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Morphology, repulsion, and ordering of red blood cells in viscoelastic flows under confinement†

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Red blood cells (RBC), the primary carriers of oxygen in the body, play a crucial role across several biomedical applications, while also being an essential model system of a deformable object in the microfluidics and soft matter fields. However, RBC behavior in viscoelastic liquids, which holds promise in enhancing microfluidic diagnostic applications, remains poorly studied. We here show that using viscoelastic polymer solutions as a suspending carrier causes changes in the clustering and shape of flowing RBC in microfluidic flows when compared to a standard Newtonian suspending liquid. Additionally, when the local RBC concentration increases to a point where hydrodynamic interactions take place, we observe the formation of equally-spaced RBC structures, resembling the viscoelasticity-driven ordered particles observed previously in the literature, thus providing the first experimental evidence of viscoelasticity-driven cell ordering. The observed RBC ordering, unaffected by polymer molecular architecture, persists as long as the surrounding medium exhibits shear-thinning, viscoelastic properties. Complementary numerical simulations reveal that viscoelasticity-induced repulsion between RBCs leads to equidistant structures, with shearthinning modulating this effect. Our results open the way for the development of new biomedical technologies based on the use of viscoelastic liquids while also clarifying fundamental aspects related to multibody hydrodynamic interactions in viscoelastic microfluidic flows.

1 Introduction

The microvascular system is a vital network of small blood vessels that intricately weave throughout tissues and organs.¹ In it, red blood cells (RBCs) flow in small capillaries to exchange nutrients and gases. Microscale flow of RBCs determines microvascular perfusion in the microcirculatory system; it is also central to a broad range of biomedical applications for blood cell separation, single-cell analysis, and to assess storage lesion in transfusion medicine.⁴ In small vessels, RBCs usually flow in a single file and dynamically adapt their shape to the vessel flow conditions, such as flow rate, geometric confinement, and rheological properties of the surrounding plasma.^{5,6} Importantly, the RBC shape in capillary flow is susceptible to alterations in the intrinsic cell properties, which can serve as a phenotypical marker for many diseases, including malaria, diabetes, sickle cell disease, or neuroacanthocytosis syndromes,8-10 impairing blood flow and microvascular RBC perfusion. 11,12 While cell count and morphology have clear implications for human health, they can also serve as an opportunity to develop fast and accessible in vitro diagnostic platforms. 13,14

Progress in our understanding of in vitro RBC single-cell flow has advanced the development of lab-on-a-chip technologies for point-of-care instruments to assess RBC deformability and evaluate RBC flow properties on the microscale. 4,15-18 In this context, microfluidic experiments 19-21 and numerical simulations²²⁻²⁴ have been employed to investigate RBC flow on the single-cell level, revealing various RBC shape configurations in microvascular flow. 25-27 At rest, healthy RBCs have a biconcave, discocyte shape with a diameter of 8 µm and a thickness of 2 µm. RBCs suspended in a Newtonian buffer and subjected to microfluidic flows display more complex steady and dynamic shapes, such as parachutes, trilobes, tank-treading, tumbling, and rolling cells. In rectangular microchannels with dimensions similar to the RBC size, this shape complexity reduces to two dominant RBC shapes: the croissant and the slipper.^{20,21} The symmetric croissant shape predominantly

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appears at flow speeds $v < 5 \text{ mm s}^{-1}$ in microchannels with a width and height of approximately 10 µm and 8 µm, respectively. In contrast, the asymmetric slipper shape emerges at elevated cell speeds above $\nu > 3$ mm s⁻¹. These cell shapes are inherently coupled to their equilibrium position in the channel cross-section. For healthy RBCs, croissant-like RBCs flow at the channel centerline, whereas slippers flow closer to the channel side walls at an off-centered position. Recently, machine-learning approaches have been employed for fast and robust shape classification of healthy and diseased RBCs in confined microfluidic flows, 28,29 demonstrating the potential of RBC shapes and equilibrium position in the channel as a biomarker for specific pathologies and to assess the cell deformability in stored blood. 4,18,23

With a few notable exceptions, 30,31 the study of RBC in small capillary flows has mainly focused on using a buffer, a Newtonian liquid, as a suspending medium. Recently, non-Newtonian liquids have gained significant attention across several microfluidic applications due to the possibility of exploiting their nonlinear rheological properties (e.g., viscoelasticity, shearthinning viscosity) for cell focusing and separation.³² However, the behavior of RBCs in non-Newtonian fluids remains largely unexplored.

In this work, we explored the flow of RBCs under confined microfluidic flows using several viscoelastic polymer solutions with distinct rheological properties as suspending liquids. We experimentally explored the impact of the rheological properties of the carrier on RBC clustering and individual shape, observing marked differences compared to the Newtonian case. We also provided the first experimental evidence of the formation of equally-spaced RBC structures in viscoelastic flows, called RBC trains, resemblant to the viscoelastic ordering phenomenon recently reported for rigid particles.33-37 Finally, we employed numerical simulations to elucidate the interplay between cell shape and viscoelastic properties in developing RBC trains.

2 Methods

2.1 Experimental

2.1.1 Sample preparation. Blood was collected with informed consent from three healthy voluntary donors using a finger prick. Blood withdrawal, sample preparation, and experiments were performed according to the guidelines of the Declaration of Helsinki and approved by the ethics committee of the

"Ärztekammer des Saarlandes" (approval number 51/18). The blood was suspended in phosphate-buffered saline (PBS) solution (Gibco PBS, Fisher Scientific, Schwerte, Germany) and centrifuged at 1500g for five minutes to separate RBCs and plasma. Sedimented RBCs were resuspended in PBS, and the centrifugation and washing steps were repeated three times. For the control sample, a bulk hematocrit concentration of 3% was adjusted in a PBS solution that contained 1 g L^{-1} bovine serum albumin (BSA, Sigma-Aldrich, Taufkirchen, Germany).

To explore the impact of fluid rheology and polymer properties on the behavior of RBCs in microchannels, we prepared aqueous solutions of several macromolecules having different molecular weights and concentration values. We prepared solutions of hyaluronic acid salt (HA, Sigma-Aldrich, Taufkirchen, Germany) with a molecular weight in the range 1.5-1.8 MDa at mass concentrations of 0.1 wt% (HA01), 0.3 wt% (HA03), and 0.5 wt% (HA05) in a PBS/BSA mixture. Additionally, aqueous solutions of poly(ethylene oxide) (PEO, Sigma-Aldrich, Taufkirchen, Germany) at 1 wt% PEO with an average molecular weight of $M_{\rm w}$ = 4 imes 10^5 g mol^{-1} (PEO400k), at 1 wt% with $M_{\rm w} = 4 \times 10^6 \text{ g mol}^{-1}$ (PEO4M), and at 0.01 wt% with $M_{\rm w} = 8 \times 10^6 \, {\rm g \ mol^{-1}}$ (PEO8M) were prepared. The polymer powder was slowly added to the PBS/ BSA mixture, and the resulting solutions were mixed by gentle agitation on a roller-mixer for 24 hours to allow the complete dissolution of the polymer. We determined the osmolality m_{osm} of the used fluids (Table 1) using a freezing point osmometer (Osmomat 3000 basic, Gonotec, Berlin, Germany). Similar to the control sample in pure PBS/BSA, a bulk concentration of 3%Ht was adjusted in all investigated fluids.

Besides the control solution, we investigated three additional fluids with a near-constant viscosity that do not exhibit significant elastic properties in shear, namely an iodixanol solution, an isotonic polyvinylpyrrolidone (PVP), and a volume replacement fluid (VRF) (Table S1, ESI†). A description of the sample preparation for the constant-viscosity fluids and a summary of their rheological properties can be found in the ESI.† Results on RBC flow in the PEO solutions and constant viscosity fluids are presented in the ESI† if not mentioned otherwise.

2.1.2 Rheological characterization. We characterized the investigated fluids using shear and extensional rheometry. Steady and oscillatory shear measurements were performed to assess the viscosity η as well as the storage G' and loss G''moduli of the used samples. For this, a stress-controlled

Table 1 Overview of the fluid properties. The zero-shear viscosity η_0 , the infinite shear viscosity η_∞ , and the longest relaxation time λ , are derived from the fitting the rheological data with the cross-model (eqn (1)). The extensional relaxation time λ_e is derived from exponential fits of the temporal evolution of the stretched fluid filaments during capillary breakup (ESI)

Sample	w (wt%)	$m_{\rm osm}~({ m mOsm~kg^{-1}})$	η_0 (mPa s)	η_{∞} (mPa s)	λ (ms)	$\lambda_{\rm e}~({ m ms})$
Control	_	297.7 \pm 1.5	1.2 ± 0.2	_	_	_
HA01	0.1	300.7 ± 1.2	7.2 ± 0.1	3.8 ± 0.4	2.0 ± 0.5	0.5 ± 0.1
HA03	0.3	307.0 ± 4.6	89.2 ± 1.3	11.9 ± 0.9	13.2 ± 0.2	2.0 ± 0.1
HA05	0.5	309.3 ± 4.5	580.8 ± 21.9	23.0 ± 0.9	51.9 ± 0.3	3.3 ± 0.2
PEO8M	0.01	298.3 ± 0.7	1.3 ± 0.2	_	_	0.8 ± 0.4
PEO400k	1	303.0 ± 4.2	11.7 \pm 0.4	_	_	0.6 ± 0.3
PEO4M	1	301.5 ± 2.1	354.1 ± 6.5	21.9 ± 0.4	57.6 ± 0.2	31.0 ± 6.5

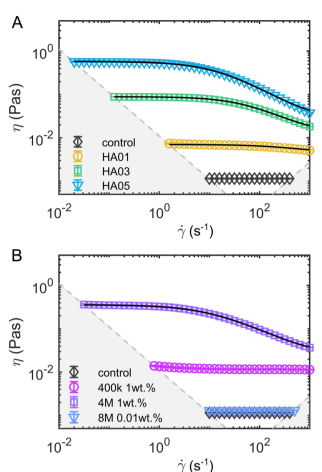


Fig. 1 Steady shear rheology of the polymeric fluids. Shear viscosity η as a function of the shear rate $\dot{\gamma}$ for (A) the HA solutions and (B) the PEO solutions. The solid black lines in the HA solution data represent fits of the viscosity curves with the cross model (eqn (1)). Dashed gray lines and areas indicate the lower torque limit of the device and the onset of flow instabilities under steady shear measurements in the cylinder geometry.³⁸

rheometer (MCR702, Anton Paar, Graz, Austria) equipped with a coaxial cylinder geometry (CC20, Anton Paar, Graz, Austria, inner radius $r_i = 10$ mm, radius ratio $r_a/r_i = 1.1$) was used at a constant temperature of T = 20 °C. While HA01 shows an almost constant shear viscosity η , HA03 and HA05 exhibit pronounced shear-thinning features (Fig. 1A). All HA solutions exhibit viscoelastic properties under oscillatory shear (Fig. S1A and B, ESI†). PEO400k and PEO4M exhibit viscoelastic features with near-constant and shear-thinning viscosity, respectively, similar to HA01 and HA05 (Fig. 1B and Fig. S2A, B, ESI†). The PEO8M solution has a viscosity similar to the control buffer. The three additional fluids (iodixanol, PVP, and VRF) show a constant shear viscosity (Fig. S3A, ESI†).

In oscillatory shear measurements (Fig. S1B, ESI†), we did not capture the terminal flow regime at low angular frequencies ω, which can be used to calculate the longest relaxation time based on the slopes of the storage G' and loss G'' moduli using standard small angle oscillatory shear (SAOS) analysis.³⁹ Therefore, we estimate the longest relaxation time λ by fitting the viscosity curves of the HA solutions (black lines in Fig. 1) with

the cross-model:

$$\eta = \eta_{\infty} + \frac{\eta_0 - \eta_{\infty}}{1 + (\lambda \dot{\gamma})^m},\tag{1}$$

where η_{∞} is the infinite shear viscosity, η_0 the zero-shear viscosity, λ a time constant that can be used as an estimate of the longest relaxation time, $\dot{\gamma}$ the shear rate, and m the factor that modulates the transition between the zero-shear viscosity region and the shear-thinning regime. For HA01, where the shear-thinning was scarcely noticeable, we further determined a relaxation time of $\lambda = 3.7 \pm 1.4$ ms using the μ -rheometer introduced previously. 40 This value agrees with the relaxation time of $\lambda = 2$ ms obtained from the cross-model (Table 1). For consistency, we use the relaxation time of $\lambda = 2$ ms for further analysis.

Moreover, we measured the shear rate dependent first normal stress difference N_1 (Fig. 2A) and shear stress τ (Fig. 2B) of the fluids using a cone-and-plate geometry (CP50/2, 50 mm diameter, 2° angle, Anton Paar, Graz, Austria). We employed a protocol with shear "peak hold" intervals, 41,42 which allowed us to correct for the normal-force transducer drift empirically. Additionally, the contribution of inertia to the normal force was subtracted.³⁹ Based on power-law fits of $N_1(\dot{\gamma})$ and the shear stress $\tau(\dot{\gamma})$, we calculated the Weissenberg number Wi = $N_1(\dot{\gamma})/\tau(\dot{\gamma})$ as a measure of the contribution of elastic effects (Fig. 2C). From these measurements, we concluded that elastic stresses dominate for the viscoelastic, shear-thinning polymer solutions (HA03, HA05, and PEO4M) at the shear rates observed in the microfluidic channel at our experimental conditions (Fig. 2C). To complete the rheological characterization of the fluids, we also performed extensional rheology experiments using the uniaxial elongation of the stretched fluid filaments (ESI†). In uniaxial extension, all HA and PEO solutions, as well as PVP, displayed elastic properties as evidenced by the formation of axially uniform cylindrical filaments during the last stage of capillary breakup (Fig. S4, ESI†).

2.1.3 Microfluidic setup. The RBC suspensions were pumped through a microfluidic chip that contains a set of four parallel channels. The rectangular channels have a width of $W = 10.1 \pm 0.4 \,\mu\text{m}$, a height of $H = 7.4 \pm 0.3 \,\mu\text{m}$, and a total length of L = 20 mm (Fig. 3A). These channel dimensions result in the formation of the well-characterized croissant and slipper RBC shapes.43 Using standard soft lithography,44 the microfluidic device was fabricated using polydimethylsiloxane (PDMS, RTV 615A/B, Momentive Performance Materials, Waterford, NY) and was bonded to a glass slide using a plasma cleaner (PDC-32G, Harrick Plasma, Ithaca, NY). The four channels shared a common inlet and outlet, which were connected with rigid medical-grade polyethylene tubing (0.86 mm inner diameter, Scientific Commodities, Lake Havasu City, AZ) to the sample and waste containers, respectively. The microfluidic device was mounted on an inverted microscope (Eclipse TE2000-S, Nikon, Melville, NY), which was equipped with blue LED illumination (SOLIS-415C, Thorlabs Inc., Newton, NJ), a USB 3.0 camera (DMK 23U1300, The Imaging Source, Bremen, Germany), and a 20× air objective (Plan Fluor, Nikon, Melville, NY) with a numerical aperture NA = 0.45.

Paper HA01 **HA03** HA05 PEO400k PEO4M Α 10² N₁ (Pa) 10⁰ 10² 10³ $\dot{\gamma}$ (s⁻¹) В 10² 10⁰ 10³ 10² $\dot{\gamma}$ (s⁻¹) C 10¹

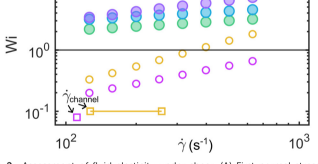


Fig. 2 Assessment of fluid elasticity under shear. (A) First normal stress difference N_1 as a function of the shear rate $\dot{\gamma}$ for the HA and PEO solutions. The horizontal dashed-dotted line in (A) indicates the experimental limit that can be resolved by the rheometer by measuring the N_1 response for the Newtonian solvent (which should be identically zero). The dashed line corresponds to the onset of secondary flows at high velocities in the used cone-and-plate geometry. 38 (B) Shear stress τ as a function of the shear rate $\dot{\gamma}$. Solid lines in (A) and (B) correspond to power-law fits of N_1 and τ , respectively. (C) Weissenberg number Wi = N_1/τ calculated from the power-law fits of N_1 and τ . For HA01 and PEO400k, we used the fit of N_1 in the high shear rate regime as an upper estimation of the maximum Weissenberg number for these fluids (open circles). The squares show the estimated shear rates $\dot{\gamma}_{\text{channel}} = v/D_{\text{h}}$ and shear rate ranges in the microfluidic channel, based on the RBC velocity v and the hydraulic channel diameter D_h .

Additionally, we use a $50\times$ air objective (NA = 0.55, LU Plan ELWD, Nikon, Melville, NY) for higher magnification of the RBC shape characteristics. A high-precision pressure device (OB1-MK3, Elveflow, Paris, France) was used to pump the suspensions through the channel at a pressure drop range of p = 50-2000mbar depending on the fluid sample. All microfluidic experiments were performed at 22 °C.

We recorded the RBC flow and obtained single-cell morphology data at two regions of interest (ROI) along the channel flow direction corresponding to $L/D_h = 120$ and $L/D_h = 1200$ from the inlet, with the hydraulic diameter of the rectangular channel $D_h = 2WH/(W + H) = 8.5 \mu m$. A low frame rate was chosen for each ROI to assess the distance s between different RBCs and RBC clusters. The frame rate was adjusted depending on the fluid viscosity so that in each frame of the image sequence, a specific cell was detected only once.

Based on the projection of the RBC flow in the x-y-plane of the microfluidic channel (Fig. 3A), we extracted the center of mass, the cell length L_{RBC} in the flow direction, the cell area $A_{\rm RBC}$, and the cell diameter $D_{\rm RBC}$ across the channel from binarized images of all single RBCs and RBC clusters using a custom MATLAB (9.14.0.2206163 (R2023a), MathWorks, Natick, MA) algorithm. Due to the absorbance of blue LED light used in this study (see Fig. 3A), and the resulting excellent contrast between the RBC and the surrounding medium, no additional image processing is required. We calculated the cell deformability index DI_{RBC} defined as $DI_{RBC} = (a - b)/(a + b)$, where a and b are the major and minor axes of the RBC shape during flow. Furthermore, we calculated the probability density distributions (PDFs) by kernel density estimation based on the histograms of each quantity (Fig. 3B-E and Fig. S1C, ESI†). The first peaks in the length and area distributions correspond to the most probable single-cell length $L_{RBC,1}$ and area $A_{RBC,1}$. These single-cell characteristics were used to assess the effect of the polymer solutions on the RBC shape. Further, the distributions of L_{RBC} and the distance s between two consecutive objects (Fig. 3A) were used to discriminate single RBCs from RBC clusters. Here, we defined a single RBC as a cell that exhibits a length smaller than the minimum in the PDF between the first and second peak (Fig. 3B) and which is further than a distance of $L_{RBC,1}$ away from any other RBC or cluster.

Based on a single-cell diameter of $D_{\text{RBC},1} \approx 8 \, \mu\text{m}$, the confinement ratio $\beta = D_{RBC,1}/D_h$ is approximately 0.94, which is much larger than the values of $0.2 < \beta < 0.45$, previously employed to investigate viscoelastic train formation of rigid particles. 36,45,46 Note that reducing the confinement ratio by increasing the channel cross-section results in the emergence of unstable shape types that display rich shape dynamics.⁴³

To calculate the RBC velocity, we performed a second set of experiments with a frame rate of up to 400 frames per second, depending on the applied pressure drop and fluid sample. Individual cell velocities were determined in this second recording by tracking the cell position over the image sequence within the field of view. Since we did not observe interindividual variations in the results, data were averaged between the three donors. For individual RBC shape and ordering

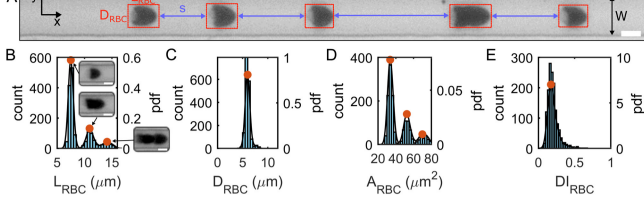


Fig. 3 Microfluidic flow of RBCs in HA solutions. (A) A representative snapshot of the RBC flowing in a HA05 solution at v = 1 mm s⁻¹. Red boxed highlight RBCs with length L_{RBC} in flow direction and diameter D_{RBC} across the channel width. Blue arrows indicate distances s between consecutive RBCs and clusters. (B)–(E) Representative histograms and probability density distributions (PDFs) by kernel density estimation of (B) the RBC length L_{RBC} . (C) the RBC diameter D_{RBC} , (D) the projection area A_{RBC} , and (E) the deformability index DI_{RBC} . Data are shown for the HA05 solution at an RBC velocity of $v = 1 \text{ mm s}^{-1}$. Red circles indicate PDF peaks corresponding to single RBCs, doublets, and triplets (see inset images) (scale bars, 5 μ m).

measurement, data analysis was performed on an average of 7725 cells (between 4629 and 20337 cells).

2.1.4 Numerical methods. To gain further insights into the main mechanism governing RBC interactions in polymeric fluid flows, we perform numerical simulations using experimentally available data. We note that the RBC shapes (for each case investigated here) are extracted from experimental data, and thus fixed. In order to reduce computational cost, the channel geometry is modeled as an axisymmetric pipe flow. Simulations are performed using two RBCs traveling at approximately terminal velocity. The fluid domain is constructed as a 5° slice, one cell wide, with a radius $R = 3.7 \mu m$ and a total length 100 µm. The domain is composed of the boundaries inlet, outlet, wall, RBC1, and RBC2. The inlet condition is set to a Poiseuille flow $\bar{U}(y,t) = U_{\text{max}}(t)(R^2 - y) + U_{\text{wall}}(t)$ where the prefactor is varied with time as

$$U_{\text{max}}(t) = \begin{cases} \frac{1 - \cos \pi t / t_{\text{ramp}}}{2} U_{\text{max}} & t < t_{\text{ramp}} \\ U_{\text{max}} & t \ge t_{\text{ramp}} \end{cases}$$
 (2)

and the outlet condition to $\nabla \bar{U} = \bar{0}$. The wall velocity condition is similarly varied with time as

$$U_{\text{wall}}(t) = \begin{cases} \frac{1 - \cos \pi t / t_{\text{ramp}}}{2} U_{\text{wall}} & t < t_{\text{ramp}} \\ U_{\text{wall}} & t \ge t_{\text{ramp}} \end{cases}$$
(3)

with no-slip boundary conditions on RBC1 and RBC2 surfaces. U_{wall} is set to $-0.805U_{\text{max}}$ in all simulations.

Note that the velocity boundary condition is chosen by performing simulations of a single RBC placed at the center of the channel; a velocity sweep is performed for each set of fluid parameters to determine the single RBC terminal velocity. For all fluid types explored here, a single RBC terminal velocity of approximately $0.805U_{\text{max}}$ is found. We ensured that the repulsive pair forces found in the viscoelastic models are insensitive to small changes in the RBC velocity near terminal velocity.

The pressure outlet boundary condition is p = 0 and at all other boundaries $\nabla p = \bar{0}$. The conformation tensor Θ boundary condition is set to $\nabla \Theta = 0$ on all boundaries. The boundary of each RBC is formed by tracing an RBC boundary taken from an image of the HA05 experiment.

For each RBC separation, the 2D boundary of the simulation is formed in CAD software and ingested into GMSH to extrude and create an unstructured mesh. The mesh is composed primarily of (in 2D) quadrilateral cells. Care is taken at the boundaries of the simulation domain to keep surfaces orthogonal. Each mesh is heavily refined near each RBC surface, with a refinement that decreases gradually with the distance to the RBCs.

Two main constitutive models are used to describe the rheological properties of the polymeric fluids, namely the Giesekus and FENE-P models. While both constitutive models are known to describe polymeric fluids possessing viscoelastic behavior and shear-thinning properties, the Giesekus accounts for microstructural anisotropy and is more appropriate for nondilute solutions. 47 Conveniently, these constitutive models are readily available within OpenFOAM, an open-source computational fluid dynamics solver, and RheoTool (for more info, please see ref. 48 and 49). For both models, the total stress σ can be written as the sum of solvent and polymeric contributions, σ_s and τ .

$$\sigma = \sigma_{\rm S} + \tau \tag{4}$$

The solvent stress tensor is formed from the solvent viscosity η_s and deformation tensor $D = \nabla u + \nabla u^T$.

$$\sigma_{\rm S} = \eta_{\rm S}(\nabla u + \nabla u^{\rm T}) \tag{5}$$

The constitutive equations governing the polymeric extrastress in the Giesekus and FENE-P models can be written

respectively as

$$\tau + \lambda \overset{\nabla}{\tau} + \alpha \frac{\lambda}{\eta_{p}} (\tau \cdot \tau) = \eta_{p} (\nabla u + \nabla u^{T})$$
 (6)

$$\tau + \frac{\lambda}{f} \overset{\nabla}{t} = \frac{a\eta_{\rm p}}{f} (\nabla u + \nabla u^T) - \frac{D}{Dt} \left(\frac{1}{f}\right) [\lambda t + a\eta_{\rm p} I] \tag{7}$$

where τ is the polymeric extra-stress tensor, ∇ is the upperconvective derivative, λ is the fluid relaxation timescale, and $\eta_{\rm p}$ is the polymer viscosity. In the Giesekus model, the mobility parameter α measures the amplitude of (microstructural) anisotropy and is related to shear-thinning properties; $\alpha = 1$ denotes maximum anisotropy and $\alpha = 0$ denoting isotropic relaxation. 50 In the FENE-P model the parameters a and f are defined using the extensibility parameter L^2 and τ as

$$f = \frac{L^2 + \frac{\lambda}{a\eta_p} \text{tr}(\tau)}{L^2 - 3}, \quad a = \frac{L^2}{L^2 - 3}$$
 (8)

All simulations are performed in the log-conformation approach, 49 where the polymeric extra-stress τ is recast (in the Giesekus model) as

$$\Theta = \log \left[\frac{\lambda \tau}{\eta_{\rm p}} + I \right] \tag{9}$$

It is common to define the dimensionless parameter $\beta' =$ $\frac{\eta_s}{\eta_p+\eta_s},$ which is a proxy for the inverse polymer concentration.

For all simulations, we chose the common model parameters $\beta' = 0.2$ and $\lambda = 0.1$. Within the Giesekus model, we set the microstructural parameter $\alpha = 0.4$ and in the FENE-P model, the extensibility parameter is set to $L^2 = 10$.

We explored Wi \in [0.1, 1.0, 5.0] for each model, where the Weissenberg number is estimated as Wi = $\lambda U_{\text{max}}/R$. The simulations are solved using a partially coupled direct solver (p and U coupled, Θ segregated) and a Crank-Nicholson time integration scheme. A time step of $dt = 1 \times 10^{-4}$ is applied in all simulations for a total time of 30λ . The parameter τ , used to vary the velocity inlet condition slowly, is set to 2.5λ .

3 Results

RBCs are highly deformable, meaning that their shape under confinement is expected to depend upon a combination of both rheological properties of the suspending liquid and experimental conditions, such as volumetric flow rate and channel size. Hence, we start by examining the formation of RBC clusters and RBC shapes in highly confined microfluidic flow (Fig. 3A). Specifically, we explore the resulting RBC clusters and shapes in different viscoelastic liquids, namely, HA and PEO, when compared to the control.

To assess the effect of fluid elasticity, we define the dimensionless Deborah number, De = $\lambda v/D_h$, where λ is the fluid relaxation time, ν is the RBC speed, and $D_{\rm h}$ is the hydraulic diameter of the rectangular microfluidic channel. Note that our

definition of De employs the RBC speed ν rather than the fluid speed: this is justified by the large confinement experienced by RBC in the microchannel (see, for instance, Fig. 3A), with the RBC occupying almost the entire channel. Depending on the Deborah number and the distance between them, two flowing objects can either experience attraction or repulsion dynamics. 35

RBC cluster formation

RBCs are well-known for forming clusters in microfluidic flows, i.e., two or more closely adhering cells. 51-54 Cell clustering is often undesirable in diagnostic microfluidic applications since it makes single-cell analysis difficult. Here, we observed a significant distinction in cluster formation depending on the fluid rheological properties (Fig. 4A). Our results show that in the Newtonian control (i.e., buffer), RBCs can be distinguished within a cluster. In contrast, RBCs in the polymeric HA solutions are closely attached. For the control fluid and HA01 an average of roughly 63% of all RBCs flow as single cells (Fig. 4B). This percentage increases to approximately 80% for HA03 and HA05. While we observe a large number of RBC clusters containing N > 3 cells in the control and HA01 samples, this fraction is significantly reduced in the higher concentrated HA03 and HA05 solutions (Fig. 4C). Moreover, we observe an increase in the number of clusters in the control buffer as the cells travel through the channel from ROI1 ($L/D_h = 120$) to ROI2 $(L/D_h = 1200)$, which is not found in the HA solutions.

PEO8M shows similar clustering behavior to the control buffer (Fig. S5, ESI†). In contrast, RBCs in PEO400k and PEO4M exhibit more closely packed clusters and an increased probability of single-cell flow. RBC suspended in the constant viscosity PVP and VRF solutions show clusters that are similar to those in PEO and HA. However, we do not observe a pronounced increase in the percentage of single cells in these fluids (Fig. S6, ESI†).

Generally, the initial aggregation process is attributed to slower RBCs acting to create a sequence of trailing cells.⁵² Once the cells are in close proximity, RBCs aggregate into clusters. This reversible aggregation was previously explained based on the bridging of macromolecules between the RBCs or induced by depletion.55,56 We find that adding polymers results in macromolecule-induced adhesion of individual cells. In combination with the bullet-like shape, these RBCs form closely packed clusters. Due to their high deformability, RBCs that adhere to a cluster deform and create large interaction surfaces between the cells. This results in high interaction energies, larger than for hard sphere suspensions. Once RBCs form a cluster of adhering cells in the polymeric fluids, either in the tubing, inlet reservoir, or channel, the RBCs stick together and are not separated by the flow.

3.2 RBC shape characteristics

Next, we characterize single-cell RBC morphology as a function of fluid type and flow conditions (Fig. 5A). To this end, we determine the single-cell length $L_{RBC,1}$, diameter $D_{RBC,1}$, area A_{RBC,1}, and deformability index DI_{RBC,1} during capillary flow in the different solutions (Fig. 5B-E). For the control, HA03, and

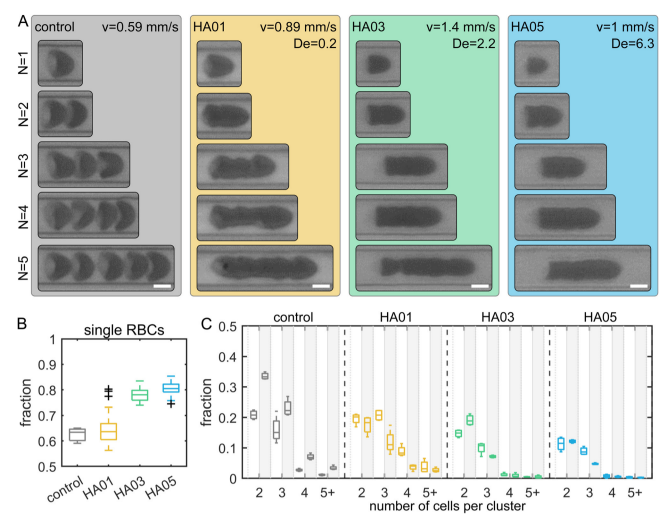


Fig. 4 RBC cluster formation in polymeric liquids. (A) Representative snapshots of clusters containing N = 1-5 RBCs (top to bottom) for the different HA samples. (Scale bars, 5 μ m.) For the control, the individual RBCs can be discriminated within the clusters. Individual RBCs adhere closely to one another for the HA solutions and cannot be discriminated within the clusters. (B) Box plots of the fraction of single RBCs that are spaced further than one RBC length apart and do not belong to an adhering cluster. (C) Box plots of the fraction of RBC clusters. The x-axes label 5+ denotes clusters with N = 5 or more RBCs. The x-axis shows the number of RBCs per cluster. Boxplots in front of the white and gray areas correspond to data in ROI1 ($L/D_h = 120$) and ROI2 ($L/D_h = 1200$), respectively. The bottom and top of each box are the 25th and 75th percentiles of the sample, respectively. The line in the middle of each box is the sample median. Whiskers go from the end of the interquartile range to the furthest observation. Data beyond the whisker length are marked as outliers with '+' signs.

HA05 cases, the average single-cell length in flow is 7.5 μm, independent of the cell velocity (Fig. 5B). In contrast, $L_{\text{RBC}.1}$ for HA01 exhibits higher values between 8 µm and 9 µm, and displays an increase with increasing flow velocity. This is due to a pronounced fraction of slipper-shaped and other elongated, non-croissant-like RBCs in the HA01 solution (Fig. S1D, ESI†). Slipper-like cells flow at an off-centered position in the microfluidic channel (Fig. S1E, ESI†) and exhibit an elongated shape, which results in a larger calculated single-cell length and deformability index (Fig. S1F, ESI†) based on their projection in the x-y-plane. Disregarding off-centered slipper-shaped RBCs leads to a smaller single-cell length (inset for HA01 in Fig. 5B). However, symmetry breaking in the rectangular crosssection results in slipper-like cells that are rotated by 90° in the channel cross-section. These cells flow at a centered position with respect to the x-y-plane, similar to croissants, and were,

therefore, referred to as sheared croissants in previous studies.²⁸ Similar to slipper-shaped cells in the x-y-plane of the channel, their pronounced tail (see arrow in the bottom panel of Fig. 5A for HA01) contributes to an increase in $L_{\rm RBC.1}$.

The single-cell diameter $D_{\rm RBC,1}$ decreases with increasing cell velocity for all samples and is smaller in the HA solutions than in the control buffer (Fig. 5C). Compared to the parabolic velocity profile observed in Newtonian liquids, the velocity profile in shear-thinning fluids is flatter around the centerline, resulting in a squeezing of the cells and forming bullet-like RBC shapes. The projection area $A_{\rm RBC,1}$ decreases with increasing HA concentration (Fig. 5D). In the control solution, $A_{\rm RBC,1}$ is approximately 42 μ m² below cell velocities of 1 mm s⁻¹, in good agreement with previous studies.²⁹ The single-cell deformability index $DI_{\rm RBC,1}$ increases as the cells elongate with increasing velocity (Fig. 5E). Note that the occurrence of slipper-shaped

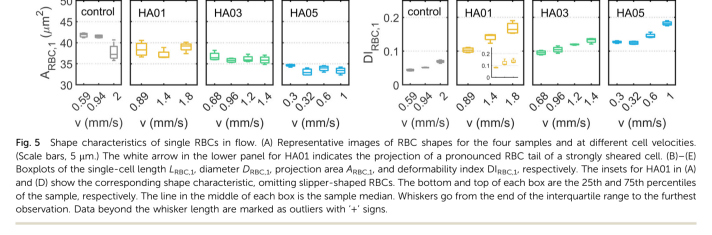
v (mm/s)

De

19 2 3. 63

HA05

Paper HA01 **HA03 HA05** Α control 9.6 mm/s s/mm 6 velocity 0.9 mm/s 4 2 mm/s ω C В De De De De De De 1,5 1,9 1,3 19 2 3. 63 √ √ δ √ δ ⁰ ½ 0,3 Ob 05 03 19 2 3. 63 control HA05 control HA03 HA01 HA03 HA01 HA05 L_{RBC,1} (μ m) D_{RBC,1} (µm) 8 6 0,80 0,89 06,00 V3 VX 03,3000 03,300 059°94 06,00 V3 VV



Ε

0.2

v (mm/s)

control

v (mm/s)

De

0,3

HA01

v (mm/s)

De

√ √ 5 √ 5 √ 5

HA03

v (mm/s)

De

19 2 31 63

HA05

v (mm/s)

De

/ 1/2 /3 1/3.

HA03

RBCs for HA01 leads again to an apparent increase in DI_{RBC.1}, similar to the observations regarding $L_{RBC.1}$. For HA03 and H05, we observe a strong correlation of the deformability index with increasing De.

RBCs suspended in PEO follow a qualitatively similar behavior regarding their shape as in HA (Fig. S2C-G, ESI†). In the

near-constant viscosity fluids, the RBC shape closely resembles the croissant-like shape found in the control buffer solution (Fig. S3B-F, ESI†). Generally, the osmolality of the suspending fluid must be considered when assessing the RBC shape and its characteristics.⁵⁷ With increasing concentrations of HA and PEO, the osmolality rises up to 309.3 mOsm kg⁻¹ (Table 1).

v (mm/s)

control

D

v (mm/s)

De

03 03 0k

HA01

The human plasma osmolality is primarily determined by the plasma sodium concentration with an RBC isotonic osmolality in a range of approximately 280–310 mOsm kg^{-1} .⁵⁸ Healthy RBCs can preserve their shape for osmolality values between 210–380 mOsm kg^{-1} ,⁵⁹ within the osmolality range of the used polymer solutions (Table 1 and Table S1, ESI†). Our results are independent of the medium's osmolality as we do not observe a strong correlation between the RBC single-cell area in capillary flow and the osmolality of the surrounding medium for the fluids used in this study (Fig. S7, ESI†). Only the iodixanol solution has an elevated osmolality (Table S1, ESI†), and suspended RBCs display a smaller shape. Hence, we do not find a pronounced effect of the medium osmolality on the RBC shape during capillary flow by

swelling or shrinking in the HA and PEO solutions. Overall, these results demonstrate that the RBC shape is highly sensitive to the rheological properties of the surrounding medium beyond its osmolality. RBCs exhibit a squeezed, bullet-like shape in the polymeric solutions compared to the Newtonian control. Surprisingly, we find that fluid elastic stresses can lead to a different shape of RBC, even in the near-constant viscosity case (HA01 and PEO400k). For these solutions, we observed a pronounced fraction of slipper-shaped RBCs below 2 mm s⁻¹, which usually only emerges at higher velocities in Newtonian buffer solutions. Even at low velocities, the emergence of slipper cells has been reported before using Newtonian fluids, such as dextran solutions, with a higher viscosity than the control buffer.21,60 This result indicates a dependence of the stable RBC shape on the external shear stress experienced by the cell.

3.3 RBC ordering and train formation

We now proceed to explore the behavior of RBC flowing in a microchannel under the large confinement of $\beta = D_{RBC,1}/D_h =$ 0.94 (Fig. 6). By looking at the experimental snapshots in Fig. 6, we notice that RBCs align along the channel centreline irrespective of the liquids in which they are suspended. However, the distance between consecutive cells is strongly affected by the properties of the suspending liquids. For instance, in both the Newtonian control buffer (Fig. 6A) and the HA01 with negligible shear-thinning properties, RBCs tend to be close to each other. For HA03 and HA05, both presenting shearthinning properties, RBCs arrange to form equally-spaced structures reminiscent of the viscoelastic particle trains recently observed in the literature. 35,36,46 To fully characterize the onset of RBC-ordered train structures, we employed the normalized distance $S^* = s/L_{RBC,1}$ with s as the distance between consecutive single RBCs or clusters (Fig. 3A). Hence, $S^* = 0$ if two objects are in close contact yet can still be distinguished as two single objects. It is well established that the bulk concentration is not an appropriate parameter to study ordering because of the fluctuations of objects entering the device. 35,61 For this reason, we employ the local RBC concentration ϕ_1 = $N_{\rm RBC}L_{\rm RBC,1}/L_{\rm f}$, in agreement with previous studies, ^{35,36} where $N_{\rm RBC}$ is the number of RBCs in the observation window and $L_{\rm f}$ is the length of the observation window. Consequently, frames of the image sequence that contain the same ϕ_1 are identified, and

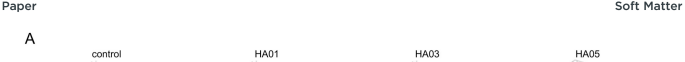
the distributions of the normalized distance S^* between two cells in these frames are evaluated. Moreover, data is analyzed in two regions of interest at $L/D_{\rm h}=120$ and $L/D_{\rm h}=1200$ downstream of the channel entrance.

The quantitative analysis of RBC distance distributions confirms the experimental observations. Notably, for HA03 and HA05, we observed the occurrence of a clear peak between $S_{\rm peak}^* \approx 1$ and $S_{\rm peak}^* \approx 2$, meaning that there now is a characteristic distance between consecutive RBCs, thus establishing the first experimental evidence of viscoelastic ordering on cells. The prominence of this peak increases with increasing local concentration, demonstrating the existence of a preferential inter-RBC distance $S^* > 0$ and the formation of RBC trains in the strongly shear-thinning, viscoelastic HA solutions (see bottom insets in Fig. 6A). This peak value is independent of the cell velocity and De (Fig. 6B), in agreement with previous studies on rigid particles.35,46 We observe an increase in the preferential inter-RBC distance further downstream at L/D_h = 1200 for HA05, where the peak in the S* distribution shifts to $S_{\text{neak}}^* = 2-5$ (Fig. 6C and Fig. S8A, ESI†). Our results also echoed previous experimental46 and numerical33 studies on the viscoelastic ordering of rigid particles, where ordered structures were observed in viscoelastic shear-thinning liquids but not in viscoelastic liquids with constant viscosity. Jeyasountharan et al.36 reported that an increase of the confinement ratio from $\beta = 0.2$ to $\beta = 0.45$ promoted the formation of equally-spaced structures over shorter channel lengths, with particles ordered at $L/D_h > 2000$ for $\beta = 0.2$ and particles ordered at $L/D_h > 1000$ for β = 0.45. In our case, with β = 0.94, we observed the formation of the RBC trains already at $L/D_{\rm h}$ = 120. According to Jeyasountharan et al.,36 the ordering for rigid particles is expected to scale with β^3 when defining S* with respect to the particle characteristic length rather than the channel diameter. Since our β is approximately two times $\beta = 0.45$, the ordering should be observed at an S* roughly eight times smaller, meaning 1000/8 = 125, in excellent agreement with our observations.

Our results are general and not only restricted to HA solutions. RBCs in the PEO8M solution show similar behavior as in the control buffer with a pronounced peak in the distance distribution at $S^* = 0$ independent of the local concentration (Fig. 6D). Similarly, the viscoelastic, near-constant viscosity PEO400k solution does not induce RBC ordering at the investigated cell velocities. As opposed to this, we observe a pronounced peak at $S^*_{\rm peak} = 2$ in the inter-RBC distribution for the viscoelastic, strongly shear-thinning PEO4M solution (Fig. 6E). RBCs in the other constant viscosity fluids do not self-assemble into equally-spaced cell trains (Fig. S8B, ESI†).

3.4 Forces between RBCs

We now use numerical simulations to gain insights into the main forces responsible for the ordering of RBCs in viscoelastic fluids. Specifically, we want to clarify whether the RBC ordering is caused by the fluid elasticity alone or whether it is a combination of suspending liquid viscoelasticity and RBC



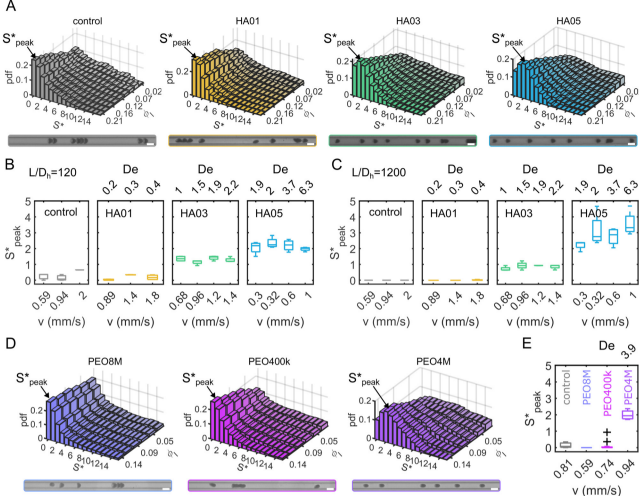


Fig. 6 Ordering of RBCs in viscoelastic confined flows. (A) Probability density distributions (PDF) of the normalized distance $S^* = s/L_{RBC,1}$ as a function of the local RBC concentration ϕ_1 . Data is shown for control ($v = 0.6 \text{ mm s}^{-1}$, De = 0), HA01 ($v = 1.8 \text{ mm s}^{-1}$, De = 0.4), HA03 ($v = 1.2 \text{ mm s}^{-1}$, De = 1.9), and HA05 ($v=1~mm~s^{-1}$, De=6.3) at $L/D_h=120$. (B) and (C) Boxplots of the peak distance S_{peak}^* at $L/D_h=120$ and $L/D_h=120$, respectively. (D) PDF of S^* for the different PEO samples as a function ϕ_1 . Data is shown for PEO8M (v = 0.74 mm s⁻¹), PEO400k (v = 0.81 mm s⁻¹), and PEO4M (v = 0.59 mm s⁻¹, De = 3.9) at $L/D_h = 120$. (D) Boxplots of the peak distance for the PEO samples at $L/D_h = 120$. The arrows in (A) and (D) indicate the peak value S_{peak}^* of the corresponding PDF at the highest local concentration. Inset images below the graphs in (A) and (D) show representative experimental snapshots with N = 8 individual objects (single RBCs or clusters) in the field of view ($\phi_1 \approx 0.21$) for the HA solutions and N=6 individual objects ($\phi_1 \approx 0.14$) for the PEO solutions. (Scale bars, 10 μm.) Data in (B), (C), and (D) is shown at a bulk RBC concentration of 3%Ht. The bottom and top of each box in (B), (C), and (D) are the 25th and 75th percentiles of the sample, respectively. The line in the middle of each box is the sample median. Whiskers go from the end of the interquartile range to the furthest observation. Data beyond the whisker length are marked as outliers with '+' signs

shapes. We performed numerical simulations using two viscoelastic constitutive models, namely the Giesekus and the FENE-P models (see Section 2.1.4). Parameters for the fluid rheological models used in the numerical simulations are obtained by fitting them to experimental data (Table 1). In addition, the RBC shape used in the simulations is obtained from the experiments (Fig. 5). In the simulations, we extract the flow field, the forces between the RBCs, and the fluids stresses around the RBCs for different configurations of surface-surface RBC distances S* and Weissenberg numbers Wi.

The streamlines of the velocity field between the RBCs (Fig. 7A) exhibit vortical structures, similar to experimental flow field measurements in the vicinity of two RBCs in capillaries.⁶² At this distance and Wi, changes in the x normal stress component of the polymeric extra-stress relative to that in an empty pipe flow $(\tau_{xx}$ – $\tau_{xx,\infty}$) develop (Fig. 7B). We observe a high-stress region that appears on the right of the trailing RBC, which gives rise to a net force that pushes the RBCs away. As the RBCs are brought together, the elastic wakes formed in front and behind each RBC interact to yield a net repulsion, similar to numerical simulations on rigid spheres.³³ Moreover, high-stress regions on both sides near the tail of the RBCs emerge. This leads to a repulsion between RBCs and the wall, which could explain the emergence of squeezed bullet-like cells observed in the experiments. Representative velocity and stress fields around a single RBC and cells spaced at various surface-to-surface distances are shown in Fig. S9 (ESI†).

Simulations allow us to calculate the force curves and effective pair potential between RBCs as a function of the

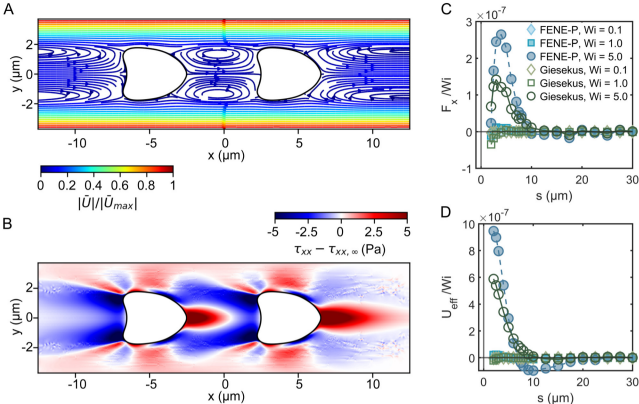


Fig. 7 Numerical simulations of two croissant-shaped RBCs in different polymer solutions. (A) Streamlines of the velocity field of the system around RBCs at a surface-surface separation of $5 \mu m$ and Wi = 5.0 using the Giesekus model. (B) Visualization of the differences in the x normal stress component for the same system as in (A). (C) The pair-force between RBCs, rescaled with the Wi of the flow, against the surface-surface displacement between the two RBCs. Positive force indicates a net repulsion between RBCs, while negative forces indicate a net attraction. As the Wi is increased to Wi > 1, RBCs develop a substantial repulsion at approximately one RBC diameter. Shear-thinning, present in the Giesekus model, modifies the turn-on of the viscoelastic repulsion. (D) Effective pair-potential between RBCs, calculated by numerically integrating the force curves in (C).

RBC distances S* for the explored models and Weissenberg numbers (Fig. 7C and D). Positive force indicates an observed repulsion between the RBCs, while negative force indicates attraction. For Wi « 1, both models show a short-ranged attraction that starts at approximately 5 µm. As Wi is increased, we find in both models that the attraction at short distances is overwhelmed by new repulsive forces. The turn-on of the repulsion is modified and weakened in the Giesekus model at intermediate and high Wi. Thus, we believe that the mechanism of the pair repulsion comes mainly from the elasticity of the fluid, and it seems that for Wi >> 1, the repulsion can be obtained despite the fluid shear-thinning properties. Nevertheless, shearthinning viscosity appears to modulate the repulsion by reducing the velocity gradients around the center line of the fluid, leading to lessened elastic stresses.

In summary, our numerical simulations demonstrate that the fluid elasticity produces significant repulsion between the cells above Wi » 1. Similarly, the Weissenberg number for the solutions where we observe RBC ordering (HA03, HA05, and PEO4M) is always larger than one at the probed shear rate in the microfluidic channel experiments (Fig. 2C). In contrast, Wi « 1 for HA01 and PEO400k, based on an upper estimation of the maximum Weissenberg number for these weakly

viscoelastic fluids (Fig. 2C). Potentially, experimental RBC ordering could also appear in these solutions at much larger shear rates when Wi >> 1, as predicted by our numerical simulations. However, this would require a higher pressure drop for the microfluidic chip and fluids used, resulting in channel de-bonding in our experimental apparatus. Using a different type of bonding method based on adhesive tape can provide higher bonding strength. Still, it can also lead to surface modification, thus altering the overall interpretations of the results. For this reason, we believe such an analysis of ordering in viscoelastic near-constant-viscosity liquids at Wi >> 1 falls behind the scope of this work.

4 Discussion and conclusions

Our results show that the interplay of fluid stresses, cell shape, and volume fraction can lead to the ordering of RBCs in microflows. We provide the first experimental evidence that fluid elasticity can lead to highly deformable, anisotropic RBCs that self-assemble into trains with a preferential inter-cell distance. Using two different polymers in combination with different polymer concentrations and molecular weights, we

experimentally show that viscoelastic, shear-thinning features of the suspending medium are required for RBC ordering independent of the molecular architecture of the monomer. This result transcends the nature of the individual polymers (e.g., HA or PEO), and can be generalized to the specific bulk rheological properties. For rigid particles, migration towards the channel centerline and ordering is commonly observed in viscoelastic fluids for confinement ratios $\beta > 0.15$. In this study, we impose velocities at which mainly croissant-shaped RBCs emerge.²⁰ Hence, RBCs preferentially flow at a stable equilibrium position at the channel centerline v/W = 0(Fig. S1C, ESI†). We show that this cell alignment along the channel flow direction is reached relatively quickly in the first region of interest $(L/D_h = 120)$ in accordance with previous studies.⁵³ In combination with the high confinement ratio of $\beta \approx 0.94$ in the present work, cell train formation of aligned RBCs is achieved already at L/D_h = 120 after the fluid inlet, which presents a sweet spot for both cell ordering and RBC shape assessment.²⁰ The observed inter-RBC distance, represented by the peak in the distribution of the normalized cell distance S_{peak}^* , is independent of De within the investigated Deborah number regime, as long as a Deborah number above the critical one describes the flow. For RBCs, a clear peak in the distribution of inter-particle distances can be observed at local concentrations larger than $\phi_1 \geq 0.1$, similar to the required local concentration for rigid beads.³⁵

Self-assembled croissant-shaped RBCs exhibit a more squeezed shape in flow with respect to the channel width and resemble the shape of a bullet in the HA and PEO solutions compared to cells in the control buffer solution. Simultaneously, the single-cell deformability index DI of these centered shapes shows a pronounced increase with the Deborah number. A similar cell stretching and the occurrence of bullet-like RBC shapes have been observed in micro-confined RBC flow using Newtonian buffer solutions. 64,65 However, these shape configurations were induced by compressing the cells in channels with a width smaller than 10 µm, hence smaller than the RBC size. In contrast, the shape changes observed in this study emerge as a consequence of the blunted velocity profile in the channel cross-section using shear-thinning fluids. While we focus on characterizing the effect of fluid viscoelastic on centered croissant-like shapes in the present study, future work will be required to investigate these effects on the off-centered slipper and other dynamic RBC shapes. This comprehensive framework will then encompass all shapes commonly observed in the so-called RBC shape phase diagram²⁰ and the potential of cell-ordering of stable and dynamic shapes.

Assessing the RBC shape and its characteristics, such as the cell's deformability index DI, during microfluidic flow has received increasing attention as a tool to evaluate pathological cell changes induced by diseases such as malaria, hemoglobinopathies, and COVID-19,8-10,29 and to assess stored blood quality. 18,66 Based on the results presented in this study, future work is required to verify if cell ordering of such impaired RBCs is possible and to evaluate how the addition of polymers and deviations from the parabolic velocity profile affect the shape of diseased RBCs. Addressing these points and exploring polymer solutions, such as the HA and PEO solutions used in this study, in microfluidic single-cell analysis techniques will further expand our understanding of self-assembling systems beyond rigid particles and enhance various biomedical applications.

Viscoelastic ordering has been found to improve the encapsulation efficiency of flowing particles well above the Poisson statistics limit, but this has only been demonstrated for rigid particles. 67,68 Our findings on RBC confirm that it is also possible to order living cells. This means that single-cell analysis of encapsulated cells using viscoelastic liquids is now feasible.

Author contributions

S. M. R. and F. D. G. designed research; S. M. R. and Y. R. performed experiments; I. G. and P. E. A. performed numerical simulations; S. M. R., Y. R., I. G., and F. D. G. analyzed data; S. M. R., Y. R., I. G., P. E. A., F. D. G., and C. W. wrote the paper; S. M. R., C. W., and P. E. A. supervised research.

Conflicts of interest

There are no conflicts of interest to declare.

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