Microfluidic study of enhanced deposition of sickle cells at acute corners and its possible role in vaso-occlusion

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Abstract

Sickle cell anemia is a blood disorder, known to affect the microcirculation and is characterized by painful vaso-occlusive crises in deep tissues. During the last three decades, many scenarios based on the enhanced adhesive properties of the membrane of sickle red blood cells have been proposed, all related to a final decrease in vessels lumen by cells accumulation on the vascular walls. Up to now, none of these scenarios considered the possible role played by the geometry of the flow on deposition. The question of the exact locations of occlusive events at the microcirculatory scale remains open. Here, using microfluidic devices where both geometry and oxygen levels can be controlled, we show that the flow of a suspension of sickle red blood cells around an acute corner of a triangular pillar or of a bifurcation, leads to the enhanced deposition and aggregation of cells. Thanks to our devices, we follow the growth of these aggregates in time and show that their length does not depend on oxygenation levels, but that their morphology changes dramatically to filamentous structures when using autologous plasma as a suspending fluid. We finally discuss the possible role played by such aggregates in vaso-occlusive events.

Key words: Aggregates; Corner flow; Adhesion; Microfluidics; Vaso-Occlusion
Introduction

Sickle cell anemia (SCA) is an inherited hemoglobinopathy (1) affecting millions of people worldwide (2). It is caused by a point mutation resulting in the production of abnormal hemoglobin called hemoglobin S (HbS) (3). When deoxygenated, HbS molecules polymerize into long fibers (4), which can span the size of the red blood cell (RBC), distorting it into the classical crescent shape at rest (5, 6). Such polymerization reduces the deformability of RBCs and reduces their lifespan (7–9), leading to anemia.

Originally, the stiffening of RBCs has been blamed for possibly causing vaso-occlusive events, qualifying SCA as a hemorheological disease (10, 11). In this scenario, the obstruction process would be a competition between hydrodynamic forces that push the cells through the smallest capillaries to provide oxygen to the tissues and the polymerization of the fibers that occlude further transport of blood cells either by steric confinement in smaller capillaries (12), or by collective jamming of larger post-capillary venules (10, 11), due to the loss of RBCs deformability and blood rheo-thickening.

However, during the last three decades, the pathophysiology of vaso-occlusion has proven to be more complex (13, 14). While nearly all patients with SCA demonstrate chronic anemia, the clinical expression of the disease differs from one patient to another. It is now well admitted that vaso-occlusive events depend not only on the mechanical properties of circulating sickle red blood cells (SRBCs) but also on their ability to adhere to each other and to the vascular walls, which depends on inflammatory and oxidative stresses (15–17).

Surprisingly, the role of the geometry where the vaso-occlusive events takes place has not been explored up to now. The microcirculatory network is highly branched and has a reticulated nature, because it is designed to perfuse efficiently and deeply into tissues. In these microcapillaries, 80% of the total pressure-drop between the aorta and the vena cava occurs (18), making it a critical place for disturbances. In addition, such microvessels generally present strong bends and wavy shapes, with short segment lengths between bifurcations, which tends to attenuate the formation of a cell-depleted layer adjacent to the vessels wall (19). These features could play a major role in vaso-occlusion when coupled to the adhesive properties of SRBCs. Even though the structure and geometry of the microcirculatory network has been poorly documented in SCA patients, preliminary results in the skeletal muscle of patients with sickle cell trait (20), in the liver (21) or in the conjunctival microcirculation (22) of patients with Sickle Cell Disease, indicate important changes: a rarefaction of microvessels in the network, a decrease of their tortuosity, and an enlargement of their diameter, all pointing to their physiological relevance which remains to be understood in the context of RBC adhesion, stiffening and flow. There is a clear need to integrate both the rheological and the adhesive points of view in a more unified vision and to study the vaso-occlusion dynamics both at the cellular scale and in flow, since the problem of obstruction is in essence a circulatory one.

We used microfluidic channels as a first step to explore the potential effects of flow geometry coupled to adhesion and polymerization on vaso-occlusion. We focused our attention to test the jamming scenario of vaso-occlusion in straight channels by designing microchannels with gas exchange that mimic physiological conditions of the microcirculation in terms of flow velocity, oxygen
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concentration and hematocrit. While none of our experiments in straight channels showed clogging at physiologically relevant velocities, we observed intense deposition of SRBCs and the formation of SRBCs aggregates around acute corners where the flow changes rapidly direction typical of bifurcations in vivo. Such aggregates might play a major role in vaso-occlusive events, linking adhesive properties of the membrane of the cells with flow characteristics.

Materials and methods

Blood samples

Blood samples were collected during the normal course of patient care at the Montpellier University Hospitals Hematology Laboratory (Montpellier - France), according to a research protocol (IRB 12/12-04) approved by the Partners Healthcare Institutional Review Board. Blood is collected in EDTA tubes and the Mean Corpuscular Hemoglobin Concentration (MCHC) as well as the HbS percentage were determined by the hematology laboratory. All the samples containing HbS came from seven different patients with a homozygous genotype (HbSS) just before their exchange transfusion. The samples used in this study and their properties are listed in table 1. RBCs are washed by successive centrifugation in a Phosphate Buffered Saline (PBS) solution containing 5 mM glucose, the final osmotic pressure being adjusted to 300 mOsm. RBCs are then resuspended at 50% hematocrit. In order to get samples with a narrow MCHC, RBCs are separated according to their density (and therefore according to the hemoglobin concentration) on a discontinuous Percoll (Sigma # 77237) density gradient, following a protocol adapted from Corbett et al. (6). The gradient is centrifuged at 4000 g for 1 hour at 4°C and then, SRBCs settling between the Percoll layers of density 1.092 and 1.101 (corresponding to a MCHC of 0.34 g/cm³) were harvested with a micropipette. To get rid of the Percoll, the SRBCs are washed 5 times in a PBS + 5mM glucose solution adjusted to 300 mOsm and finally resuspended at 50% hematocrit for storage at 4°C. Blood samples were used within a maximum of 5 days after venipuncture. Before any experiment, the SRBCs were diluted to a hematocrit value of 25% typical of the physiological microcirculation.

Samples for experiments which address the question of the presence of free HbS in solution were prepared using healthy RBCs washed as previously described. Before the experiment, RBCs were suspended in a buffer which consisted in a 3-day-old supernatant of washed SRBC sample (PBS + 5mM glucose at 300 mOsm), containing a small amount of free HbS in solution, due to hemolysis (sample 6).

Samples used to investigate the effects of plasma proteins, platelets and white blood cells were prepared from washed SRBCs which are suspended in autologous plasma containing platelets and white blood cells. The resulting white blood cell count was estimated by visual counting under a microscope and was of the order of 0.1%. 
### Table 1: List of blood samples used during the experiments

<table>
<thead>
<tr>
<th>sample</th>
<th>% HbS</th>
<th>extra free HbS</th>
<th>autologous plasma</th>
<th>patient</th>
<th>test</th>
</tr>
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<tr>
<td>1</td>
<td>79.7</td>
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<td>patient 1</td>
<td>clogging</td>
</tr>
<tr>
<td>2</td>
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<td>no</td>
<td>patient 2</td>
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</tr>
<tr>
<td>3</td>
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<td>no</td>
<td>patient 3</td>
<td>clogging</td>
</tr>
<tr>
<td>4</td>
<td>57.9</td>
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<td>patient 4</td>
<td>aggregate formation</td>
</tr>
<tr>
<td>5</td>
<td>69.0</td>
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<td>no</td>
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</tr>
<tr>
<td>6</td>
<td>0</td>
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<td>no</td>
<td>healthy 1</td>
<td>aggregate formation</td>
</tr>
<tr>
<td>7</td>
<td>58.2</td>
<td>no</td>
<td>yes</td>
<td>patient 4</td>
<td>aggregate formation</td>
</tr>
<tr>
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<td>62.8</td>
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<td>no</td>
<td>patient 6</td>
<td>aggregate formation</td>
</tr>
<tr>
<td>9</td>
<td>90.0</td>
<td>no</td>
<td>no</td>
<td>patient 7</td>
<td>aggregate formation</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>no</td>
<td>no</td>
<td>healthy 1</td>
<td>control (no HbS)</td>
</tr>
<tr>
<td>11</td>
<td>0</td>
<td>no</td>
<td>no</td>
<td>healthy 2</td>
<td>control (no HbS)</td>
</tr>
<tr>
<td>12</td>
<td>0</td>
<td>no</td>
<td>no</td>
<td>healthy 3</td>
<td>control (no HbS)</td>
</tr>
</tbody>
</table>

### Microfluidic channels

Devices are fabricated using the classical soft-lithography with replica molding method (23) (the detailed procedure is described in the supporting materials). We mold the devices using a silicone elastomer, polydimethylsiloxane (PDMS), which is porous to gases (24). We use this feature to control the oxygen concentration in the channels via a two-layer PDMS structure. The first layer of 60 µm in thickness is engraved with channels of 30 µm in height where RBCs should flow. This layer is bonded to a glass microslide by plasma bonding. The plasma treatment has to be fast (few seconds) because the fast decrease of permeability of PDMS with the exposure time to the corona discharge (25). The second layer, dedicated to the gas flow, is plasma bonded on the top of the first one and engraved with a long serpentine channel of 500 µm in width and 80 µm in height. It covers the entire surface of the chip (see figure 1) to maximize gas exchange. RBC channels and gas channels are thus separated by a 30 µm thin PDMS membrane through which oxygen diffuses. The geometries of the microfluidic chips are shown in the figure 1. They all contain a “deoxygenation chamber” which consists of a large rectangular section 2 cm long and 4 mm wide, supported by pillars to prevent the top surface from collapsing onto the glass slide. Downstream from the deoxygenation chamber, we have the microcirculation section. On the first chip (Chip 1), the microcirculation section consists of a region containing triangular pillars to mimic corner flows, followed by a section of 16 parallel channels 1 cm long and 30 µm wide (see figure 1b) to test the clogging hypothesis. On the second chip (Chip 2), the microcirculatory section consists of successive branchings of 20 µm wide channels (figure 1c).
Figure 1: **Microfluidic channel geometry.**

**a.** Overlay of the geometry of the gas channel (light gray) and the channel network for the RBC suspension (black). The gas channel, a coil of 500 µm wide and 100 µm high, covers the entire surface of the chip to promote and control gas exchange within the RBC channels.

**b.** Geometry of RBC channels of chip 1. RBCs are deoxygenated during their flow through the gas exchange chamber. In the zoom-in window, microcirculation section consisting of a region of triangular pillars followed by 16 parallel channels 1 cm long and 30 µm wide.

**c.** Chip 2: The microcirculation section consists of 20 µm wide channels which successively branch. Scale bars 2 mm.
RBCs deoxygenation protocols

To optimize the deoxygenation rate, the gas exchange surface is large and the membrane separating the gas channels from RBCs flow is 30 µm thick. In such a device, gas exchange is known to occur over tens of seconds (24, 26, 27). This time scale has to be compared with the time for a RBC to flow along the deoxygenation chamber which is of the order of minutes. When the RBCs reach the microcirculatory part, they are already deoxygenated if pure nitrogen is injected in the serpentine channel. We initiate the gas flow 30 min before the injection of RBCs to saturate the PDMS chip. The gas flow is maintained throughout the experiment. RBCs are therefore deoxygenated while flowing.

Experimental setup

Observations are performed on a inverted microscope and the temperature surrounding the microfluidic chips is maintained at 37°C by an air flow in all experimental conditions. The gas channel is directly linked to a pressure controller (Fluigent-4C, 4 independent channels) and RBCs are injected from a pressurized tube which is linked to the pressure controller. Microfluidic channels have been designed to obtain a configuration with characteristic length, velocity and hematocrit conditions similar to that encountered in the microcirculation (18). The total pressure difference used to induce the flow is 130 mbar. Given the typical length of the devices we use, this value yields a pressure drop per unit length of the order of 0.1 Pa/µm typical of the microcirculation (28). Characteristic velocities of RBCs measured in the middle plane of the 30 µm channels are, depending on this input pressure, of the order of 7 mm/s to mimic the post-capillary venules blood circulation (28). Around the triangular pillars studied here, the velocity ranges from 100 to 400 µm/s, closer to capillary flow velocities (29).

Results and discussion

We first test the clogging hypothesis at the microcirculation scale using Chip 1. While no occlusion has been observed in the 30 µm wide microchannels, even after 2 hours of flow, we observe the deposition of SRBCs into large aggregates at acute corners in the microfluidic device. These aggregates contain both SRBCs and healthy RBCs, since our samples are all from exchange transfusion patients. Nevertheless, we verified that both healthy and sickled RBCs do not enter the microchannels as clusters and as we will show later, neither deposition nor aggregation in flow occurs with healthy RBCs only.

Role of the flow geometry

We focus our attention on the triangular pillars whose acute corners both upstream and downstream mimic locally the fast variation of direction of the streamlines observed at bifurcations in the microcirculation. We observe the formation of SRBCs aggregates on one pillar first at the two corners situated downstream (Chip 1) and later at the opposite corner upstream (Chip 1 & 2) as
shown in figures 2(a-c) and 3. The deposition is slow and a delay time of 20 minutes is observed before the onset of growth which results from successive and accumulative adhesion of SRBCs. The cells form layers and the flow pushes slowly the whole aggregate on each side towards the center of the triangular pillar base. This leads to the formation of a single large aggregate after about one hour of flow (figure 2 c. and Movie S1). Deposition at the upstream front of the pillar results in the formation of an aggregate on each side (figure 3 a-c). This deposition is observed to be slower than the one downstream. When observed in branching channels (Chip 2), these aggregates reach a critical transverse size up to half of the total channel width (figure 3 d). This is due to a continuous increase of the shearing force during deposition due to channel lumen reduction. Above this critical size, we observe some additional SRBCs which transiently bind to the aggregate with a residence time ranging between 2-13 seconds. The intricate interaction between the flow and the aggregate development at branches lead us to focus this study of deposition around the pillars walls where the flow is not confined.

Interestingly, no deposition is initiated on the substrate or the ceiling of the chips on the timescale of the experiments. Deposited aggregates are observed waving in the flow and are found to be best focused around the half-height of the channel. They can break or detach from the pillars but this remains a rare event.

We characterize the temporal evolution of the size of different SRBCs aggregates at the rear of the pillars using samples 4, 8 and 9 at 25 % hematocrit (similar to venules hematocrit) as shown in Figure 2g. SRBCs are deoxygenated while flowing. Initially, only few RBCs adhere behind the pillar, and during 15-20 minutes there is no major evolution. Then, an aggregate develops from each corner of the triangular pillar and finally merge together after one hour. This sequence is shown for sample 4 in Fig. 2(a-c). The resulting aggregate still grows to reach a steady length of several tens of microns. The aggregate redefines the boundary of the triangular pillar, reducing the local curvature of the streamlines and hindering further deposition. The aggregate length saturates when the angle of the corner initially close to 60° reaches approximately 120°. Control experiments were performed on the same timescales with healthy RBCs (same MCHC). They show that only few RBCs stick at the corners of the pillars (blue curve on figure 2 g, pictures in Fig. 2(d-f) and Figure S2).

Our microfluidic assay points out that acute corners are a location of much higher capture frequency and for aggregates to grow as also observed for biofilm growth of dilute solutions of bacteria (30). The formation of SRBCs aggregates on the vertical walls of the pillars, where the flow changes direction, is to be correlated with the few in vivo images of the onset of deposition available in the literature (31–33). In these studies, aggregates are present at blood vessel bifurcations where strong bends are present, similar to the flow geometry around the triangular pillars in our microfluidic chips.

The hydrodynamics of concentrated suspensions around corners at low Reynolds number seems to play a fundamental role in the cellular deposition. To describe the flow in this region, we use cross-correlation image velocimetry (using ImageJ and the Particle Image Velocimetry plugin) around one pillar to obtain the local average velocity field and the associated streamlines for SRBCs flowing in the middle plane of the channel. The streamlines are reported in figure 4a, together with
Figure 2: **Morphology and time evolution of aggregates at the rear of pillars.** a-c SRBCs from sample 4: pictures of aggregate formation at the rear of a pillar. For $t = 10$ min, only a few cells are in adhesion on the PDMS pillar. After 30 min an aggregate has grown on both corners at the rear of the pillar. After 60 min the two aggregates merge together and the length remains stable. Scale bars 20 $\mu$m. d-f Healthy RBCs deposition on pillars (sample 10). Only few cells stick to the pillar. Scale bars 20 $\mu$m. g. Temporal evolution of the length of SRBCs aggregate at the rear of pillars (sample 4, red circle; sample 8, orange triangle; sample 9, green square. 25 % hematocrit). The aggregate length is defined as the maximum extent from the base of the pillar and the fluctuating end. For $t < 15$ min, only few cells are in adhesion on the PDMS pillar. Then the growth initiates and aggregates can reach a size of about 40 $\mu$m. For each curve, the length is averaged over 7 different aggregates. Control experiment, with healthy RBCs: only few of them stick to the PDMS pillar (blue diamonds, sample 10, 25 % hematocrit).
Figure 3: **Morphology of SRBC aggregates at the front of pillars.** a-c Aggregation overtime at the front of a pillar (sample 4). d. Aggregate at a branching point in 20 µm wide channels (geometry Fig. 1c.), observed after 2 hours of flow. In this geometry, the aggregate can grow up to half of the channel width (sample 5). Scale bars 20 µm.

The trajectories for 7 SRBCs followed by hand at different initial distances from one of the pillar edge. We compare these lines with those of the suspending fluid flow only (without cells, white streamlines in figure 4a), computed using an in-house 3-D flow solver (34) (YALES2BIO, details in supporting material) around a pillar of the same geometry. The creeping flow (the Reynolds number is of the order of 0.0028) is computed around an infinite array of pillars, using periodic conditions in the directions tangential to the wall. The comparison reveals that downstream of the pillar, the SRBCs streamlines are contracted towards the pillar basis compared to the 3-D computed flow of a simple Newtonian fluid in the same geometry. On average, there is a net transverse drift of the cells towards the base (on each side of the pillar) in the region of maximum rotation of the streamlines. This observation could explain the higher collision rate of the SRBCs in this region leading to a higher deposition rate. Furthermore, the cells can explore the otherwise excluded region behind the pillar around the corner. Deformable objects such as RBCs circulating near a flat wall are known to be driven away from the wall by a lift force (35, 36) at the origin of the cell-free layer in the microcirculation. This force prevents RBCs from adhering to vessels wall. This is not the case here, notably due to the presence of corners and multi-body interactions. The contraction of the streamlines we see, compared to the 3D Newtonian case seems, to indicated the non Newtonian nature of this effect.

To illustrate the link between flow geometry, inverse-lift and multi-body interactions, we perform complementary simulations of one and two deformable two-dimensional capsules flowing around a pillar using our in-house flow solver. The capsules are supposed to be inextensible and resist bending. Their dynamics and deformation is computed using a full fluid-structure interaction approach and results are displayed in figure 4 b and c. All the technical details and assumptions are provided in the supporting materials. In the case of a single capsule, the shortest distance between the capsule membrane and the wall always increases, except at the corner, where the flow
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slightly accelerates. On the other hand, a capsule may follow a different trajectory if a second capsule follows it. In the corner case displayed in figure 4b, the front capsule stays closer to the corner and remains close to the pillar wall beyond the corner in a zone of lower speed. If the cells are close to each other at the right moment when passing the corner, their “hydrodynamic collision” produces a transitory pairing (due to lubrication forces) which drives the cluster in an otherwise forbidden region during rotation. We then compared this situation with a second test case shown in figure 4c with the same capsules in a simple 2-D Couette flow between two flat walls (configuration shown in figure S5(b)). The model, grid resolution and the characteristic parameters (shear rate, capillary number, Reynolds number, bending coefficient) are strictly identical to the case of flowing capsules around a pillar. In this Couette flow, the shear rate is constant and identical to the one seen by the capsules when they flow along the lateral wall of the pillar. Couette flow simulations were used as control cases, to confirm that the specific results obtained around the pillar are related only to the flow geometry. In the Couette flow, capsules simply interact by switching positions, as observed in computations of two interacting vesicles or capsules (37) in the presence of a wall. The specific phenomenon shown in the paper of capsules exploring regions very close to the pillar wall is thus the result of a combination between geometrical characteristics of the corner and multi-body effects. This behavior increases the probability for the cells to stick to each other and to adhere at this specific location and may participate in the formation of an aggregate at the corner.

In general, it has been demonstrated that the highly bifurcating topology of the microcirculatory network causes RBCs from side branches to be frequently introduced into the region closest to the wall at each branch, as shown for instance in the vascular bed of the rat spinotrapezius muscle (19), preventing the development of a significant cell free layer in networks. This phenomenon is also observed in vitro around our pillars or at our microfluidic bifurcations. Such modifications would also promote adhesion, but further studies both in vitro and in vivo are necessary to understand how flow patterns such as the bifurcation angle and flow direction could promote deposition and lead to occlusion.

Role of oxygen

We test the role of oxygen on the formation of aggregates using Chip 1 (figure 5). Starting from a long aggregate formed under deoxygenated conditions, we replace the nitrogen in the gas channel with air. First, the aggregate becomes more floppy due to the depolymerization of HbS within the SRBCs. Meanwhile, the cells detach and the aggregate shortens during 10 minutes. Finally, the aggregation process restarts in oxygenated conditions and the aggregate recovers the same size reached under nitrogen conditions. More generally, oxygen does not prevent aggregate formation at the pillar corners. Indeed, when we initiate the flow under oxygenated conditions, we observe the growth of RBC aggregates of the same size than in the absence of oxygen (see Figure S3). It is interesting to notice that aggregates made of oxygenated SRBCs exhibit a different appearance. Oxygenated SRBCs are more deformable due to the absence of HbS polymer and thus they better align according to the streamlines into layered stacks of cells (Figure 5 d and Movie S2). Noteworthy as in the deoxygenated case the aggregate surrounds the pillar, reducing locally the curvature and
Figure 4: **SRBCs flow around corners.** - **a.** Triangle symbols represent individual trajectories of SRBC. Red streamlines are the SRBC flow field computed by PIV. The SRBC flow field contracts at the rear of the pillar compared to the creeping flow of a Newtonian fluid (white streamlines). **b.** Simulation of deformable 2-D capsules flowing around a corner: the distance $d$ of a capsule to the pillar wall, normalized by the capsule characteristic size $R$, is represented over time. The comparison between the trajectory of a single capsule (black curve) and trajectories of two consecutive capsules (red and blue curves), points out multibody effects: in the displayed case, consecutive capsules flow closer to the corner wall than a single capsule. Moreover, when two consecutive capsules flow around the corner, the second capsule may push the first one closer to the wall at the corner (inset 2b) and at the rear (inset 2d), as in the simulation shown here. **c.** Simulation of deformable 2-D capsules flowing along a flat wall in a Couette flow: same legend as **b.**
Figure 5: Oxygen does not prevent aggregate formation - a. Time evolution of SRBCs aggregates at the rear of a pillar in the absence or presence of oxygen (sample 4). Under deoxygenated conditions, the aggregate grows as previously described, to reach a stable length of 45 µm. Then, nitrogen is turned off in the gas channel and replaced by oxygen at $t = 80$ min. HbS polymer in SRBCs depolymerizes, which results in more deformable SRBC and a rearrangement of the aggregate. The length of the aggregate decreases to 20 µm before recovering its initial length 20 minutes after oxygen injection. b-d Snapshots of the aggregate morphology in presence or absence of oxygen, the time refers to the evolution in a. Blood sample number 4 at 25% hematocrit is used. Results of a control experiment with healthy RBCs are also shown (blue triangles, sample 10).

Role of the presence of free hemoglobin and the products of hemolysis

People suffering from sickle cell anemia have a higher amount of free hemoglobin in their blood plasma due to a higher intravascular hemolytic rate (38). Furthermore, some studies report on the possibility for hemoglobin to bind to the RBC membrane (39, 40). We investigate in this section whether the presence of products of hemolysis (PHemo) which contain free HbS could be an enhancing factor for the formation of cellular aggregates. In the channels, we inject healthy RBCs suspended at 25 % hematocrit in the supernatant of initially washed SRBCs samples kept in the fridge (PBS + 5 mM glucose + PHemo see Materials and methods section) which contains hindering further growth.
free HbS due to hemolysis (sample 6). The evolution of the average aggregate length over time is then reported in figure 6. Aggregates grow after about 20 minutes, as previously observed in the case of SRBCs, but reach a constant length of 18 ± 1.5 µm after 80 minutes, 3 times shorter than for the SRBCs aggregates. When the same experiment is performed with RBCs suspended in a supernatant of healthy RBCs (see Materials and methods) containing free HbA instead of free HbS and all the same products of hemolysis, we do not observe aggregate formation similar to the former case (blue curve in figure 6).

Free HbS and its derivative products seem to play a role in the adhesive nature of RBCs. However, this is not sufficient to obtain aggregates on the order of 50 µm. HbS is usually submitted to a high rate of auto-oxidation, particularly when it is not encapsulated in RBCs. The generation of oxygen free radicals may cause severe RBCs membrane damage, hence changing the rheological and adhesive properties of flowing RBCs. It was previously reported that oxidative stress is able to decrease RBCs deformability (41), increase RBCs aggregation strength and promote the adhesion of RBCs to endothelial cells (42, 43). Indeed, it may be speculated that increased adhesion and RBC aggregate formation at a corner in the presence of free HbS could be related to increased oxidative stress. These questions need further investigation in order to better understand the pro-adhesive properties of SRBCs.
Effect of autologous plasma in combination with platelets and white blood cells?

Since the 90s, numerous factors have been identified as potentially involved in vaso-occlusive mechanisms (14, 17). These include the presence of proteins in the plasma, platelets and WBCs. We explore the potential role of these elements on the aggregate formation in our microfluidic chips by adding autologous plasma containing platelets and WBCs to SRBCs (blood sample 7). The final WBCs count was around 0.1%. We observe the formation of cellular aggregates but the morphology is completely different. In this case, aggregates form long chains, waving in the flow, which can reach several hundreds of microns (figure 7 b and Movie S3). SRBCs in these aggregates are elongated and adopt a tear drop shape, forming tethers. On figure 7, we can distinguish the membrane tube linking the pillar to the first RBC as well as a WBC in the aggregate. Contrary to previous observations, the growth of these aggregates appears to be randomly distributed in time. They can form within the first minutes of flow and on a timescale of seconds. In the control experiment with healthy cells, we do not obtain such structures.

We still do not know the respective role of platelets or WBCs in the formation of these structures and the origin of the membrane forming the filaments. However, recent observations show a preferential adhesion of WBCs at bifurcations (44), independent of the endothelium state, which could result from local hydrodynamical conditions. In addition, RBCs-WBCs interactions are known to be important in post-capillary venules (32) and WBCs from SCA patients seem to be more adhesive to SRBCs (45) and are capable of forming aggregates of few cells in whole blood (46). Moreover, the drastic morphology change we observe for the aggregate could be correlated to the known activation of platelets observed during blood storage at 4°C (47, 48). It has been reported in particular that activated platelets are able to form aggregates in a shear micro-gradient (49), in locations analogous to the corner of the pillars of the present device, but for much higher Reynolds number flow conditions, not relevant for the microcirculation. Further experiments are required to determine the specific role of the platelets or WBCs during aggregate formation. A first step would be to label their membrane with different dyes to track their positions within an aggregate.

Vaso-occlusion: two hypothetical scenarios

All the observations reported above provide information to propose two hypothetical scenarios for vaso-occlusion to occur at bifurcations in vivo, as illustrated in figure 8a. Aggregation present at corners could either reduce vessels diameter (as seen at bifurcations in figure 3d) until occlusion, and/or large aggregates could be shed randomly in time disturbing the flow downstream. We observed in rare occasions the breaking of the aggregate (figure 8b and Movie S4) and its passage through a 30 µm channel. We succeeded in recording one passage event in a slightly different chip where the region of 30 µm channels was replaced by an array of 20 µm channels flowing at different intersection angles into a main output channel (see geometry in Supplemental Information). This passage is shown in figure 8 c. When the aggregate enters upstream of the field of view, the entire flow in the channel decreases. When the aggregate reaches the field of view, we observe downstream of it a depletion in SRBCs, while upstream the SRBCs density is higher. This critical situation leads to a dramatic drop of the average flow velocity (down to 15% of its initial value as shown in figure
Figure 7: Morphology of SRBCs aggregates in the presence of autologous plasma, platelets and WBCs - a. SRBCs from sample 7. The aggregates are long chains fluctuating in the flow. They can reach several hundreds of microns long. We can distinguish the membrane tube linking the pillar and the first RBC. The picture has been taken after the RBCs had cleared the channel but there is still remaining media flowing in the channel. This allows a better observation of the aggregate structure. l defines the length of the aggregate. Scale bar 20 µm. b. Aggregates length in presence of autologous plasma, platelets and ≃0.1 % WBCs, sample 7 (green diamonds). The control correspond to healthy RBCs (blue triangles, sample 10).
and could also promote further HbS polymerization and aggregation of the SRBCs behind the aggregate. This would lead to flow regimes close to the ones where jamming was observed in microfluidic devices (26). However, we were never able to observe total occlusion in our devices, but this is consistent with the fact that blood samples came from patients which are regularly transfused to prevent frequent vaso-occlusive crises. In addition, other factors listed by Switzer et al. (38) like platelets aggregation, vasoconstrictive molecules (endothelin family proteins), decreased nitric oxide bioavailability, will certainly act together with the hydrodynamic conditions at corners to promote deposition and aggregation of SRBCs and to participate in a vaso-occlusive crisis.

**Conclusion**

Microfluidics is a suitable tool to study blood circulation at the microcirculatory level. In this study, we reported on SRBCs aggregation at bifurcations and around acute corners. We have shown that the geometry of the flow is a key factor for the formation of SRBCs aggregates. The onset of aggregation results from an intricate interplay between geometry, collective motions and pro-adhesive properties of SRBCs and remains to be fully explored and understood.

Moreover, the origin of SRBCs adhesiveness at the heart of our observations is still an open question and measuring the strength of adhesion would give some better insights on its importance in the context of cellular deposition in flow. In addition, it would be interesting to extend this microfluidic study to different hematocrit and healthy/sickled RBCs proportions, as well as to research actively the signs of such corner-flow-enhanced deposition and aggregation in vivo and its consequences on vaso-occlusion.

Finally, we hope that our preliminary results will motivate future studies on the role of flow geometry in cell capture not only during vaso-occlusion but also in other physiologically relevant situations like the inflammatory response.

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**Authorship contributions**

E.L., G.M. and M.A. designed the study. E.L. realized the experiments. S.M. performed the numerical simulations. All the authors discussed the results and wrote the paper. The authors declare no competing financial interests.
Figure 8: **Suggested scenario** - a. The formation of microthrombi at a corner or at branching points can drastically impair the hemodynamics, due to a possible severe restriction of the lumen area. b. Example of aggregate detachment (blood sample 8 at 25 % hematocrit): when reaching a critical size, cellular aggregates can break. The scale bar are 20 µm. c. Free flowing aggregate in microcirculation. Relative velocity of an aggregate flowing along a 20 µm wide channel. Initial velocity is 200 µm/s and SRBCs were from blood sample 8 at 25 % hematocrit. White arrows represent the direction of the flow.
References


