**Functionalized supported membranes for quantifying adhesion of P. falciparum-infected erythrocytes**

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**ABSTRACT** The pathology of *Plasmodium falciparum* malaria is largely defined by the cytoadhesion of infected erythrocytes to the microvascular endothelial lining. The complexity of the endothelial surface and the large range of interactions available for the infected erythrocyte via parasite-encoded adhesins make analysis of critical contributions during cytoadherence challenging to define. Here, we have explored supported membranes functionalized with two important adhesion receptors, ICAM1 or CD36, as a quantitative biomimetic surface to help understand the processes involved in cytoadherence. Parasitized erythrocytes bound to the receptor-functionalized membranes with high efficiency and selectivity under both static and flow conditions, with infected wild-type erythrocytes displaying a higher binding capacity than do parasitized heterozygous sickle cells. We further show that the binding efficiency decreased with increasing intermolecular receptor distance and that the cell-surface contacts were highly dynamic and increased with rising wall shear stress as the cell underwent a shape transition. Computer simulations using a deformable cell model explained the wall-shear-stress-induced dynamic changes in cell shape and contact area via the specific physical properties of erythrocytes, the density of adhesins presenting knobs, and the lateral movement of receptors in the supported membrane.

**INTRODUCTION**

Tropical malaria is an infectious disease caused by the unicellular eukaryotic parasite *Plasmodium falciparum*. An estimated 229 million people were infected with *P. falciparum* in 2019, of which 409,000 patients died of severe complications (1). The pathology of *P. falciparum* malaria is associated with the intraerythrocytic life cycle of the parasite. As the parasite develops within red blood cells, it changes the structure and function of the host cell (2,3). Most notably, infected erythrocytes acquire cytoadhesive properties and sequester in the deep vascular bed by adhering to the endothelial lining of microcapillaries (2,3), thus escaping clearance by the spleen. Sequestered infected erythrocytes can obstruct tissue perfusion and elicit localized inflammatory reactions, among other pathophysiological sequelae, which together contribute to severe disease.

**SIGNIFICANCE** Adhesion of infected erythrocytes to the microvascular endothelial lining is the key event that defines pathology of *Plasmodium falciparum* malaria. To dissect critical contributions involved in the complex interaction between parasitized erythrocytes and the endothelial surface, supported membranes functionalized with important adhesion receptors ICAM1 or CD36 were used as a well-defined biomimetic surface. In combination with computer simulations, our experiments revealed that the cytoadhesion of parasitized heterozygous sickle cells is reduced compared to infected wild-type erythrocytes not only because of reduced receptor-specific adhesion but also because of changes in the cell shape. Thus, our work highlights how molecular and cellular aspects synergize in the adhesion of *P. falciparum*-infected erythrocytes.
Cytoadhesion is mediated by parasite-encoded immune-variant adhesins, of which a family of antigens, collectively termed *P. falciparum* erythrocyte membrane protein 1 ( PfEMP1), plays a dominant role (2–4). PfEMP1 variants can bind to a wide range of host receptors, including receptors on the surface of endothelial cells such as CD36, intercellular adhesion molecule 1 (ICAM-1), or endothelial protein C receptor (2–4). In addition, PfEMP1 variants can mediate binding of infected erythrocytes to uninfected red blood cells, leukocytes, platelets, and serum proteins (4). Each infected erythrocyte carries ~13,000 PfEMP1 molecules on its surface, presented in groups of three to four on parasite-induced membrane elevations, termed knobs (5). Knobs have a profound effect on the biomechanical properties of the membrane because they cause strain hardening by suppressing membrane fluctuations via increased membrane-cytoskeleton coupling (6,7).

Previous studies have shown that the sickle cell trait, which is caused by a single amino acid substitution of valine for glutamic acid at position 6 in the β-globin chain of the hemoglobin tetramer, protects carriers from severe malaria-related disease and death and that this protection is associated with altered knob morphology and a reduced cytoadhesion capability (8,9). In particular, parasitized erythrocytes containing the sickle cell hemoglobin S or related structural hemoglobinopathies present fewer and enlarged knobs (8,10), which, together with an overall reduced number of presented PfEMP1 molecules (5,9), results in a reduced ability to engage in cytoadhesive interactions. Yet, the underlying biophysical mechanisms are largely unknown.

The complexity of the endothelial surface with its multiple receptors, together with the range of interactions available to the infected erythrocyte via parasite-encoded adhesins, poses an investigative challenge and asks for a more amenable and better controllable system to define and compare critical contributions of different host receptors during cytoadhesion. Supported membranes (11,12) have been widely used to study membrane organization and receptor-ligand interactions, including cell adhesion phenomena (13–15). Because supported membranes can be fabricated from pure components, their composition can be controlled and adjusted according to experimental needs. This includes the option of functionalizing supported membranes with defined receptor proteins and varying the receptor orientation and density to represent better the variable nature of these interactions rather than the binary (binding or not binding) classification that is sometimes assumed. Given that supported membranes maintain the thermodynamic and structural properties of a free bilayer, they facilitate surface-sensitive techniques, such as cell detachment experiments via ultrasonic pressure waves to assess the intrinsic binding strengths (14,16) and reflection interference contrast microscopy (RICM) (17,18) to inform on cell-adhesive contact zones and their dynamics during shear stress or as a function of the receptor density. Thus, supported membranes offer a versatile tool to study aspects of cytoadherence that are not readily approachable in cell-cell binding assays. Despite these benefits, only one study has thus far applied the supported membrane technology to the *P. falciparum* system. By functionalizing supported membranes with chondroitin–4-sulfate (CSA), the main receptor for parasite cytoadhesion in the placenta, Rieger et al. (13) could show that binding of infected erythrocytes to CSA is cooperative, shear stress induced, and dependent on the receptor density under physiological flow conditions.

Here, we have further explored the application of supported membranes as a well-defined biomimetic surface to study quantitative aspects of cytoadhesion of *P. falciparum*-infected erythrocytes. To this end, we functionalized supported membranes with two representative host receptors presented on endothelial cells, namely CD36 and ICAM-1. We show that binding of infected erythrocytes to ICAM-1 or CD36 functionalized membranes is specific and strongly dependent on the receptor density, the mechanical loading, and the red blood cell type under both static and flow conditions. RICM revealed that the adhesion contact area of infected erythrocytes is very dynamic and defined by wall-shear-stress-induced shape transitions. We finally developed computer simulations using a deformable cell model to demonstrate that receptor redistribution due to cell shape changes explains the experimental data.

**MATERIALS AND METHODS**

**P. falciparum culture**

The *P. falciparum* IT4 (alias FCR3) strain was kept in continuous culture using HbAA and HbAS erythrocytes (19). Briefly, blood cultures were maintained at a hematocrit of 3.5% and a parasitemia not higher than 5% in Roswell Park Memorial Institute (RPMI) 1640 medium (supplemented with 2 mM L-glutamine, 25 mM HEPES, 100 μM hypoxanthine, 20 μg mL−1 gentamicin, and 10% (v/v) heat-inactivated human AB serum). Cultures were kept under controlled atmospheric conditions (37°C, 3% CO₂, 5% O₂, 92% N₂, and 96% humidity). Binding capability to CD36 and ICAM-1 was ensured by repeatedly selecting for knobs using the gelatin flotation method (20), followed by a panning procedure over TNF-activated human dermal microvascular endothelial cells (PromoCell, Heidelberg, Germany) (21). Trophozoite stage parasites were enriched using the magnetic cell sorting method (22), yielding a parasitemia >95%. The number of infected erythrocytes was determined using a cell counter from Beckmann Coulter Diagnostics (Brea, CA).

**Preparation of supported membranes**

1,2-Dioleoyl-sn-glycero-3-phosphocholine (DOPC), 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC), and 1,2-dioleoyl-sn-glycero-3-(N-(5-amino-1-carboxypentyl)liminodiacetic acid)succinyl] (nickel salt) (DOGS-NTA (Ni²⁺)), were purchased from Avanti Polar Lipids (Alabaster, AL). Recombinant human CD36 and human ICAM-1 with histidine tags were purchased from Thermo Fisher Scientific (Waltham, MA). The glass slides (Gerhard Menzel, Braunschweig, Germany) were cleaned with the modified Radio Corporation of America (RCA) method (23), then bonded to bottomless culture dishes or to μ-Slide VI0.4 flow chambers (ibidi, Martinsried, Germany) using polydimethylsiloxane produced from the base and the curing agent (SYLGARD184; Dow Corning, Midland, MI).
Stock solutions (5 mg mL\(^{-1}\) in CHCl\(_3\)) of DOPC and DOGS-NTA (Ni\(^{2+}\)) were mixed at different molar ratios \(\chi\), dried under a nitrogen stream, and stored in vacuum. The lipids were resuspended (1 mg mL\(^{-1}\)) in Hepes buffered saline (HBS) buffer (150 mM NaCl, 10 mM HEPES (pH 7.5)) and sonicated using a tip sonicator (Misonix, Farmingdale, NY), for 30 min to create small unilamellar vesicles. Residual titanium particles were removed by the centrifugation at 13,400 \(\times\) g for 10 min. Supported membranes were prepared by vesicle fusion by injecting lipid suspensions into the sealed chambers for 1 h at room temperature, where supported lipid membranes were prepared by vesicle fusion by injecting lipid suspensions and sonicated using a tip sonicater (Misonix, Farmingdale, NY), for 10 min. The areas of adhesion contact and tight binding zone were determined by using the camera. To investigate the influence of shear stress on the adhesion contact and tight binding zone, 100 consecutive images with an integration time of 30 ms were collected for at least three different positions after applying a given shear stress \((r = 0.05–1.0\) Pa) for 10 min. The mean-square amplitude (MSA) of the light intensity was calculated for each pixel inside the projected cell contact area. The regions with MSA values below the threshold level (MSA < 0.02) were categorized as tight binding zones. The space occupied by a parasite was removed manually before analysis, as shown in Fig. 4A.

**Determination of cell-surface contact**

The areas of adhesion contact and tight binding zone were determined by using label-free RICM as reported previously (14). The sample was illuminated with monochromatic, linear polarized light (\(\lambda_{\text{ex}} = 546\) nm) through a 63×/1.25 Antiflex oil-immersion objective with a built-in \(\lambda/4\) plate (Zeiss). Internal stray light is suppressed using an additional linear polarizer shifted 90° in front of the camera. To investigate the influence of shear stress on the adhesion contact and tight binding zone, 100 consecutive images with an integration time of 30 ms were collected for at least three different positions after applying a given shear stress \((r = 0.05–1.0\) Pa) for 10 min. The mean-square amplitude (MSA) of the light intensity was calculated for each pixel inside the projected cell contact area. The regions with MSA values below the threshold level (MSA < 0.02) were categorized as tight binding zones. The space occupied by a parasite was removed manually before analysis, as shown in Fig. 4A.

**Mesoscopic simulations**

Modeling binding and detachment of infected erythrocytes in shear flow requires two key components: 1) a deformable erythrocyte model and 2) hydrodynamics. We employed multiparticle collision dynamics for hydrodynamics (29,30), combined with a widely used deformable erythrocyte model (31).

The solvent comprises \(N\) point particles with mass \(m\), sorted in a cubic grid with side length \(a\). The dynamics of solvent particles evolve in two steps: 1) streaming and 2) collision. For the streaming step, we have

\[
r_t(r + \Delta t_{\text{cd}}) = r_t(r) + v_t(t) \Delta t_{\text{cd}},
\]

where \(\Delta t_{\text{cd}}\) is the collision time step and \(v_t(t)\) is the \(t\)th particle velocity. The collision step that allows for momentum exchange among the solvent particles is

\[
v_t(t + \Delta t_{\text{cd}}) = v_{\text{cm}}(t) + \mathbb{R}(\alpha)(v_t(t) - v_{\text{cm}}(t)),
\]

where \(v_{\text{cm}}(t)\) is the center of mass velocity of the grid cell to which particle \(i\) belongs and \(\mathbb{R}(\alpha)\) is the stochastic rotation matrix. It is generated for each grid cell by choosing a random axis and a fixed angle \(\alpha\). Solvent density \(\rho\), collision time step \(\Delta t_{\text{cd}}\), and the rotation angle \(\alpha\) determine the shear viscosity of the solvent (32).

The deformable red blood cell is modeled as two-dimensional triangular meshwork of springs with \(N_e\) edges and \(N_t\) triangles. The total potential energy is

\[
V(\{x_i\}) = V_{\text{in-plane}} + V_{\text{bend}} + V_{\text{area}} + V_{\text{volume}}.
\]

The first term is the in-plane elastic energy of the network:

\[
V_{\text{in-plane}} = \sum_{j=1}^{N_e} k_g T_{\text{in}} \left( \frac{3x_j^2 - 2x_j^3}{4p(1 - x_j)} \right) + \frac{k_p}{l_j},
\]

where the first term is the attractive worm-like chain potential and the second is the short-ranged repulsive potential, which, together, results into a spring-like potential. In an attractive potential, \(p\) is the persistence length, \(l_m\) is the maximal extension of the spring, and \(x_j = l_j/l_m\).
The second term is the bending energy of the triangular mesh:

$$V_{\text{bend}} = \sum_{j=1}^{N_b} k_b \left( 1 - \cos(\theta_j - \theta_0) \right),$$

and

$$V_{\text{area}} = \frac{k_a (A - A_0)^2}{2A_0} + \sum_{j=1}^{N_q} k_a \left( \frac{A_j - A_0^j}{2A_0^j} \right)^2$$

$$V_{\text{volume}} = \frac{k_v (V - V_0)^2}{2V_0},$$

where $k_b$, $\theta_j$, and $\theta_0$ are the bending modulus, the angle between the two faces sharing the same edge, and the preferred angle, respectively.

The last two terms in Eq. 3 correspond to surface area and volume constraints:

$$V_{\text{area}} = \frac{k_a (A - A_0)^2}{2A_0} + \sum_{j=1}^{N_q} k_a \left( \frac{A_j - A_0^j}{2A_0^j} \right)^2$$

and

$$V_{\text{volume}} = \frac{k_v (V - V_0)^2}{2V_0},$$

where $k_a$ and $k_q$ are global area and local area constrain coefficients and $k_v$ is the volume constraint coefficient. All the three coefficients were chosen such that the membrane is nearly incompressible and that the total surface and volume fluctuate within 1% throughout the simulation. For a perfectly hexagonal network, an in-plane elastic modulus, such as a shear or Young’s modulus, depends on the choice of model parameters. The shape of the infected cell is chosen to be a discocyte because the trophozoite stage cell is known to have almost retained its initial discocyte shape. The drastic changes in shape start to occur 24–30 h postinvasion (33). The mass of the membrane vertex is set to $\mu m$. The size of an erythrocyte is chosen to be $D_0 = \sqrt{A/\pi}$. The mean edge length is close to 0.5 $\mu m$. The coupling between solvent particles and membrane vertices is introduced via two ways: 1) membrane vertices are included in the collision scheme and 2) solvent particles bounce back at the triangular plaquettes, separating the inside and the outside fluid. The solvent particle density is chosen to be $\rho = 12.5$ and $d_{H_2O} = 0.01 \sqrt{m^2/k_BT}$, which results in the viscosity $\eta = 104\sqrt{m^2/k_BT}$. For a typical shear rate of $\gamma = 0.1$, the Reynolds number becomes $Re = R^2 \rho \eta / \mu < 0.1$. Model units to physical units can be converted by equating the relevant length, force, and timescales (31). For example, the erythrocyte diameter in simulations sets the length scale, i.e., 7.8 $\mu m$ is equivalent to 6.5 $\mu m$, which makes $a = 0.82 \mu m$. Simulation time is approximated with physical time by matching the relaxation time, $\tau = \pi a^2 / \kappa_{\text{sw}}$. The shear modulus of both iHbAA and iHbAS cells is set to 25 $\mu N/m$, taking the value measured by optical tweezers (34). The bending modulus for iHbAA cells is set to $4.1 \times 10^{-19} J$, and for iHbAS, it is $5.6 \times 10^{-19} J$, as determined by flicker spectroscopy (6).

Bond dynamics are employed between the membrane vertices (which also act as receptors) and the ligands that are uniformly distributed on the substrate. The choice of knobs is done using a random procedure which ensures that each trajectory is made with a different knob configuration. At each integration time step $dt_{\text{int}}$, possible bond associations and bond dissociations are performed among vacant receptors and vacant ligands. For bond association, the receptor forms a bond with an available ligand if the distance is less than a critical distance $r_0 = 0.375 \mu m$ with a constant on rate $k_{\text{on}}$. The off rate is chosen in the range of 10–1000 s$^{-1}$. For bond dissociation, we have employed Bell’s model (35), in which the off rate depends on the force $k_{\text{off}} = \kappa_{\text{off}} e^{-\frac{F_0}{|F|}}$, where $F_0$ is the internal bond force scale, which is close to 10 pN for the PITEP-1:CD36 bond (36). The unstrained off rate is chosen in the range of 1–10 s$^{-1}$. We modeled the receptor-ligand bond with a cable force model (37,38): $F(l) = kg(l - l_0)$ when $l$ is greater than $l_0$, else zero, where $k_0 = 2 \mu N/m$ and $l_0 = 0.15a$. At each step, the probabilities for bond association and bond dissociation are computed to be $P_{\text{on}} = 1 - \exp(-k_{\text{off}}\Delta t_{\text{int}})$ and $P_{\text{off}} = 1 - \exp(-k_{\text{off}}\Delta t_{\text{int}})$, respectively, and the unstrained off rate is chosen to be 10 s$^{-1}$.

For measuring binding events, we employed Stokesian dynamics of spherical particle in shear flow to avoid extensive computational complexity. The details of the algorithm are laid out in (37,38). Here, we avoid using bond dynamics as we assume that when the cell reaches close to the substrate (less than 50 nm), it would bind to the substrate, which is a fair assumption because the density of ligands on the substrate is very high.

**Computer simulations setup**

All flow simulations are performed in a three-dimensional rectangular box of the size $36a \times 20a \times 20a$. Periodic boundary conditions are employed along the $x$ and $y$ directions, whereas hard wall boundary conditions are employed in the $z$ direction. Solvent particles make bounce-back collisions with hard walls, but this is not sufficient to account for the no-slip boundary condition at the walls, which is resolved by adding ghost particles in grid cells that are cut by the walls (39). Shear flow $\gamma = \gamma_{\text{sw}}$ is employed along the $z$ direction. Shear flow is generated by adding an offset velocity $v_{\text{sw}}$ to the ghost particles that reside in the grid cells that are cut by the wall at $z = L_z$, which generates shear flow with shear rate $\gamma = v_{\text{sw}}/L_z$. Before starting the flow, we place the cell close to the substrate and let it adhere until it reaches the steady state, and then after, we initiate the flow.

**Statistics**

The presented data were obtained from three independent experiments using blood from more than two different donors both for iHbAA and iHbAS erythrocytes. If not stated otherwise, the data points and error bars represent mean values and standard deviations, respectively.

**Ethical clearance**

The experiments were approved by the ethics review board of Heidelberg University and Mannheim University. The experiments were performed after obtaining informed consent from all voluntary donors in accordance with the relevant guidelines and regulations.

**RESULTS**

**Detachment of adherent cells depends on receptor density and red blood cell type**

Throughout the study, we used the *P. falciparum* strain IT4 (alias FCR3), which was repeatedly panned over primary human dermal microvascular endothelial cells to enrich for parasite populations adhering to two representative host receptors on the surface of endothelial cells, namely CD36 and ICAM-1, before the experiments. A detailed transcriptional analysis revealed predominant expression of the three PTEMP1 encoding var genes, IT4var13, IT4var25, and IT4var66 (40). These var genes belong to the subtypes UpsB and UpsC and are predicted to encode ICAM-1 and/or CD36 binding PTEMP1 variants (41).

We first determined the adhesion strength of *P. falciparum*-infected erythrocytes at the trophozoite stage (24–32 h post-invasion) on supported membranes (12–14) displaying CD36 or ICAM-1 at different densities (Fig. 1 A), using an ultrasonic pressure wave generated by a picosecond laser pulse (Fig. 1 B; (13,16)). Before the experiments, we confirmed that the nonspecific adhesion of uninfected erythrocytes to membranes displaying CD36/ICAM-1 is negligibly small.
Because the ultrasonic pressure wave interacts with the cells for only ~80 ns but at MPa strength, the cellular response is purely elastic. Because of the noninvasive nature of this technique, cells remain viable during the course of the experiment and are able to readhere after detachment.

Because the ultrasonic pressure exerted on cells depends on the radial distance from the initial focus point $P_{\text{Focus}}$ (Fig. S2), the experimentally obtained fraction of adherent cells remaining on the surface normalized by the number of initially adherent cells $\chi_A$ can be analyzed as a function of the radial distance and, hence, the local pressure $P$. The obtained $\chi_A-P$ relationship can be fitted with an error function:

$$\chi_A(P) = \frac{\chi_0}{2} \left(1 - \text{erf} \left( \frac{P - P^*}{\sqrt{2}\sigma} \right) \right).$$

$\chi_0$ is the initial level, and $P^*$ and $\sigma$ are the critical pressure and its standard deviation corresponding to the transition from the adherent to the detached state.

We found that the critical pressure to detach prebound cells declined with increasing intermolecular receptor distances $<d>$ (Fig. 1, C and D). For instance, the $P^*$-values (indicated by arrows in Fig. 1, C–F) decreased more than twofold, from 5.6 to 2.2 MPa, as the intermolecular CD36 distance increased from 11 to 34 nm. Similarly, $P^*$ declined from 3.6 to 2.1 MPa as the intermolecular distance of ICAM-1 increased from 11 to 34 nm. Infected HbAS erythrocytes revealed a detachment behavior on CD36- and ICAM-1-functionalized surfaces qualitatively similar to that of parasitized HbAA red blood cells, albeit the curves were shifted to lower pressure values (Fig. 1, E and F; see also Fig. S3, A and B for an extended $<d>$ range), which was also suggested by previous reports (8,10,40). We further noted that both iHbAA and iHbAS exhibited sharper transitions from the adherent to the detached state on CD36 compared with ICAM-1.

Dynamic binding of infected erythrocytes under shear stress

We next investigated the dynamic aspects of adhesion under physiological hydrodynamic shear stress, using a microfluidic device (Fig. 2 A). The number of bound cells $N_B$ was determined for each condition and normalized by unit area ($\text{mm}^2$) and volume of applied blood suspension (milliliters) to allow for the direct comparison of data collected at different volume fluxes.

Fig. 2 B shows the number of bound cells $N_B \ [\text{mL}^{-1} \text{mm}^{-2}]$ for iHbAA and iHbAS erythrocytes to surfaces functionalized with CD36 at $<d>_{\text{CD36}} = 11 \text{ nm}$, plotted as a function of shear stress $\tau$. The number of binding events gradually decreased with increasing shear stress. For example, the binding of infected HbAA erythrocytes decreased by one order of magnitude from $\Delta N_{\text{iHbAA/CD36}} = 185$ to 15 $\text{mL}^{-1} \text{mm}^{-2}$ as the shear stress increased from $\tau = 0.03$ to 0.10 Pa. Similar to our results with the pressure wave assay, infected HbAS erythrocytes exhibited the same tendency, albeit at an overall lower binding capacity. For example, the number of binding events at $\tau = 0.03 \text{ Pa}$, $\Delta N_{\text{iHbAS/CD36}} = 117 \text{ mL}^{-1} \text{mm}^{-2}$, was reduced by ~30% as compared with parasitized wild-type red erythrocytes.
HbAA and HbAS erythrocytes were more pronounced in the case of ICAM-1 than of CD36 ($\Delta N_{\text{ICAM-1}} = 79 \text{ mL}^{-1} \text{ mm}^{-2}$ obtained at $\tau = 0.03 \text{ Pa}$).

To assess the effect of receptor density on cytoadhesion, we performed flow experiments under systematic variation of $<d>$ from 6 to 67 nm (Fig. 2, D and E). In the case of CD36 (Fig. 2 D), the binding of infected HbAA erythrocytes monotonically decreased with increasing $\tau$ and increasing $<d>$ but did not reach zero at the largest interligand distance $<d> = 67 \text{ nm}$ and $\tau = 0.03 \text{ Pa}$, possibly because of lateral diffusion and accumulation of receptors on the surface of supported membranes retaining their intrinsic lateral fluidity. In the case of ICAM-1 (Fig. 2 E), the number of bound cells first rose to a maximum as $<d>$ increased before it fell to a basal value. This phenomenon was particularly evident at lower wall shear stresses. In the case of infected HbAS erythrocytes, the number of cells bound to CD36 (Fig. 2 F) and ICAM-1 (Fig. 2 G) was distinctly lower compared with wild-type HbAA erythrocytes. Uninfected red blood cells did not bind to the supported membranes at any of the conditions tested, excluding the possibility of nonspecific cell adhesion (Fig. S1).

**Persistence of cytoadhesion under shear**

We next investigated the cytoadhesion strength of prebound cells under increasing wall shear stress. To this end, infected erythrocytes at a hematocrit of $10^6$ cells mL$^{-1}$ were flushed over CD36 or ICAM-1 functionalized membranes for 15 min at a low wall shear stress ($\tau = 0.03 \text{ Pa}$). The shear stress was then increased from 0.05 to 4.0 Pa in a stepwise manner, covering the physiological range (Fig. 3 A; (42)). Fig. 3, B–E depict the surface density of infected HbAA erythrocytes $N_R$ [mm$^{-2}$] remaining attached to CD36 and ICAM-1, respectively, as a function of the shear stress $\tau$. Sigmoidal relationships emerged for both intermolecular distances investigated ($<d> = 11$ and 34 nm) and for both parasitized HbAA and HbAS red blood cells. However, there were consistently fewer cells attached at an intermolecular distance of 34 nm than at 11 nm, and infected HbAS erythrocytes displayed two- to threefold lower $N_R$-values under all experimental conditions. It is worth pointing out that the majority of remaining cells could resist hydrodynamic pressures of up to 0.5 Pa, suggesting that cytoadhesion is persistent once bond formation has been established.

To compare the sustainability of cytoadhesion in flow quantitatively, each data set was fitted with an error function (solid lines in Fig. 3, B–E),

$$N_R(\tau) = \frac{N_R^0}{2} \left( 1 - \text{erf} \left( \frac{\tau - \tau^*}{\sqrt{2}\sigma} \right) \right) + 1,$$

yielding the critical shear stress for the cell detachment $\tau^*$. On surfaces displaying CD36 at $<d> = 11 \text{ nm}$ (Fig. 3, B
Adhesion of malaria-infected erythrocyte

Changes in cell shape and cell-substrate contact by shear

Cells exposed to shear stresses undergo viscoelastic deformation and adapt their adhesion contacts (43). These cellular factors might be equally important as molecular ones, especially when considering the persistence of cytoadhesion after the establishment of initial adhesion. Extending our ensemble observation of cell adhesion and detachment, we monitored changes in the cell-surface contacts caused by shear stresses ($\tau = 0.05$–1.0 Pa), using RICM. As the intensity of light detected at each pixel reflects the separation distance between the cell membrane and the substrate, we categorized two regions as follows (Fig. 4): 1) the adhesion contact area as defined by calculating the gradient of the time average of $100 \times$ RICM images (regions surrounded by purple solid lines) and 2) the tight binding zone as identified by thresholding the MSA of intensity and distance fluctuation at MSA < 0.02 (regions shaded in dark gray).

The left row of Fig. 4 $A$ depicts the RICM images (left) and the extracted cell-surface contacts (right) of infected HbAA erythrocytes on a CD36 surface ($<d>=11$ nm) at $\tau = 0.10, 0.25,$ and 1.0 Pa. The red arrow indicates the direction of flow. The adhesion contact area increased with rising shear stress. At $\tau = 0.10$ Pa, the adhesion contact area resembled a “U shape,” with no adhesion contact on the side facing the flow. When the shear stress was increased to $\tau = 0.25$ Pa, the opening closed, and the adhesion contact zone adopted an “O shape.” Concomitant with the changes in contact area, the cells altered their morphology as $\tau$ increased. At low $\tau$, the cells sustained discocyte-like shapes. No adhesion contact was established in the middle of the cell. Once the shear stress reached $\tau = 1.0$ Pa, the cell and the adhesion contact area took on a “raindrop shape,” indicating that the tip of the cell facing the flow was pinned. We note that these shape transformations were driven by shear flow.

Infected HbAS erythrocytes exhibited qualitatively similar changes in response to increasing shear stress. To
highlight the difference between infected HbAA and HbAS erythrocytes, the contact area and the tight binding zone were plotted as a function of the shear stress (Fig. 4 B). The contact area of infected HbAA erythrocytes increased from 11 to 25 $\mu m^2$ as the shear stress rose from $\tau = 0.05-1.00$ Pa, respectively. The contact area of infected HbAS erythrocytes initially exhibited a similar increase up to 17 $\mu m^2$, but there was no further increase at $\tau > 0.25$ Pa. It should be noted that changes in the binding zones of both infected HbAA and HbAS erythrocytes were less dependent on shear stress, suggesting that the unbinding from CD36 bond is not shear dependent. This finding agrees with the data presented in Fig. 3 F.

As presented in Fig. 4 C, the adhesion contact area of an infected HbAA erythrocyte on ICAM-1 surfaces also underwent the shape transition from a U through an O to a raindrop shape, and concomitantly, the adhesion contact area grew from 11 to 25 $\mu m^2$ (Fig. 4 D) with increasing shear stress from 0.05 to 1.00 Pa. This finding is comparable to that made on CD36 surfaces (Fig. 4 B), with the exception that the ICAM-1 tight binding zone exhibited a remarkable increase from 4 to 12 $\mu m^2$ with increasing shear stress, which was not seen to the same extent on CD36 surfaces. A statistical analysis confirmed that the sizes of the tight binding zones were statistically different between ICAM-1 and CD36 surfaces at 1.00 Pa (12 vs. 5.6 $\mu m^2$, respectively; $\chi^2$ test, $p = 0.01$), although the starting values at 0.05 Pa were comparable. As shown in Fig. 4 D, infected HbAS erythrocytes exhibited a qualitatively similar shear-induced adhesion behavior. The contact area and binding zone quickly increased after a rise in shear stress from 0.05 to 0.10 Pa. At $\tau \geq 0.25$ Pa, the gains in adhesion became...
Assessing shear-induced cell deformations by mesoscopic computer simulations

Recently, mesoscopic computer simulations have been established as an instructive method to interpret results from quantitative flow chamber experiments (37,40). Using triangulated surfaces for cell shape and multiparticle collision dynamics for hydrodynamic flow, one can simulate how adherent cells deform and move on a substrate (Fig. 5 A). However, these kinds of simulations are very time consuming. If shape changes are not the main focus, one can use Stokesian dynamics simulations of spherical particles in shear flow to get better statistics. Here, we used both approaches to simulate the effect of shear flow and receptor distance on the cytoadhesion of iHbAA and iHbAS cells. We first used the deformable cell model. To reflect the naturally occurring variability in knob density, we have used a range within which we varied the knob density instead of fixing it. We used 6–8 knobs \( \mu m^{-2} \) for iHbAA and 3–5 knobs \( \mu m^{-2} \) for iHbAS (40). For each realization, we chose the number of active vertices randomly within a range that corresponds to the respective knob density range. In Fig. 5, B and C, we show the probabilities of the cells to adhere against the shear stress \( \tau \) as a function of ligand distance and hemoglobin variant, respectively. Each point for a given shear stress is an average of 10 different simulations performed, and for each realization, the knob density is chosen randomly. The probability is simply given by the ratio of number of realizations when the cell survived to the total number of realizations. The critical shear stress for the binding-unbinding transition determined with the error function is 0.22 and 0.15 Pa for iHbAA and iHbAS, respectively. The values are reasonably close to the experimental values (see Fig. 3, F and G).

To estimate the binding events and its dependence on shear stress, we used Stokesian dynamics simulations. For each shear stress, we run multiple simulations with different initial heights (up to 25 \( \mu m \)), and we counted how many cells fall close to 50 nm to the substrate without exceeding 5 mm along the flow direction. This simulation setup is designed to imitate the binding events measurements in the flow chamber experiments described above. The results are not sensitive to the cutoff distances. In Fig. 5 D, we show the number of cells that succeed in making contact with the substrate for varying shear stress. The dependence agrees well with Fig. 2 A. Because no bond dynamics is employed, we captured the phenomenon qualitatively, but not quantitatively. We conclude that the difference between binding events of iHbAA and iHbAS in experiments is mainly due to differences in knob density.

Our computer simulations with deformable cells can also shed light on the contact zone with the substrate and its dynamic changes under the shear. Fig. 6 A represents the side view (upper panels) and the bottom view (lower panels) of an infected erythrocyte undergoing a shear-induced deformation simulated by mesoscopic computer simulations. The gray scale represents the absolute mean curvature. The adhesion contacts, surrounded by solid black lines in the lower panels, exhibit a transition from a U shape to an O shape, which reproduces the experimental results (Fig. 4, A and C). In Fig. 6, B and C, we show the change in the adhesion contact area for both iHbAA and iHbAS cells. iHbAA cells show a monotonic increase, and iHbAS cells are less prominent than iHbAA, which is due to relatively small knob density. These results agree well with
the experiments (Fig. 4, B and D). It should be noted that the change in area given in μm² is only a semiquantitative indicator because the adhesion contact area strongly depends on how the adhesion contact is defined. In simulations, we measured the contact area as the membrane surface that is bonded with the substrate. Thus, this would be equivalent to the binding area in experimental measurements.

At the highest shear, the adhesion contact became homogeneous. We noted that the experimentally observed sharp tip of the raindrop shapes cannot be reproduced with homogeneous coverage of knobs. We speculate that such sharp tips are the result of an accumulation of receptors moving in the supported membranes, resulting in a few strong adhesion contacts that hold the entire cell. When the supported membrane consists of lipids in fluid phase, the lipids, as well as recombinant proteins coupled to anchor lipids, undergo free lateral diffusion (46). In our previous account, we showed that the kinetics of cancer cell apoptosis driven by the ligand-receptor pairs becomes distinctly slower when the recombinant proteins were anchored on a membrane in a gel phase (15). To recapitulate the effect caused by the accumulation of receptors, we modeled the cell with few strong adhesions and many weak adhesions. For strong adhesions, more than one receptor is used per knob, and the bond strength is increased by a factor of 10. In Fig. 6 D, we show cell snapshots (both top view and bottom view) for a cell with one strong adhesion along with few hundreds of weak adhesions. The sharp tip appeared on the front side of the cell. It resembles the cell snapshots from Fig. 4, A and C at higher shear stress. We also simulated cells with a higher number of strong adhesions, resulting in conformations with multiple sharp edges. These simulations were performed at shear stress τ = 0.5 Pa, and on and off rates were chosen such that the cells do not detach from the substrate. Some of the features of the raindrop shape could also be seen in recent simulations when the cell has very few adhesion contacts (47).

To further verify our hypothesis, we compared the shape of adhesion contacts on 1,2-palmitoyl-sn-glycero-3-phosphocholine (DPPC) membranes in gel phase, displaying CD36 and ICAM-1 at <d> = 11 nm. As presented in Fig. S5, the adhesion contact did not take a raindrop shape even at the highest shear (1.00 Pa). This result indicates that the fluidity of supported membranes, which shares a common feature with the plasma membrane of endothelial cells, allows for the lateral accumulation of ligand-receptor pairs, resulting in the strong pinning near the tip of the cell facing to the flow. The capability of supported membranes to control both inter-receptor distance and membrane fluidity helped us unravel how the individual molecular interactions manifest themselves on the cellular level.

![FIGURE 6](image-url)

(A) Side (upper row) and bottom (lower rows) views of an infected erythrocyte simulated with the deformable cell model under different shear conditions. The regions surrounded by solid black lines coincide with adhesion contacts. (B) Change in contact area under varying shear stress for iHbAA cells; the same is displayed for (C) iHbAS cells. For both (B) and (C), the interligand distance is <d> = 80 nm. (D) Snapshots (left: top view, right: bottom view) of adherent iHbAA cells under high shear stress flows. The shear flow direction is from left to right. The circle on the bottom viewed snapshot indicates the position of strongly adhered knob. The strong adhesion contact or knob is modeled with higher spring constant and multiple receptors rather than one receptor per knob as in default case. The shear stress is τ = 0.5 Pa. To see this figure in color, go online.
Our study suggests that receptor-functionalized supported membranes are a valuable complementary tool to study the processes underpinning cytoadherence of *P. falciparum*-infected erythrocytes. Unlike microvascular endothelial cells, which can present multiple cytoadhesion receptors at variable density and distribution on their surface (48–51) in a manner affected by the activation state (52,53), supported membranes offer the option of controlling the receptor identity, orientation, and density. Thus, cytoadhesion studies can be performed under controlled conditions, providing additional insights into the dynamics and kinetics of cytoadherence, which are not easily acquirable by using microvascular endothelial cells or receptor-coated petri dishes.

Previous studies using microvascular endothelial cells have shown that the number of adhering infected erythrocytes and their adhesion strength increase with the receptor density (48–51). Our study now provides a quantitative understanding of this observation by revealing that adhesion parameters declined with increasing receptor distance over a range of 10–100 nm for both CD36 and ICAM-1 at all wall shear stresses investigated. The red blood cell type, the knob density, and the amount and type of PfEMP1 variant presented on the cell surface are variables that can affect this relationship (see below for further discussion).

According to Sanchez et al., each knob carries on average three to four PfEMP1 molecules placed 18–14 nm apart from one another at the tip of the knob, as shown for the PfEMP1 variant VAR2CSA that confers binding of infected erythrocytes to placental CSA (5). Interestingly, the receptor distance supporting optimal cytoadhesion seems to be in the same nanometer range. Whether the spatial arrangement of PfEMP1 molecules on knobs has evolved to match the spacing of receptor molecules and hence to support efficient cytoadherence or whether these are two coincidental events will require further investigation. Further studies would also be needed to investigate whether the positioning of VAR2-CSA on knobs is a general property of other PfEMP1 variants.

Supported membranes further allowed us to assess dynamics in contact footprint using RICM. In flow chamber experiments, infected erythrocytes respond to external shear stresses by adapting their shape before the hydrodynamic pressure peels them off from the surface once the wall shear stress exceeds a critical value. We now observed that the shape transition and the gain in contact area depend on the knob density and the intermolecular receptor distance, and it is enforced by the accumulation of mobile ligand-receptor pairs caused by a vertical pulling force, as suggested by a previous study using artificial model systems and theoretical models (54). By comparing the adhesion contacts on membranes in fluid phase (Fig. 4) and in gel phase (Fig. 55), we experimentally confirmed that the shear-induced pinning near the tip of the cell facing to the flow originates from the lateral accumulation of receptor molecules. Supported membrane systems offer a unique advantage over substrates displaying nanopatterns of receptor molecules because they are capable of fine adjusting not only inter-receptor distance but also membrane fluidity. In this context, it is further worth mentioning that cell-cell adhesion assays support the view of receptor accumulation at contact zones. For example, the engagement of CD36 with infected erythrocytes results in CD36 clustering via a process that involves actin cytoskeleton remodeling and dephosphorylation of Thr92 in the ectodomain of CD36 (52). Similarly, ICAM-1 molecules can also cluster and are recruited into lipid rafts upon interaction with a ligand (44), in addition to ICAM-1 being upregulated (10- to 20-fold on human brain microvascular endothelial cells and ~1000-fold on human umbilical vein endothelial cells) in response to inflammatory mediators (49). Other studies have shown that the lifetime of the contact area (containing clusters of ligand-receptor pairs) might be several orders of magnitude higher than the half-life of the single bond dissociation rate (55,56), as dissociated molecules can easily find new interaction partners particularly on substrates with high receptor densities (45). On the basis of these findings, we propose that the same processes as described above occur in vivo and that the shear-induced increases in contact area and tight binding zone contribute to the fact that cytoadhering infected erythrocyte can withstand hydrodynamic pressures of 1.0 Pa and more without peeling off.

The increase in contact area and tight binding zone with increasing flow seems counterintuitive. However, our mesoscopic computer simulations explain this behavior by the specific cellular and biomechanical properties of the infected erythrocyte, including membrane mechanics and knob density (6,40), and the lateral recruitment and accumulation of receptors at sites of contact. In other words, the shape transition and the gain in contact area under conditions of increasing wall shear stress are ascribed to the ability of infected erythrocytes to respond to hydrodynamic pressure with changes in cell shape, which, in turn, increases the cell-surface area that can interact with the support. The ability to engage in binding events then depends on the knob density and the intermolecular receptor distance, and it is enforced by the accumulation of mobile ligand-receptor pairs caused by a vertical pulling force, as suggested by a previous study using artificial model systems and theoretical models (54). By comparing the adhesion contacts on membranes in fluid phase (Fig. 4) and in gel phase (Fig. 55), we experimentally confirmed that the shear-induced pinning near the tip of the cell facing to the flow originates from the lateral accumulation of receptor molecules. Supported membrane systems offer a unique advantage over substrates displaying nanopatterns of receptor molecules because they are capable of fine adjusting not only inter-receptor distance but also membrane fluidity. In this context, it is further worth mentioning that cell-cell adhesion assays support the view of receptor accumulation at contact zones. For example, the engagement of CD36 with infected erythrocytes results in CD36 clustering via a process that involves actin cytoskeleton remodeling and dephosphorylation of Thr92 in the ectodomain of CD36 (52). Similarly, ICAM-1 molecules can also cluster and are recruited into lipid rafts upon interaction with a ligand (44), in addition to ICAM-1 being upregulated (10- to 20-fold on human brain microvascular endothelial cells and ~1000-fold on human umbilical vein endothelial cells) in response to inflammatory mediators (49). Other studies have shown that the lifetime of the contact area (containing clusters of ligand-receptor pairs) might be several orders of magnitude higher than the half-life of the single bond dissociation rate (55,56), as dissociated molecules can easily find new interaction partners particularly on substrates with high receptor densities (45). On the basis of these findings, we propose that the same processes as described above occur in vivo and that the shear-induced increases in contact area and tight binding zone contribute to the fact that cytoadhering infected erythrocyte can withstand hydrodynamic pressures of 1.0 Pa and more without peeling off.
adhesion bonds are broken simultaneously. Effects occurring under physiological wall shear stress conditions, such as morphological changes due to remodeling of the cytoskeletons (57) and rate-dependent bond rupture forces (56), can therefore be ignored. Thus, the ultrasonic pressure wave assay depicts a more realistic view of the intrinsic adhesion strengths than do standard wash-off experiments using wall shear stresses in the physiological range. Note that the bond rupture force for a receptor-ligand pair depends exponentially on the loading rate (56), which explains the very high cell detachment pressures observed in the ultrasonic pressure wave experiments.

We acknowledge that we cannot attribute the adhesion phenomena described in this study to specific PfEMP1 variants, owing to the fact that the parasite population is heterogeneous with regard to the var genes expressed. The generalizability of our findings is also constrained by the range of PfEMP1 variants tested in our infected erythrocyte populations. It is, therefore, speculative to conclude that the PfEMP1-CD36 and the PfEMP1/ICAM-1 interactions behave like slip bonds, although the monotonic decrease in binding events at each <d> with increasing shear stress would be consistent with a slip bond. A catch-slip bond behavior has recently been proposed for the PfEMP1/ICAM-1 interaction, based on single-cell pull experiments conducted using an atomic force microscope (58,59). This observation was made at a very low loading rate, and it is possible that there exists, in the low shear stress regime, a threshold beyond which a catch-slip bond becomes indistinguishable from a slip bond (55). In comparison, the interaction between the PfEMP1 variant VAR2CSA and placental CSA has been described as a cooperative, shear-enhanced catch bond-like behavior (13). It would be interesting to use our system to measure binding characteristics of other PfEMP1-ligand interactions.

Throughout our study, we noted the distinct cytoadhesion characteristics of infected HbAS erythrocytes. Compared with age-matched infected HbAA erythrocytes, the infected sickle cell trait HbAS erythrocytes exhibited two- to fourfold lower adhesion efficiencies on both CD36 and ICAM-1 surfaces at wall shear stresses below 0.05 Pa. At wall shear stresses exceeding 0.05 Pa, the adhesion efficiencies of infected HbAS and HbAA erythrocytes converged. These findings are in line with previous studies showing reduced cytoadhesion of infected HbAS erythrocytes to microvascular endothelial cells under hydrodynamic conditions prevalent in microvessels (8,9,40). The convergence of the adhesion curves is explained by the fact that only 35–45% of total hemoglobin in sickle cell carriers is hemoglobin S (60). It is, therefore, conceivable that blood from sickle cell carriers contains a fraction of wild-type HbAA erythrocytes and that these cells, when infected, adhere under higher wall shear stresses. Along these lines, Cholera et al. reported that a small fraction of infected erythrocytes from patients with the sickle cell trait display knob sizes and knob density, as well as similar amounts of PfEMP1 on the surface, comparable with parasitized wild-type erythrocytes (8).

The reduced adhesion capability of infected HbAS erythrocytes is also reflected by a 30% smaller contact zones and significantly lower pressures to detach prebound cells. As suggested by both experiments and computer simulations (9,33,40), one of the determinants for the weaker cytoadhesion capacity of infected HbAS erythrocytes is their aberrant knob size and density (5,8,40). Infected HbAS erythrocytes have fewer knobs ($\sigma_{iHbAS} = 3 \pm 2$ knobs $\mu m^{-2}$) compared with parasitized HbAA erythrocytes ($\sigma_{iHbAA} = 14 \pm 4$ knobs $\mu m^{-2}$). Moreover, the diameter of knobs expressed on iHbAS, $\phi_{iHbAS} \approx 191 \pm 99$ nm, are larger than those on iHbAA, $\phi_{iHbAA} \approx 72 \pm 21$ nm (5,40). Taking these values, one can calculate a dimensionless knob size $\epsilon = 2\pi(\phi/2)^2\sigma$. As expected, this value is higher for infected HbAS erythrocytes ($\epsilon_{iHbAS} \sim 0.16$) compared with infected HbAA erythrocytes ($\epsilon_{iHbAA} \sim 0.12$). Yet, despite an ~33% smaller dimensionless knob size, the number of PfEMP1 molecules per knob is comparable between both red blood cell types (5), indicating some deeper molecular aberrations, such as impaired export of PfEMP1 molecules and other parasite-encoded factors into the host cell compartment (61), impaired anchoring of PfEMP1 molecules to the membrane skeleton of the host cell via knobs (8,10,62) and/or defects in parasite metabolic activity (63).

In summary, our study supports the use of receptor-functionalized membranes as a complementary tool to study quantitative aspects of cytoadhesion, such as the effect of the receptor distance and mobility on adhesion efficiency. The supported membranes also allowed us to assess the intrinsic binding strengths and to inform on cell-adhesive contact zones and their dynamics under shear stress.

**SUPPORTING MATERIAL**

Supporting material can be found online at https://doi.org/10.1016/j.bpj.2021.07.003.

**AUTHOR CONTRIBUTIONS**

M.T., U.S.S., and M.L. designed the research. B.F., C.L., J.C., C.P.S., and M.C. performed the experiments. B.F., A.Y., M.L., and M.T. analyzed the data. A.K.D. performed the computer simulations with help from U.S.S.; M.L., U.S.S., B.F., and M.T. wrote the manuscript. All authors participated in discussion and manuscript editing.

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SUPPORTING CITATIONS

Reference (64) can be found in the Supporting material.

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