number of beads regardless of blood, however there is a much larger range of number of alginate-only beads encapsulated within the hyaluronic acid layer (Fig. 4D). This may be due to difficulty identifying alginate-only beads from contrast images. To confirm limited swelling of the hyaluronic acid-containing hydrogel we determined that there is no significant change in volume over 1 week in storage at 4 °C (Fig. 4E). By measuring the protein content of the storage buffer as an indication of blood breakdown release, data revealed that MHB can be stored for ~3 weeks before significant protein detection (Fig. 4F). Mean protein concentrations at 3 weeks were half of detected levels for blood beads without the A4 HA₃₅/CS hydro-

gel layer (308 mg mL⁻¹ compared to 712 mg mL⁻¹, Fig. 1E). Detection of signal in control model haematoma samples at 3 weeks perhaps indicates a breakdown in the polysaccharide in the HA₃₅/CS hydrogel and interference in the BCA detection assay. To ensure model consistency, model haematomas were formed within 24 hours of blood donation and bead formation and were used for further investigation and cell culture within 2 weeks of fabrication. Analysis of hemin concentration in the buffer and dissolved MHB after 3 weeks reveals that MHB only had about 2% of the total hemin detected in whole blood samples (Fig. 4G) whereas the blood beads had equal amounts of hemin as whole blood (Fig. 1F). This implies a faster breakdown of the haemoglobin and loss of hemin than in blood beads.

Beads release toxic compounds in a dose dependent manner

Blood beads were added in increasing volumes to wells for a 24 h incubation at 37 °C to show a dose-dependent effect on protein release (Fig. 5A). This resulted in dose-dependent cell





Fig. 3 Whole blood can be mixed with alginate to form consistent, but quick releasing beads. (A) Images of alginate-only (AB) and (B) blood beads (BB) in 6 well plate and under 10 \times magnification. Scale bars 500 μ m (C) Diameter and (D) volume of AB and BB ($n = 20$). Box shows median value and extends to 25/75%, whiskers min/max values. (E) Protein concentration measured by BCA assay in the HBS that beads were stored in at 4 $^{\circ}$ C ($n = 3$) mean \pm s.d. (F) Hemin measurements from whole blood and storage buffer and dissolved BB after 3 weeks at 4 $^{\circ}$ C ($n = 2$).

death in macrophage-like THP-1 cells when cultured for 24 hours (Fig. 5B). The largest influence on cell death was caused by the number of blood beads, and to a lesser extent, the concentration of CaCl₂ used to crosslink the alginate regardless of alginate type (A1 and A4). Blood beads made with A4 and 0.25 M CaCl₂ were selected for further experiments and dosed at 50 μ L into 200 μ L wells.

In order to determine what factors released from the beads in culture are causing toxicity, further analysis was performed

to analyse the contents of the media that may be damaging the cells. Reactive oxygen species (ROS) are released from the haematoma and from surrounding damaged cells. After 24 hours at 37 $^{\circ}$ C detectable ROS levels from blood beads and model haematomas with blood beads were 100 \times less than undiluted plasma at 2.52 μ M and 2.89 μ M respectively (Fig. 5C). Hemin is commonly used to damage cells in culture and previous studies in the literature have shown 100 μ M of hemin results in 50% neuronal cell death.⁸ Fig. 5D shows





Fig. 4 Encapsulating BB into A4 HA₃₅/CS hydrogel slows blood breakdown release. (A) Images of AB (red arrows) and (B) BB within the A4 HA₃₅/CS hydrogel layer in a 6 well plate and under 2.5× magnification. Scale bars 500μm (C) average volume of the A4 HA₃₅/CS hydrogel with either AB or BB. (D) Average number of beads encapsulated in the A4 HA₃₅/CS hydrogel MH. (E) Average volumes of MH over 1 week in storage at 4 °C. (F) Protein release into buffer over 3 weeks storage at 4 °C (*n* = 3). Bars show average and s.d. Data was analysed using a one-way ANOVA for each time point to determine differences from HBS control (***P* = 0.005). (G) Hemin concentrations in buffer and dissolved gels after 3 weeks (*n* = 2).

detection was greater from the model haematomas with blood beads than blood beads, and there was less of a difference dependent on time of incubation. Hemin released from the model haematomas with blood beads was ~20 μM, almost half of the levels detected in plasma. Blood beads released ~10 μM. Haemoglobin, another toxic compound released from the haematoma when red blood cells lyse, was detectable at the same levels as plasma after 24 hours (Fig. 5E). Our study shows that beads release 130 μM haemoglobin after 24 hours in culture,

and plasma contains as much as 170 μM. Data suggests that there is a faster degradation of haemoglobin in model haematomas with blood beads than in blood beads, and this results in the quick release of hemin which breaks down and is undetectable after storage for 3 weeks.

Human fibrinolysis LEGENDplex assay analysis revealed that when incubated at 37 °C for 6 hours, blood beads and model haematomas with blood beads released significantly less fibrinogen, anti-thrombin, factor XIII, IL-6 and tissue





Fig. 5 Blood beads release cytotoxic factors (A) Protein concentration released at 24 hours in 37 °C in a dose dependent manner. (B) Concentration CaCl_2 solution (aq.) and of volume of BB solution added, has a dose response effect on macrophage-like THP-1 cell death ($n = 3$) after 4 hours incubation. (C) ROS detection after 4 and 24 hours in 37 °C ($n = 3$). (D) Hemin detection in the buffer after 4 and 24 hours at 37 °C ($n = 3$). (E) Haemoglobin detection after 4 and 24 hours in 37 °C ($n = 3$). (F) LEGENDplex (Fibrinolysis 5 plex and Thrombosis 3 plex) assays determine concentration in the media after 4–6 hours at 37 °C with BB and MHB ($n = 3$, $n = 1$ for plasma reading of thrombosis factors). Data analysed using a one-way ANOVA and comparisons made to plasma value.



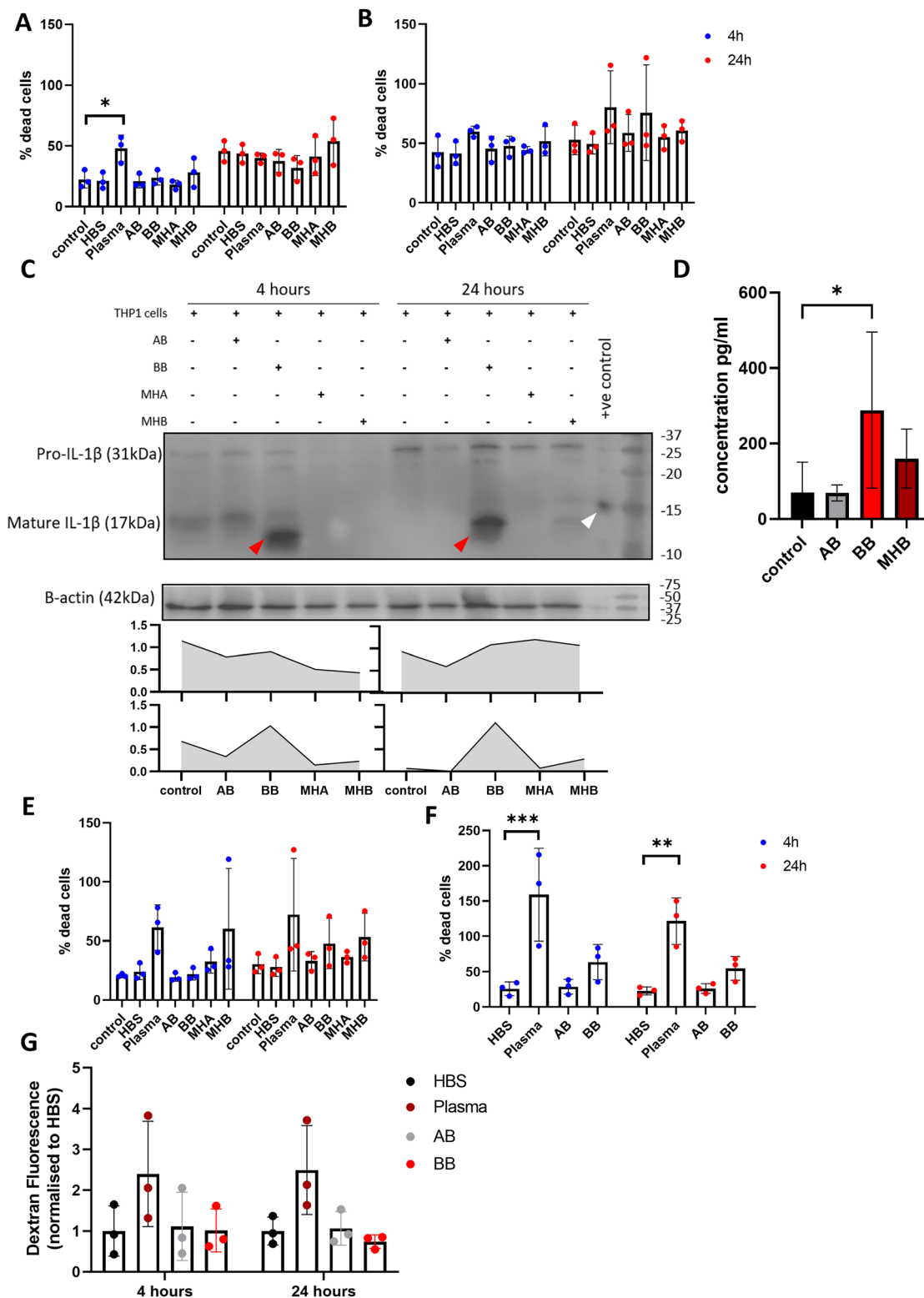


Fig. 6 Cellular response to blood beads and model haematomas with blood beads incubation. LDH cell death assay of (A) monocytic THP-1 cells, (B) macrophage-like THP-1 cells when incubated with BB/MHB for 4 and 24 hours ($n = 3$). $*P = 0.0407$ (C) western blot analysis and quantification of bands determined by ratio to β -actin. Pro-IL-1 β increases in THP-1 cells after incubation for 24 hours, but no difference in mature IL-1 β over time. BB incubation stimulates IL-1 β maturation (red arrows, positive control is denoted by a white arrow) ($n = 3$) (D) ELISA analysis of IL-1 shows significant increase in macrophage-like THP-1 cells with BB, and to a lesser extent with MHB culture after 24 hours ($n = 4$) $*P = 0.0467$. (E) LDH cell death assay of SH-SY5Y neuronal cells and (F) HMBECs when incubated with BB/MHB for 4 and 24 hours ($n = 3$). $**P = 0.0033$, $***P = 0.0002$. (G) Quantification of dextran leakage through HMBECs layer on culture inserts showed no effect of BB/MHB on the function of the cells to establish tight junctions.



factor than found in donor plasma (Fig. 5F). Levels of plasminogen were not significantly different from plasma, and prothrombin released from blood beads was significantly higher than plasma controls. IL-8 (CXCL8) was not detected in plasma however, was detected to be released from the models. IL-8 is a chemokine that promotes angiogenesis and pro-inflammatory roles that damage the endothelial cells, implying perhaps donor leukocytes survive and are active within the hydrogel. These data imply that blood beads exhibit fibrinolytic and thrombotic factors which may contribute to the toxic effects observed in culture however this is less than detected physiologically.

Characterising the cellular response to toxic factors

To investigate the effect of beads in culture with different cell types, 50 μL blood beads and 3 \times model haematomas with blood beads were added to monocytic THP-1 cells in suspension, stimulated macrophage-like THP-1 cells, SH-SY5Y neuronal cells and brain endothelial hCMEC/D3 cells. Blood beads and model haematomas with blood beads had no effect on cell death of monocytic THP-1 cells or macrophage-like THP-1 cells at either time point (Fig. 6A and B). IL-1 β is activated and released in a two-step process and is a potent down-stream pro-inflammatory cytokine. Western blot analysis showed that monocytic THP-1 cells produced pro- and mature IL-1 β in response to blood beads (Fig. 6C), however this was undetected by ELISA and therefore unreleased from cells. Macrophage-like THP-1 cells release IL-1 β detected by ELISA when treated with blood beads, and a lesser extent with model haematomas with blood beads (Fig. 6D). In literature, it has been observed before that heme,²⁸ red blood cells and haemoglobin²⁹ all induce IL-1 β release from phorbol-12-myristate-13-acetate (PMA) – stimulated THP-1 cells. This implies that perhaps the blood insult is the first priming step for IL-1 β maturation, however a second activation stimulus is required for release, such as the PMA stimulation of cells.³⁰ These results support the conclusion that blood beads and model haematomas with blood beads is a good model to study the immune response to haemorrhage. Blood beads and model haematomas with

blood beads had no effect on cell death of SH-SY5Y neuronal cells or hCMEC/D3 cells at either time point (Fig. 6E and F). SH-SY5Y cells did not produce or release any IL-1 β nor was IL-6 and TNF α detected in any of the cell types. To determine whether blood beads impacted brain vascular cell function, we conducted a barrier permeability assay on hCMEC/D3 cultures (Fig. 6G). Incubation with blood beads had no effect on the permeability of the cells showing that the barrier function of these cells is not impaired.

Discussion

In this study we have shown that blood can be encapsulated into alginate beads and stored for up to 24 hours. When introduced to cell culture, blood beads were proven to mimic the toxicity of blood on cells in a dose-dependent manner. This model presents a novel method of investigating pathophysiology of blood extravasation and the associated immune response that can be applied with medical relevance for pre-clinical modelling of ICH. Application of such a model would allow for investigation of pathology in a human system, to screen compounds for anti-inflammatory or cyto-protective effects.

When considering the use of this model in the pre-clinical ICH pipeline, it is important to compare with the current *in vitro* models commonly used. Hemin and haemoglobin, as breakdown products of RBCs within the haematoma, have been previously used to model ICH *in vitro*. There is limited understanding of pathology when trying to recreate ICH *in vitro* as dosing cell culture with individual factors does not include the full neurotoxicity of whole blood. In this study we have shown that after 4 hours of incubation at 37 $^{\circ}\text{C}$, blood beads released 8 μM hemin and 117 μM of haemoglobin. Model haematomas with blood beads released 20 μM of hemin and 11 μM of haemoglobin. Previously, researchers have shown that similar doses of hemin have had considerable deleterious effect on cell survival and are capable of stimulating ROS release from cells (Table 1). In our study we showed

Table 1 A comparison of the outcomes of hemin and haemoglobin treatment in cell culture

Neurotoxic factor	Dose	Cell type	Outcome	Ref.
Hemin	20 μM for 2 hours	THP-1	40% reduction in cell viability 8 \times increase in ROS production	Imoto <i>et al.</i> 2018 ³¹
Hemin	100 μM for 14–18 hours	Primary mouse neurons	50% reduction in cell viability	Zille <i>et al.</i> 2017 ⁸
Haemoglobin	1.5 μM for 24–28 hours	Primary mouse neurons	25% reduction in cell viability	Zille <i>et al.</i> 2017 ⁸
Hemin	5 pg mL^{-1} LPS for 24 hours 10 μM hemin	Macrophage-like THP-1	No cytotoxic effect Reduced glycolysis, HIF-1 α , cell metabolism and phagocytosis	Carrasco-Pozo <i>et al.</i> 2020 ³²
Hemin	0.01–1 $\mu\text{g mL}^{-1}$ LPS for 3 hours 30 μM hemin 4 μM ATP for 1 hour	Human primary macrophages Macrophage-like THP-1 Mouse bone marrow-derived macrophages	1000 pg mL^{-1} IL-1 β release NLRP3 inflammasome activation No change in ROS Increased autophagy	Nurmi <i>et al.</i> 2017 ³³



that blood beads stimulated vastly more ROS release than in previous studies,³¹ released 10× the concentration of haemoglobin used in literature⁸ and was capable of initiating IL-1β release without a liposaccharide (LPS) priming step, which highlights the discrepancies between using single neurotoxic factors and whole blood at initiating a translational injury response that recreates human disease.

Formulation of the beads in this study was simplified for selection however, there are many parameters that can be further altered to investigate the influence on size, volume, permeability, and stability of the beads and subsequent outcomes in culture. In principle, high G/M ratios and high cross-linking cation concentration lead to a high crosslinking density and a stiff hydrogel network which slows the diffusion of loaded molecules. Design criteria in this study focused on the use of non-animal derived polymers, generating mechanical properties to match brain tissue, and encapsulating and releasing human blood. Within whole blood, the pathogenic molecules are a range of sizes and alginate hydrogels typically have a pore size of ~5 nm.³⁴ We selected Ca²⁺ as crosslinking cations based on previous knowledge in modelling mechanics of soft tissues *in vitro*.^{35,36} The size of fabricated blood beads can be increased by using a larger extrusion nozzle, decreasing the vibration frequency and increasing the flow rate during encapsulation.³⁷ Alginate type and concentration influences the diffusion of large molecules through the hydrogel matrix.²⁵ We used alginates with different G/M ratios to best match the stiffness of soft tissues.³⁸ With this strategy we demonstrated that high M alginates (A4) form softer hydrogels with properties similar to brain tissue stiffness and that are also suitable for diffusion of blood components.

We investigated the use of hyaluronic acid-coated chitosan nanoparticles for the inclusion of hyaluronic acid in alginate hydrogels as a component of the ECM, to overcome well known issues of network instability with polymeric hyaluronic acid. We used two different molecular weights of hyaluronic acid and found a slight increase in hydrogel stiffness with nanoparticles made with high MW hyaluronic acid (~1.5 kPa A4-HA₁₈₀/CS vs. ~1.1 kPa A4-HA₃₅/CS). Of note, while using HA₃₅/CS alginate-based hydrogels, volumes remained consistent for the duration of the experiment. As expected, adding a higher volume of blood beads to cell culture resulted in a higher amount of blood breakdown products available for diffusion.

Further use of this model with different cell types, in different culture settings, or with 3D cultures may reveal new insights into the pathology of blood extravasation in the brain and help reveal different proinflammatory and cell damage/death pathways that are activated following ICH. We can use this model to investigate therapies that target specific cell types in a human system, prior to further pre-clinical testing. Investigation into patient-specific responses or using genetic analysis of primary patient-derived cell cultures may identify specific products of injury that may be used as disease biomarkers or for targetable therapies. Further characterisation of this model can reveal insights into the mechanisms behind

haematoma breakdown and resorption, by investigating culture with microglia and neuronal stem cells. Broadly, the encapsulation of blood into alginate can be employed in other culture systems to investigate the cellular response to haematoma, whilst providing a reversible and controllable blood insult.

By adding models to the pre-clinical toolkit for studying ICH, we hope to close the gap between animal models and clinical trial to improve translational outcomes and benefit patients.

Materials and methods

Reagents and materials

Reagents

HEPES buffered saline (HBS)

Recipe

150 mM NaCl
20 mM HEPES
pH 7.4

Materials

Sodium alginate #71238
Sodium alginate-LN
Sodium alginate-LT
Sodium alginate-MC
Hyaluronic acid MW 35 kDa (HA₃₅)
Hyaluronic acid MW 180 kDa (HA₁₈₀),
Chitosan MW 36 kDa #9012-76-4

Supplier

Sigma
PROALG
PROALG
PROALG
Medipol SA
Medipol SA
Sigma

Preparation of alginate

HBS was made fresh and sterilised. A 10 mg mL⁻¹ alginate solution was prepared using the following alginate types:

Notation	Species	Commercial source	Country of origin	Reported G/M ratio
A1	<i>Laminaria hyperborea</i>	Sigma 71238 Lot# BCCD8789	Norway	2.3
A2	<i>Lessonia nigrescens (bertorona)</i>	LN - PROALG	Chile	=1
A3	<i>Lessonia trabeculata</i>	LT - PROALG	Chile	>1
A4	<i>Macrocystis pyrifera</i>	MC - PROALG	Chile	<1

After complete hydration, each solution was warmed to 37 °C and filter sterilised using 0.22 μm polyethersulfone filter (SLGP033RS, Millipore).

Encapsulation of blood in alginate beads

All experiments involving human samples were conducted following the guidelines of ethical involvement of human participants published by the University of Manchester (UK) and with ethical approval from the University of Manchester Research Ethics Committee (#16181-28064). Peripheral blood samples were freely repeatedly donated by 5 healthy volunteers who gave informed consent, in tubes containing 10% anti-coagulant citrate dextrose (ACD) (Sigma C3821) and 0.1% PPACK Dihydrochloride (Millipore 520222) and immediately



processed for encapsulation. Equal volumes of 10 mg mL⁻¹ alginate solution was mixed with blood or HBS for controls, and loaded into a 10 mL syringe and encapsulated into beads as previously described.³⁷ In brief, blood/alginate suspension was extruded at 8.9 mL min⁻¹ through a 150 µm nozzle. Droplets were generated using a vibration frequency of 5500 Hz and the stream was disrupted by electrostatic potential of 1.0 kV. Droplets were collected into a rotating, sterile CaCl₂ bath (speed 100 rpm) and incubated for 10 minutes to allow for ionic crosslinking. Alginate hydrogel beads were then poured through a 50 µm cell strainer and washed twice with HBS. Beads were stored at 10% volume in HBS at 4 °C (2.5 mL of beads and 25 mL of HBS). All blood beads (BB) in experiments were compared to alginate-only beads (AB) formulated at the same time.

Making hyaluronic acid/chitosan nanoparticles

Hyaluronic acid (HA) MW 35 kDa (HA₃₅) and 180 kDa (HA₁₈₀), (Medipol SA) was dissolved in dH₂O to 1.5 mg mL⁻¹ and stored at 4 °C. Chitosan (CS) MW 36 kDa (Sigma 9012-76-4) was made to 0.69 mg mL⁻¹ in 4.6 mM HCl (aq.) and adjusted to pH 5.0. Hyaluronic acid coated chitosan nanoparticles (HA_{MW}/CS) were made as previously described.²³ In brief, 250 µL of CS was added to 250 µL of dH₂O in a round bottomed tube and stirred at 1000 rpm for 10 min under sterile conditions. Following this, 500 µL was then added to 500 µL of HA solution and stirred at 1000 rpm for 30 min at RT. Dynamic light scattering (DLS) using a Malvern Zetasizer Nano ZS and software (Malvern Instruments Ltd, Malvern, UK) was used to determine hydrodynamic diameter and polydispersity index (PDI) at 298 ± 0.1 K, recorded at a backscattering angle of 173 °C.

Protein release assay

Samples for assay were collected from either the storage buffer (HBS) or dissolved alginate samples at the appropriate time point. AB and BB were dissolved using 100 mM HEPES and 500 mM trisodium citrate dehydrate adjusted to pH 7.3. Protein concentration was determined using the Pierce BCA protein assay kit (Thermo Scientific 23225) according to kit protocol. Colourimetric detection of absorbance at 562 nm was performed using the Biotek plate reader and Gen5 software. Experimental replicates (*n*) were defined as beads formed from different blood donors.

Blood breakdown release assays

BB and MHB were incubated in HBS at 37 °C for 4 and 24 hours. Buffer samples were collected and assayed for ROS release (Abcam ab238535), haemoglobin release (Sigma MAK115), hemin release (Abcam ab65332), fibrinolytic and thrombotic factor release using the LEGENDplex panels (BioLegend 740760 and 740908 respectively), all according to kit instructions. Control whole blood/plasma samples were used from the same donation. Detection of absorbance/fluorescence was performed using the Biotek plate reader and Gen5 software. LEGENDplex samples were analysed using the

BD FACSVerser flow cytometer and data analysed using the open source LEGENDplex software (<https://www.biolegend.com/en-us/legendplex?tab=software>). Experimental replicates (*n*) were defined as beads formed from different blood donors.

Proton nuclear magnetic resonance (¹H-NMR) analysis of alginates

Prior to ¹H-NMR analysis, sodium alginate (A1 and A4) was partially degraded by acid hydrolysis as described in standard test method ASTM F2259-10.³⁹ In brief, 0.1% w/v alginate solution was prepared under sterile conditions and adjusted to pH 5.6, then pH 3.8 each time heated to 100 °C for 1 h. Final pH was adjusted to 7.4 and samples were freeze-dried at -80 °C, 0.01 mbar (Scanvac Coolsafe Pro55 with Vacuubrand Vacuum pump). Samples (10 mg) were fully hydrated at RT in pure deuterium oxide (D₂O, 7789-20-0, Acros Organics) and freeze-dry repeated. A 10 mg mL⁻¹ alginate in D₂O solution was prepared and 700 µL was tested. ¹H NMR scans were performed using the Bruker Avance III 400 MHz spectrometer (Bruker) at 78 °C with 30° pulse proton angle, relaxation delay 1 s and acquisition time 3.984 s using a 5 mm prodigy cryo-probe. The total number of scans was 16. Obtained spectra were processed using Bruker Topspin (version 3.6.3). ¹H-NMR traces were annotated as previously described⁴⁰ and the area under the curve was used to calculate the G : M ratio using eqn (1).³⁹

$$\begin{aligned} G &= 0.5[G1 + G5G + 0.5(GG5M + MG5M + M1G)] \\ M &= M1M + 0.5(GG5M + MG5M + M1G) \end{aligned} \quad (1)$$

Rheology

Rheological tests were performed to measure the elastic (*G'*) and viscous (*G''*) moduli of alginate-based hydrogels. Oscillatory rheology tests were performed using a Discovery HR-2 rheometer (TA Instruments, UK) equipped with a 20 mm parallel plate geometry. Frequency sweeps were performed at 0.5% strain, 1–10 rad s⁻¹ frequency range at 37 °C with the gap (plate-to-plate) adjusted to the height of each sample (5 mm). Retrieved data were analysed using DIN 51810-2 recommended settings in RheoWin Data Manager (version 4.87.0002) to determine linear viscoelastic range (LVR).

Model haematomas

In sterile conditions, AB or BB were well mixed and 2 mL of each transferred to a tube. Beads were left to settle and HBS removed. 1 mL of 10 mg mL⁻¹ A4 and 1 mL HA/CS nanoparticles were added and gently mixed until a homogenous suspension was obtained. The solution was loaded into a 2 mL syringe and extruded through a 30G gauge needle into a rotating 75 mM CaCl₂ solution bath, allowing crosslinking at RT. After 10 minutes, MH were poured through a 50 µm cell strainer to remove the CaCl₂ and washed twice with HBS. Model haematomas were stored in 1:10 volume of HBS at 4 °C.



Imaging

Images were acquired using a fluorescent inverted microscope (Leica DMI6000, Leica Microsystems, UK) coupled with a 5.5 Neo sCMOS camera (Andor, UK), and equipped with 2.5× (PLAN 2.5×/0.07, Leica) and 10× objectives (PL 10×/0.3 PH1, Leica). Images were acquired and analysed using μ Manager software (v.1.46, Vale Lab, UCSF, USA) and ImageJ.

Cell culture

The human leukaemia monocytic cell line, THP-1, was cultured in RPMI-1640 medium supplemented with 2 mM L-glutamine, 100 μ g mL⁻¹ penicillin–streptomycin (Pen-Strep, Sigma), and 10% (v/v) heat-inactivated fetal bovine serum (FBS) (Gibco). Cells were grown in T75 flasks and incubated in a humidified 5% CO₂ (v/v) atmosphere at 37 °C and routinely split for optimal culture numbers. Cells between passages 11 and 23 were plated 1 million cells per mL to a 96 well plate the day of loading haemorrhages. THP-1 cells between passages 11 and 19 were plated 1 mil mL⁻¹ to a 96 well plate and differentiated using 250 nM of phorbol-12-myristate-13-acetate (PMA) overnight. Media was changed in the morning when plating the models to PMA-free supplemented RPMI.

Immortalised human neuroblastoma cell line SH-SY5Y, were cultured between passages 13 and 20 were raised in DMEM supplemented with 100 μ g mL⁻¹ penicillin–streptomycin (Pen-Strep, Sigma) and 10% (v/v) FBS. Cells were trypsinized with a solution of 0.5% (w/v) trypsin and 0.2% (w/v) EDTA (Sigma) to split cultures. Cells were plated 0.0375 million cells per mL to a 96 well plate and allowed to adhere overnight. Media was changed the following morning prior to adding the models.

Immortalised human cerebral microvascular endothelial cell line, hCMEC/D3, was purchased from Merck (UK). Cells were cultured between passages 8 and 10, and plated on pre-coated surfaces with rat tail collagen type I (Merck), (1 : 100 in PBS) at 37 °C for 1 h. Cultures were maintained in complete Endothelial Cell Growth Medium MV (PromoCell, C-22020) supplemented with 1 ng mL⁻¹ recombinant human basic fibroblast growth factor (Merck), 100 units per ml penicillin, and 100 μ g mL⁻¹ penicillin–streptomycin (Pen-Strep, Sigma). Cells were seeded in the upper compartment of collagen-coated inserts (24-well, 0.4 μ m pores, PET translucent, CellQART) at a density of 0.5 million cells per ml in 100 μ L 24 h prior to experiments.

Estimation of blood content

For AB/BB treatment, 50 μ L media was removed from 200 μ L in 96 well plates and equal volume of homogenous bead solution added. At 10% v/v density this was estimated to contain ~5 μ L of blood held within the beads. For MHA/MHB treatment, 3× MH were isolated using a micro spatula spoon and added to the wells. MHB were made with 2 mL BB at 10% v/v equalling 200 μ L blood per batch. 2 mL A4-HA₃₅/CS solution resulted in ~100 MH assuming equal distribution of blood beads equals 2 μ L of blood held in each MHB.

Cell death, ELISA and microscopy

At 4 and 24 hours following incubation, 100 μ L of media was removed and analysed for hIL-1B using ELISA kit and instructions (Biotechne DY201). Cell death was analysed using a lactate dehydrogenase (LDH) release assay (CytoTox 96 Promega G1780) on supernatant samples following bead incubation as per the kit instructions. Colourmetric detection of absorbance was performed using the Biotek plate reader and Gen5 software. Experimental replicates (n) were defined as experiments performed on different passages of cells.

Dextran permeability of hCMEC/D3 monolayers

Plasma, AB and BB were added to the lower compartment of the well for 24 h. Following treatments, media was changed to phenol-red free Opti-MEM medium (Gibco) and fluorescein isothiocyanate–dextran 70 kDa (Sigma) was added to the upper compartment at 1 mg mL⁻¹ in a total volume of 100 μ L. 30 min later media was collected from the lower compartment. Fluorescence of the media samples was measured using a Fluorstar reader with excitation at 492 nm and emission at 520 nm (BMG Labtech). Fluorescence was normalised to the buffer control group. Experimental replicates (n) were defined as experiments performed on different passages of hCMEC/D3 cells.

Quantitative PCR

Cells were pelleted and RNA extracted using Trizol and chloroform/isopropanol DNA precipitation. Complementary DNA sequences were generated from 800 ng of RNA and reverse transcriptase from the High-Capacity RNA-to-cDNA kit (Applied Biosystems). Gene expression analysis was performed using RT-qPCR with sample cDNA and SYBRGreen qPCR Master mix (Thermo Fisher Scientific). Primers used for IL-1 β and TNF, are listed below and compared to relative quantities of house keeper genes *HPRT1* and *GAPDH*. Data was acquired using the Applied Biosystems StepOne software and quantified using 2^{- $\Delta\Delta$ C_T}.⁴¹ Experimental replicates (n) were defined from different cell passages.

Primers	Sequence
Fw_hs-HPRT1	CAGGCGAACCTCTCGGCTTT
Rv_hs-HPRT1	GGGTCGCCATAACGGAGCC
Fw_hs-GAPDH	TCGGAGTCAACGGATTTGGTC
Rv_hs-GAPDH	AGTTGAGGTCAATGAAGGGT
Fw_hs-IL1B	ACGATGCACCTGTACGATCACT
Rv_hs-IL1B	CACCAAGCTTTTTTGCTGTGAGT
Fw_hs-TNF	TGCACCTTGAGTGATCGGC
Rv_hs-TNF	AGCTTGAGGGTTTGCTACAAC

Western blot

Cells were incubated with beads and model haematomas for 4 and 24 hours before being pelleted, supernatant removed and dissociated using a 1% Triton X lysis buffer and protease inhibitor. Protein was loaded into 12% resolving gel and separated by SDS-PAGE before blotting using a semi-dry transfer system (BIO-Rad). Primary antibody (goat anti-IL-1 β , RnD Systems) was added 1 : 1000 overnight at 4 °C and a secondary



HRP antibody (anti-goat, Agilent Technologies) (1 : 5000) used to visualise bands with ECL detecting reagent (Amersham) in the G : Box with GeneSys software (Syngene). Membranes were washed and incubated with anti- β actin (Sigma A3854) 1 : 20 000 as a loading control and imaged again. Gel bands were quantified using ImageJ gel analysis tool and area under the curve.

Statistical analysis

All data was analysed using GraphPad Prism. Box and whisker plots show the median value and extend to 25th and 75th percentile, whiskers show the minimum and maximum values. Analysis of multiple groups was performed using a one-way or two-way ANOVA with appropriate multiple comparisons. Error bars show s.d. Analysis of fibrinolytic factor release was performed using unpaired Student's *T*-tests.

Abbreviations

A1–4	Alginate samples 1–4
AB	Alginate-only beads
BB	Blood beads
MHA	Model haematomas made with alginate-only beads and A4 HA ₃₅ /CS gels
MHB	Model haematomas made with blood beads and A4 HA ₃₅ /CS gels
HA ₃₅	35 kDa Hyaluronic acid
HA ₁₈₀	180 kDa Hyaluronic acid
CS	Chitosan
HBS	HEPES buffered solution

Author contributions

SC is responsible for the project conceptualisation, methodology, investigation, data acquisition, analysis, validation and visualisation and the writing of the original manuscript. VST is responsible for the data presented in Fig. 1A and 6G. CB is responsible for the NMR data acquisition and analysis in Fig. 1B. AT is responsible for the conceptualisation of the project, provision of resources and supervision for SC. All authors were involved in the reviewing and editing of the final manuscript.

Data availability

All raw data pertaining to this study is available without restriction from the authors. Data acquired using models fabricated with A4-HA₁₈₀/CS hydrogels is available as ESI Fig. 1.†

Conflicts of interest

The authors declare no competing interests.

Acknowledgements

This study was funded by the NC3Rs (NC/V002082/1) to SC. The authors would like to thank the University of Manchester NMR core facility for use of the Bruker BioSpin spectrometer, Dr Katharina Edkins for use of the Discovery HR-2 rheometer and Dr Gareth Howell and the University of Manchester Genomic Technologies Core Facility for cell sorting for LEGENDplex assays.

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