Clustering of microscopic particles in constricted blood flow

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Abstract

A mixed suspension of red blood cells (RBCs) and microparticles flows through a cylindrical channel with a constriction mimicking a stenosed blood vessel. Our three-dimensional Lattice-Boltzmann simulations show that the RBCs are depleted right ahead and after the constriction. Although the RBC mean concentration (hematocrit) is 16.5% or 23.7%, their axial concentration profile is very similar to that of isolated tracer particles flowing along the central axis. Most importantly, however, we find that the stiff microparticles exhibit the opposite behavior. Arriving on a marginated position near the channel wall, they can pass through the constriction only if they find a suitable gap to dip into the dense plug of RBCs occupying the channel center. This leads to a prolonged dwell time and, as a consequence, to a pronounced increase in microparticle concentration right in front of the constriction. For biochemically active particles such as drug delivery agents or activated platelets this clustering may lead to physiologically relevant hot spots of biochemical activity.
I. INTRODUCTION

Blood mostly consists of highly deformable red blood cells (RBCs) occupying from 15 up to 45 volume percent [1–4]. When flowing through a channel or blood vessel, RBCs preferentially travel in the low-shear zone around the channel center. Stiff particles such as white blood cells, platelets or synthetic microparticles are thereby forced to flow near the channel walls. This well-known effect, called margination, is of physiological relevance, e.g., for platelets where a near-wall position allows them to quickly stop bleeding in case of vessel injury. For synthetic particles such as drug delivery agents a near-wall position is essential to directly deliver their cargo to endothelial cells, thus minimizing unwanted distribution of pharmaceutical agents through the vascular system.

In 1980 margination was first observed experimentally [5]. Since then, a large number of studies investigated the phenomenon by means of experiments [6–14] or simulations [14–36]. All these works focus on spatially constant geometries like straight pipes, rectangular channels or plane-Couette systems. However, more complex geometries such as bifurcations or constrictions (stenoses) are a common feature of the vascular system. At the same time, investigations of marginated particles near such geometrical features are scarce. For RBCs a recent detailed study in cylindrical microchannels with stenosis exists [37]. Ref. [38] presented 2D simulations near aneurysms while ref. [39] investigated the recirculation zones after a sudden expansion. Near a stenosis ref. [40] presented local velocity fields while other studies treated the dynamics of platelets in 2D and plane-Couette geometries [41, 42].

Here, we use 3D Lattice-Boltzmann simulations to study margination of stiff spherical microparticles in a red blood cell suspension flowing through a cylindrical channel with a constriction mimicking the geometry of a stenosed blood vessel. While the radial concentration profile is found to be qualitatively unaffected (RBCs in the center, microparticles near the wall) we find that the concentration profile along the axial direction varies significantly for both RBCs and microparticles. Right in front of the constriction we observe a decrease in RBC concentration which can be almost quantitatively explained by the flow acceleration using passive point particles flowing along the central axis as the most simple model system. Microparticles, on the other hand, exhibit the opposite behavior: their concentration in front of the constriction increases significantly. This effect can be traced back to the near-wall trajectories on which microparticles travel in the straight channel section before
arriving at the constriction and seems to be strongly reinforced by the cylindrical geometry since a recent study in planar geometries [42] did not report it. In order to pass through the constriction, the particles need to leave this position and squeeze into the densely packed red blood cell layer. This lowers the microparticles’ passing rate leading to a pronounced peak in microparticle concentration ahead of the constriction.

MATERIALS AND METHODS

We perform our simulations by means of a 3D Lattice-Boltzmann method (LBM) implemented in the simulation package ESPResSo [43–45] extended by an immersed boundary method (IBM) algorithm [46, 47]. The LBM uses the spatially and temporally discretized Boltzmann equation to calculate fluid behavior [48–50]. The simulation package ESPResSo enables us to add thermal fluctuations with a temperature $T = 300$ K. The fluid in which the particles are suspended is chosen to mimic blood plasma with a density of $1000 \text{ kg/m}^3$ and a viscosity of $1.2 \times 10^{-3} \text{ Pa s}$.

Both RBCs and microparticles are realized via the IBM. The RBC diameter is chosen to be $7.82 \mu m$. The red blood cell membrane is modeled as an infinitely thin elastic sheet which is discretized by a set of nodes that are treated as Lagrangian points flowing with the fluid and that are connected by triangles. The forces between the fluid and the membrane nodes are calculated every time step by interpolating between the fluid nodes using an eight-point stencil. For the elastic properties we use the Skalak model [51–53] with a shear modulus $k_S = 5 \times 10^{-6} \text{ N/m}$ and an area dilatation modulus $k_A = 100k_S$. Bending forces are computed via the Helfrich model with a bending modulus $k_B = 2 \times 10^{-19} \text{ Nm}$ [2, 54] using the bending algorithm denominated method A in [54]. We note that, while this rather simplistic algorithm is clearly not the most accurate method, it should be sufficient for the present purpose where the major focus is on the behavior of the microparticles, not the RBCs themselves. In addition, we use an empirical volume conservation potential [34].

The spherical microparticles have a diameter of $a = 3.2 \mu m$ and are thus in a range for which strong margination was recently reported [28]. Numerically, they follow the same model as the RBCs, but with 1000 times higher shear and bending moduli. In addition, they are stiffened by a 3D lattice inside the particle with lattice constant of one LBM grid cell. The grid is connected to the shell by elastic springs and a bending potential. The
stability of our particle model is documented in [33]. A soft-sphere potential with cut-off radius of one grid cell between the particles and between the wall and a particle ensures numerical stability.

The constriction is modeled by two straight cylinders with different radii and a transition zone between them. To provide a smooth transition the radius in the transition zone follows a third order polynomial such that the function describing the boundary and its first derivative are continuous. To control the flow rate a pressure gradient is applied. The Reynolds number calculated from the centerline velocity, the RBC radius $R_{RBC}$, and the kinematic viscosity of the fluid $\nu$ is $Re = \frac{R_{RBC}v}{\nu} = \mathcal{O}(10^{-3})$. The time step is chosen to 0.06$\mu$s.

At the beginning of the simulation the RBCs and the microparticles are inserted with a regular pattern. The system is then run until all remnants of this regular initial configuration have vanished and a steady state is achieved. Concentration profiles are obtained by time-averaging over a rather long part of the simulations ($\sim 90\%$ of the total simulation time) starting from these random configurations. The systems are simulated over about 10s.

The axial and radial concentration is calculated by dividing the system in axial and radial bins. At every time step RBCs and microparticles are sorted into corresponding bins using their center-of-mass. The concentration is calculated by dividing the averaged number of particles inside one bin by the volume of the bin.
RESULTS AND DISCUSSION

A straight channel as comparison and two examples of the investigated systems with stenosis are shown in figure 1. In all cases the cylinder radius is \( R_C = 13.4\mu \text{m} \) and in (b) and (c) the ratio between the radius of the constricted part and the main cylinder is 1/2. The length of the region with constant lower radius here is 2.6\( \mu \text{m} \) but will be varied below. The length of the straight channel is 47\( \mu \text{m} \) and 117\( \mu \text{m} \) for the constricted channel. We verify in the supplementary material that varying the channel length has no influence on the results. The hematocrits are 16.5\% in (a) and (b) and 23.7\% in (c). The mean RBC velocity is 1.7\( \frac{\text{mm}}{\text{s}} \) in the main channel in (a) and 1.3\( \frac{\text{mm}}{\text{s}} \) and 0.8\( \frac{\text{mm}}{\text{s}} \) inside the constriction in (b) and (c), respectively.

Center-of-mass distribution in comparison to straight cylinder

We first extract two-dimensional concentration profiles in the \( x-r \) plane, where \( x \) is the coordinate along the channel and \( r \) is the radial position in cylindrical coordinates. Local concentrations of RBCs and microparticles are calculated from their center-of-masses as
described in the Methods section. The result for the straight channel is shown in figure 2 (a) for RBCs and in (b) for microparticles. The RBC concentration is rather homogeneous and the RBCs are located around the channel center since their deformability causes the RBCs to migrate to the center [55–57]. Near the channel wall, the well-known cell-free layer with almost zero RBC concentration is visible [58–60]. As can be seen in figure 2 (b) the microparticles are expelled into this cell-free layer leading to a sharply peaked concentration distribution of the marginated microparticles. The origin of microparticle migration into the cell-free layer is commonly attributed to heterogeneous collisions with the RBCs [17, 19, 22, 26, 31, 33, 36, 42].

The general picture looks similar for the constricted channel with Ht=16.5% as shown in figure 2 (c) and (d) (see supplementary material for Ht=23.7%). Besides a reduced cell-free layer inside the constriction, which agrees well with ref. [37], the RBC concentration in the channel center is not significantly influenced by the constriction. The peak at large radii in the microparticle concentration is however more blurred compared to the straight channel, especially right behind the constriction. This can be explained since the microparticles require a certain time (or, equivalently, distance) in order to regain their fully marginated state after passing through the constriction.

Axial distribution in one-component suspensions

In order to assess more quantitatively a possible influence of the constriction, we average the 2D concentration profiles of figure 2 over the radial position to obtain one-dimensional axial concentration profiles. To discriminate between the generic influence of the constriction and the influence of margination we start by considering single-component suspensions where margination is not present. In figure 3 (a) we show the concentration profile for a pure RBC suspension which has been created by replacing all microparticles by RBCs in the system of figure 1 (b). Far from the constriction the concentration exhibits a constant plateau as expected in a straight channel. Upon entering the constriction, the concentration first decreases and then rises again until a roughly constant plateau inside the constriction is reached. At the exit, the concentration decreases a second time and soon after recovers its straight channel level. Removing the red blood cells from the system in figure 1 and leaving only the 18 microparticles leads to the concentration profile in figure 3 (b). The micropar-
Figure 3. Concentration as a function of the axial position for a pure RBC suspension (a) and for a pure microparticle suspension (b). Despite very different particle properties and mean concentrations both profiles are remarkably similar. (c) The axial flow velocity on the centerline for the pure liquid without thermal fluctuations (green curve). The concentration distribution for a train of passive tracer particles following this velocity profile (red curve) well reproduces the concentration profiles of the single-component suspensions in (a) and (b). The channel geometry is illustrated by the black line.

Particle concentration profile looks remarkably similar to that of the pure RBCs pointing to a common generic origin.

In order to explain these observations, we revert to the most simple model system: passive tracer particles flowing along the central axis through the constriction. For this, we first extract the velocity profile from a simulation without any particles and thermal fluctuations (Fig. 3 (c) green curve). We then compute the position as a function of time for a train of tracer particles which precisely follow this velocity. Due to periodic boundary conditions the particles repeatedly pass the constriction allowing us to compute the time-averaged concentration profile, which is shown as the red line in figure 3 (c). The profile of this simple system looks almost the same as the ones from the more complex RBC and microparticle suspensions. This furnishes a straightforward explanation for the concentration profiles: upon entering the constriction the fluid velocity rises due to mass conservation leading to an acceleration of a flowing particle. The acceleration increases the distance to the following particle and thus causes a concentration decrease. Inside the constriction, this effect however is (over-)compensated by the reduced channel radius leading to a plateau which is slightly higher than the straight channel value. At the exit of the constriction the opposite dynamics is observed. It remains remarkable that the concentration profile of the RBC suspension in figure 3 (a) and the microparticle suspension in figure 3 (b) can be well reproduced by passive tracer points. The only exception is that the RBC densities in front of and behind the...
constriction are slightly asymmetric while those of the particle train are perfectly symmetric as expected under Stokes flow conditions. Similar asymmetries in RBC concentration have also been observed by [37] and (in planar channels) by [42]. The most likely source for the asymmetry lies in the RBC deformability.

**Clustering of microparticles at constrictions in mixed suspensions**

We now consider the mixed RBC/microparticle suspension with Ht=16.5% as shown in figure 1 (b). Due to their position in the channel center the RBCs in this system are not affected by the small number of microparticles and exhibit the same behavior as in the pure RBC system (Fig. 3 (a)) as demonstrated by the red curve in figure 4 (a). In stark contrast,
the microparticle concentration shown by the blue line in figure 4 (a) is completely changed by the presence of the red blood cells. Instead of a decrease, the concentration in front of the constriction now exhibits a pronounced increase. At the exit of the constriction the same dip in the concentration appears as seen before for the pure microparticle suspension in figure 3 (c).

The increase of the microparticle concentration in front of the constriction, which is our main result, can only be explained by the interplay of margination and constriction. Due to their deformability the RBCs migrate to the channel center and are thus able to pass relatively unhindered through the constriction. The dense central plug of RBCs however acts as a barrier for the microparticles. Arriving on a marginated position at the channel walls, in order to pass the constriction, they need to penetrate the RBC plug. The collisions with the RBCs counteract this motion resulting in a longer dwell time in front of the constriction. This causes the peak in the microparticle concentration.

Indeed, this mechanism is demonstrated by the radial concentration profiles in figure 4 (c). Before and after the constriction the microparticles are clearly located inside the cell-free layer well separated from the RBCs. Entering the constriction the particles move towards the center and the particle peak starts to overlap with the broad RBC profile. Finally, inside the constricted part the microparticles and RBCs are located at similar radial positions, although the microparticles do retain a near-wall position in agreement with the planar situation investigated in [42] (although the authors did not observe the increase in concentration before the constriction). Regarding the RBCs themselves, we find that the cell-free layer thickness transiently increases at the exit of the constriction as in [37, 42]. The pausing in front of the constriction also reflects itself in the microparticle velocity in figure 4 (b). Whereas the RBC velocity monotonously increases ahead the constriction, the microparticle velocity exhibits a clear dip.

Thus, the effect of the same constriction is directly opposite for RBCs and marginated microparticles: while the RBCs are diluted, the concentration of the microparticles considerably increases in front of the constriction.
Figure 5. (a) Concentration and (b) velocity as a function of axial position for a mixed suspension with hematocrit $H_t=23.7\%$. (c) Concentration as a function of the radial position for various sites along the constriction. The RBCs (red curve) exhibit the same behavior as for $H_t=16.5\%$ in figure 4 except that the concentration inside the constriction is lower than in the main channel. The microparticles cluster in roughly the same amount ahead of the constriction. The channel geometry is illustrated by the black line.

**Influence of hematocrit**

By increasing the number of RBCs in the system to 151, we investigate the influence of a higher hematocrit $H_t=23.7\%$ as shown in figure 1 (c). The axial concentration for $H_t = 23.7\%$ in figure 5 (a) proofs that microparticle clustering also appears at higher $H_t$. However, differences occur in RBC concentration: in contrast to the lower $H_t$ in figure 4 (a), RBCs are diluted inside the constriction compared to the main channel. This effect is not captured by our simple isolated particle model in figure 3 (c) showing the importance of collective RBC-RBC interactions at higher $H_t$.

In figure 5 (c) the radial concentration profiles are shown for $H_t = 23.7\%$. The dipping of microparticles into the RBC plug is present here in the same manner as for $H_t = 16.5\%$ in figure 4 (c). While the velocity shown in figure 5 (b) is in general lower due to the larger
amount of cells, a dip in the microparticle velocity ahead the constriction is again observed. The results for Ht=23.7% suggest that an upper boundary for the extent of microparticle clustering exists. A possible explanation may be provided by the discrete motion of the RBCs through the constriction\cite{37}. Due to the limited space inside the constriction, even for higher Ht the number of simultaneously passing RBCs is not strongly increased. As a consequence the probability for microparticles to find a gap and thus the clustering effect is nearly the same at both hematocrits.

**Influence of constriction properties**

To investigate the influence of the constriction properties we conduct one simulation with a longer constriction, one with a narrower constriction and a third one with a longer (smoother) transition zone at Ht=16.5%.

In the first simulation, the constricted part is four times longer than in figure 1 (b). The observed axial concentration distribution is shown in figure 6 (a). The microparticle concentration increases in front of the constriction in the same manner as for the short constriction (Fig. 4). Inside the constriction the microparticle concentration decreases slowly. We thus conclude that the clustering effect is independent of the length of the constriction.

The narrower constriction is chosen such that the ratio of the radii of the constricted part and the straight part equals 3/8 instead of 1/2 before. The increase in the microparticle concentration is significantly stronger (Fig. 6 (b)) compared to figure 4. The explanation is straightforward: to pass the constriction a microparticle now has to migrate closer to the centerline. As explained before the collisions with the RBCs counteract this motion leading to a longer dwell time in front of the constriction.

Finally, we double the length of the transition zone between the main cylinder and the constricted part leading to a smoother transition. The observed concentration profile of the RBCs in figure 6 (c) is similar to that in figure 4 except that the decrease ahead and behind the constriction is less pronounced. This difference can be reproduced by the point particle method which shows the same changes in the concentration profile (data not shown). Importantly, the amount of microparticle clustering is not influenced by the smoother transition.
Figure 6. Concentration as function of the axial position for the RBCs (red curve) and microparticles (blue curve) for (a) a four times longer constriction, (b) a narrower constriction with a radii ratio equal 3/8, and (c) a twofold longer transition zone. The channel geometry is illustrated by the black line.

These results lead us to suggest that the radius of the constriction is the important parameter that determines the amount of microparticle clustering: the narrower the constriction the stronger the clustering. However, microparticle clustering is independent of constriction length and transition zone. Below we show that the clustering in front of the constriction occurs for ellipsoidal microparticles, as well.

Influence of particle shape and size

Figure 7. Center-of-mass density depending on the axial position for the RBCs (red curve) and the stiff ellipsoids (blue curve) in a red blood cell suspension flowing through a constriction. The ellipsoidal microparticles cluster, but not in the same amount as the spherical microparticles. An image of an ellipsoid is inserted. The channel geometry is illustrated by the black line.

Finally, we investigate the behavior of stiff ellipsoids to study a possible influence of the particle shape and size. For this, we use prolate ellipsoids with a ratio of two between the
long and the short axis (Fig. 7 inset). Along the short axis the ellipsoid diameter equals the diameter of the spheres used before. We find that also ellipsoids exhibit clustering in front of the constriction as shown in Fig. 7. Compared to the spherical microparticles the increase in concentration ahead of the constriction is however less pronounced. The slightly reduced clustering of the ellipsoids can be explained by the increased size of the ellipsoids along one principal axis. Due to this increased size the ellipsoids can dip more easily into the RBC plug and migrate towards the channel center. Nevertheless, the ellipsoid data suggests that clustering in front of constriction is a robust effect, which is qualitatively independent of the particle shape.

CONCLUSION

We investigated the axial concentration of red blood cells and microparticles in a mixed suspension flowing through a constricted cylindrical microchannel as a model system for a stenosis in the microcirculation. We found that the constriction causes a pronounced peak in the microparticle concentration right in front of the constriction. This clustering of microparticles is explained by the marginated near-wall position, on which the microparticles arrive at the constriction and the corresponding more central position of the RBCs. As a consequence, microparticle passage through the constriction is severely hindered by the dense plug of RBCs occupying the channel center. Hence, the microparticles get stuck in front of the constriction until they find a suitable gap among the RBCs leading to a prolonged dwell time, which in the end causes the increased concentration. Interestingly, the red blood cells themselves behave directly opposite showing a dip in concentration before the constriction. Furthermore, we investigated the dependency on hematocrit, stenosis, particle size, and shape. The clustering effect is qualitatively robust.

Our discovery of microparticle clustering will be especially relevant for biochemically active particles such as drug delivery agents, where the zone right in front of constrictions may become a hot spot of biochemical activity. We furthermore expect that similar clustering effects will also be present for other particles undergoing margination, such as blood platelets, which may increase the likelihood of thrombus formation ahead of vessel stenoses [36, 61].
SUPPLEMENTARY MATERIALS

See supplementary materials for influence of the total channel length. Furthermore, we provide the velocity and axial concentration profiles along the constriction for the simulations with different stenosis properties and with ellipsoidal microparticles.

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