Membrane fluctuations and acidosis regulate cooperative binding of ‘marker of self’ protein CD47 with the macrophage checkpoint receptor SIRPα

Jan Steinkühler1,2, Bartosz Różycki3, Cory Alvey1, Reinhard Lipowsky2, Thomas R. Weikl2, Rumiana Dimova2 and Dennis E. Discher1,*

ABSTRACT
Cell-cell interactions that result from membrane proteins binding weakly in trans can cause accumulations in cis that suggest cooperativity and thereby an acute sensitivity to environmental factors. The ubiquitous ‘marker of self’ protein CD47 binds weakly to SIRPα on macrophages, which leads to accumulation of SIRPα (also known as SHPS-1, CD172A and SIRPA) at phagocytic synapses and ultimately to inhibition of engulfment of ‘self’ cells – including cancer cells. We reconstituted this macrophage checkpoint with GFP-tagged CD47 on giant vesicles generated from plasma membranes and then imaged vesicles adhering to SIRPα immobilized on a surface. CD47 diffusion is impeded near the surface, and the binding-unbinding events reveal cooperative interactions as a concentration-dependent two-dimensional affinity. Membrane fluctuations out-of-plane link cooperativity to membrane flexibility with suppressed fluctuations in the vicinity of bound complexes. Slight acidity (pH 6) stiffens membranes, diminishes cooperative interactions and also reduces ‘self’ signaling of cancer cells in phagocytosis. Sensitivity of cell-cell interactions to microenvironmental factors – such as the acidity of tumors and other diseased or inflamed sites – can thus arise from the collective cooperative properties of flexible membranes.

This article has an associated First Person interview with the first author of the paper.

KEY WORDS: Adhesion, Vesicles, Elasticity, Cooperative binding, GPMV

INTRODUCTION
Cells use adhesion molecules to interact with and thereby identify other cells, but recent theoretical and simulation results indicate that the physical act of binding is highly cooperative and mediated by membrane fluctuations in a generalized law of mass action (Krobath et al., 2009). Differences in 3D versus 2D binding have been studied with cells adhering to supported lipid bilayers (Zhu et al., 2012; Tolentino et al., 2008) and with lipid vesicles bearing reconstituted binding molecules (Fenz and Sengupta, 2012; Bihr et al., 2014, Chan et al., 2007), but cell experiments are complicated by cytoskeleton-driven remodeling (Huppa et al., 2010) and liposomes lack the molecular richness of cell membranes. Here, we use cell-derived giant vesicles that lack cytoskeletal interactions (Sezgin et al., 2012), and we focus on the universally expressed protein CD47 in membranes as it mediates adhesion to SIRPα (also known as SHPS-1, CD172A and SIRPA) on a rigid surface. Wide-ranging (bio)physical experiments provide evidence of equilibration and are linked closely to multiscale simulation.

CD47 binding to SIRPα on macrophages ultimately inhibits engulfment of ‘self’ cells by macrophages (Oldenborg et al., 2000) and is now being pursued clinically in cancer therapy by antibody blockade of this ‘macrophage checkpoint’ (reviewed in Alvey and Discher, 2017). ‘Self’ cells also signal to T-cells (via interactions of PDL1 with PD1) through a phosphatase pathway also downstream of SIRPα, and antibody blockade of the PDL1-PD1 T-cell checkpoint has already proven remarkably effective in the clinic against some cancers. As with the majority of cell-cell signaling molecules, trans interactions of two integral membrane proteins in juxtaposed plasma membranes leads to a dependence of membrane adhesion on membrane conformation. An unfavorable membrane geometry such as occurs with chemically rigidified discocyte-shaped red blood cells, limits CD47 signaling and increases phagocytosis (Sosale et al., 2015a). For distances between membranes that are similar to the sizes of adhesion molecules, the membrane-membrane interaction depends on (membrane) flexibility and fluctuations as well as conformations of binding molecules (Xu et al., 2015). Studies here of a native membrane protein interacting with its immobilized receptor provides some of the first experimental evidence of cooperative binding due to suppression of membrane fluctuations. The findings should apply to immunological synapse complexes and many membrane-bound signaling complexes beyond CD47-SIRPα at the phagocytic synapse.

RESULTS AND DISCUSSION
Cell-derived vesicles retain CD47 and thermally fluctuate
Membrane sheets of several hundred square micrometers in area were detached from the cortical cytoskeletons of CD47-GFP expressing HEK cells (Fig. 1A, left) via a method previously used to study lipid segregation (Sezgin et al., 2012). These sheets spontaneously bud and seal to form giant unilamellar vesicles that are more specifically referred to as giant plasma membrane vesicles (GPMVs). GPMVs possess a near-native composition of lipid and membrane protein (Bauer et al., 2009) and retain lipid and protein mobility and conformation (Baumgart et al., 2007; Sezgin et al., 2012; Veatch et al., 2008) as well as optically resolvable shape fluctuations (Fig. 1B, inset).

Phase-contrast microscopy with μs exposure times shows the amplitudes of Fourier modes q of membrane fluctuations at

Received 12 February 2018; Accepted 9 May 2018
the equatorial plane (Fig. 1B). The $q^{-3}$ dependence agrees with the Helfrich theory of elastic membranes fluctuating at finite temperature, as previously applied to pure lipid vesicles in a fluid state (Gracia et al., 2010; Lipowsky, 1991). The mechanical properties of GPMVs can thus be reduced to a few parameters, such as bending rigidity, tension and preferred curvature, that are independent of detailed molecular composition and the biological state(s) of membranes.

**CD47 is properly oriented in GPMVs and functional for binding**

CD47-GFP makes GPMVs uniformly fluorescent (Fig. 1A, Fig. S1A), and a red fluorescent anti-CD47, which binds to the extracellular domain of CD47, suggests proper orientation. The two signals not only correlated (Fig. 1C) but also confirm a variation of CD47 levels between GPMVs. A range of CD47 concentrations could thus be studied in adhesion experiments.

To assess CD47-SIRPα binding, we physically adsorbed the extracellular domain of SIRPα (fused to a large single domain protein, GST) or the surface-blocking agent bovine serum albumin (BSA) on coverslips (Subramanian et al., 2006). SIRPα homogenously adsorbed based on fluorescence imaging (Fig. S1B). Gravitational settling of GPMVs onto BSA-coated coverslips showed near-spherical vesicle shapes and therefore little interaction (Fig. 1A, top right). On SIRPα-coated coverslips in contrast, GPMVs deformed strongly as they adhered (Fig. 1A, bottom right): the upper unbound cap is spherical whereas the flattened lower membrane segment is where CD47 accumulated in binding the immobilized SIRPα (Fig. 1A). After initial contact between a GPMV and the SIRPα surface, CD47-GFP diffused into the adhesion zone in a ring-like fashion and became homogeneous over time (Fig. 1A).

Experiments were then done on GPMVs with equilibrated and homogenous adhesion, lacking any domain formation and thus implying a single species of CD47-SIRPα adhesion complexes.
(Weikl et al., 2009). Diffusion dynamics in these two segments were measured by fluorescent recovery after bleaching (FRAP) of micrometer-sized spots.

2D binding affinity determined by FRAP
FRAP dynamics in small bleached spots differed between unbound and bound membrane segments. In unbound segments, CD47 fluorescence recovered over seconds and yielded a diffusion constant of $D_{\text{CD}47}=0.12\pm0.02 \, \mu\text{m}^2/\text{s}$ ($n=3$) as determined from fitting to a standard model for unhindered diffusion (Soumpasis, 1983). In stark contrast, CD47 within the adherent zone recovered $\sim100$-times slower (Fig. 1D). CD47-SIRP$\alpha$ unbinding events are likely followed by a short distance of free diffusion before re-binding (Sprague et al., 2004). Such trajectories are governed by a reaction-diffusion model (Sprague et al., 2004; see Material and Methods). By fitting the model to the measured time-intensity traces the 2D binding affinity $[R]K_{2D}=k_{\text{on}}/k_{\text{off}}$ is extracted. Here, $[R]$ is the density of (unbound) SIRP$\alpha$ on the glass surface. The fit to the model gives $[R]K_{2D}$ data for individual vesicles expressing different densities of CD47. The results are independent of the exact value of $[R]$, but $[R] \sim 4000$ molecules$/$$\mu\text{m}^2$ (see Fig. S1B). Strikingly, CD47-SIRP$\alpha$ complex concentration and binding affinity are positively correlated, which is indicative of cooperative binding (Fig. 2A). Computer simulations help to clarify the cooperative effects in CD47-SIRP$\alpha$ binding and unbinding.

Cooperative binding by suppression of membrane fluctuations
Our multiscale modeling approach combines: (i) simulations of a coarse-grained CD47-SIRP$\alpha$ complex (Fig. 2B) to determine the effective spring constant of the complex, and (ii) simulations of large membrane segments adhering via CD47-SIRP$\alpha$ complexes, modeled as elastic springs (Fig. 2C,D). In simulations of a CD47-SIRP$\alpha$ complex, the transmembrane domain of CD47 is embedded in a small membrane patch parallel to the substrate and connected by a short flexible linker to its extracellular domain. This domain is bound to SIRP$\alpha$ in the correct orientation based on the crystal structure of the complex (Protein Databank, PDB-2JJS). SIRP$\alpha$ is connected via a linker to substrate-bound GST in the experiments. Flexible linkers and protein interactions have been successfully applied to several multi-protein complexes (Kim and Hummer, 2008), including ERK2-HePTP (Francis et al., 2011) and ESCRT complexes (Boura et al., 2011, Boura et al., 2012). Variation of the separations between the membrane patch and substrate observed in simulations (Fig. S2) result from the molecular architecture

Fig. 2. Multiscale modeling of experiments to clarify interactions. (A) Binding constant $K_{2D}$ as a function of complex concentration $[RL]$ from experiments (data points) and modeling (lines) for different values $\Delta l$ of repulsive membrane-substrate interactions. Modeling is based on Eqn 1, with fit values $K_{\text{on}}=(956\pm24)/[R]$, $(470\pm10)/[R]$, $(378\pm8)/[R]$ and $(326\pm8)/[R]$ for $\Delta l=1, 3, 4, 5 \, \text{nm}$, respectively. Binding constant $K_{2D}$ has units of concentration $[R]$ of unbound SIRP$\alpha$. Error bars indicate 95% confidence intervals for fitting the reaction-diffusion model±s.e.m.). Each data point corresponds to measurement of an individual GPMV in at least two independent experiments. (B) Simulation snapshots of coarse-grained CD47-SIRP$\alpha$ complex. Separation between fluid membrane patch (cyan) and substrate (grey) varies in simulations mainly because of conformational changes of the unstructured linker that covalently connects extracellular SIRP$\alpha$ domain (blue) to substrate-bound GST (red). (C,D) Snapshots of adhering membranes with area $3\times3 \, \mu\text{m}^2$ for respective concentrations $[RL]=30, 80 \, \mu\text{m}^{-2}$ of CD47-SIRP$\alpha$ complexes at $\Delta l=4 \, \text{nm}$. Black dots indicate complexes. (E) Distributions $P(l)$ for local separation between membrane and substrate obtained from averaging over many membrane snapshots for various $[RL]$. Dashed black line represents $K_{2D}(l)$ (arbitrary units).
and flexibility of the CD47-SIRPα complex. The mean length ($l_0=17.2 \text{ nm}$) of these separations is the preferred membrane-substrate separation of the complex, the standard deviation $\sigma=1.2 \text{ nm}$ reflects flexibility and an effective spring constant $k_0=\kappa_0 T/\sigma^2$ of the complex, where $\kappa_0$ is Boltzmann’s constant and $T$ is temperature ($\sim300 \text{ K}$).

In our simulations of large adhering membrane segments (Fig. 2C,D), membranes are elastic surfaces with bending rigidity $\kappa=10 k_0 T$ as measured experimentally, and discretized into quadratic patches of size $15 \times 15 \text{ nm}^2$ (Weikl and Lipowsky, 2006). Membrane patches that contain CD47-SIRPα complexes are bound to the substrate with separation $l_0$ and spring constant $k_0$. Repulsive interactions between the protein layer on the substrate and the membrane are taken into account by allowing only local separations $l>(l_0-\Delta l)$ between membrane patches and substrate, with $\Delta l=0$. From simulations, we obtain distributions $P(l)$ of local separations that depend on concentration [RL] of CD47-SIRPα complexes (Fig. 2E) because bound complexes constrain membrane shape fluctuations reflected by $P(l)$.

From the distributions $P(l)$, and from the preferred length $l_0$ and spring constant $\kappa$ of the complexes, we obtain the binding constant $K_{2D}$ from Xu et al. (2015):

$$K_{2D} = \int \hat{K}_{2D}(l)P(l)dl,$$

with $\hat{K}_{2D}(l) = K_{\text{max}} \exp[-(l-l_0)^2/\sigma^2]$ up to a pre-factor $K_{\text{max}}$, which we treat as a fit parameter. $K_{2D}$ increases with bond concentration [RL] (Fig. 2A), because distributions $P(l)$ become narrow and shift towards $l_0$ with increasing [RL] (Fig. 2E). Modeling results agree with experimental data points for different values of $\Delta l$ for the repulsion between membrane and protein-coated substrate (Fig. 2A). We vary $\Delta l$ because neither the thickness of protein layers nor the roles(s) of GPMV proteins in steric repulsion are known. Strong increase of $K_{2D}$ with [RL] reflects cooperative binding of CD47-SIRPα, which results from smoothing of membrane fluctuations by bound complexes (Krobath et al., 2009; Hu et al., 2013). Smoothing facilitates formation of additional CD47-SIRPα complexes.

**Slight acidity inhibits CD47-SIRPα cooperativity and favors phagocytosis**

To further validate our model, we explored the effects of membrane fluctuations (and hence cooperativity) in relation to the bending rigidity of the membrane. We discovered that subjecting cells to a slight acidity of pH 6 induces stiffening of the GPMV (Fig. 3A). In both simulations and experiments, such rigidification led to a substantial decrease in cooperativity (Fig. 3B). Simulations and experiments are linked via the measured bending rigidity, which lumps the complex cell (plasma membrane) response to a pH change into a single parameter. Absolute values of $[R]K_{2D}$ data points are also significantly reduced, as confirmed by an independent but more approximate determination of $K_{2D}$ using fluorescent intensities of adherent and free membrane segments (Fig. 3C). Some alternative explanations for the decreased 2D binding affinity can be ruled out by further experiments: first, binding of soluble SIRPα to red blood cells did not exhibit a strong pH dependence in the 3D binding affinity (Fig. S3A). Second, density changes in absorbed protein $[R]$ do not change with pH, perhaps because of a change in conformational and rotational free-energy loss of CD47 and/or SIRPα during binding (Xu et al., 2015). Regardless of the exact cause of the 2- to 4-fold decrease in 2D binding affinity with decreased pH, the loss of cooperativity is likely to be more important for signaling, because CD47 densities vary over one to two orders of magnitude across...
otherwise healthy cells and yet CD47 signaling remains effective (Subramanian et al., 2007). Changes in the collective binding affinity of CD47 for SIRPα could ultimately inhibit a cell’s ability to efficiently signal through this pathway.

Tumor tissues commonly experience acidosis (Tannock and Rotin, 1989). Opsonization of lung cancer-derived cells (A549) with an IgG to present an ‘eat me’ signal to macrophages normally does not efficiently engage phagocytosis because of expression of CD47 on the cancer cell surface (Sosale et al., 2015b). However, opsonization under acidic conditions caused an increase in phagocytosis (Fig. 4A), consistent with the biophysical data, showing diminished cooperativity and binding capacity of CD47 at slightly acidic conditions of pH 6 (Fig. 3B,C). When A549 cancer cells are cultured long term at pH 6, CD47 levels were elevated slightly acidic conditions of pH 6 (Fig. 3B,C). When A549 cancer cells are cultured long term at pH 6, CD47 levels were elevated (Fig. 4B), indicating a cellular signaling cascade that works against reduction in ‘self’ signaling by upregulation of CD47. Together, these data suggest that the measured 2D binding affinities reveal physiologically relevant parameters that would have been missed by using a classical binding affinity assay.

Conclusion
Here, we studied protein complex formation on reconstituted cytoskeleton-free plasma membrane and demonstrated that membrane shape fluctuations on nanoscales lead to cooperative binding. In contrast to our plasma membrane vesicles, cellular plasma membranes are bound to a cytoskeleton that confines shape fluctuations on length scales larger than the cytoskeletal mesh size (~50-100 nm) (Morone et al., 2006), but hardly affects fluctuations on lateral scales of tens of nanometers (the typical distance between neighboring bond complexes) that lead to cooperative binding (Weikl et al., 2009; Hu et al., 2013). Therefore, these nanoscale thermal fluctuations, and thus the binding cooperativity, should persist in living cells. Cell membrane fluctuations appear to be driven by active processes and by thermal motion (Biswas et al., 2017; Turtlier et al., 2016; Monzel et al., 2015), and both can enhance binding cooperativity. In general, fluctuations and displacements of single, unbound membranes should be distinguished from distance fluctuations between two membranes adhering via protein complexes (Lipowsky, 1995). The distance fluctuations of two adhering membranes or of one membrane adhering to a surface are constrained by the protein complexes on length scales larger than the typical distance between neighboring complexes, which leads to binding cooperativity (Fig. 2). GPMVs are known to exhibit many aspects of native cell membranes, and yet the membrane mechanical properties can be described using a just few parameters that connect experiment to theory and simulation. Membrane fluctuations are reduced when the plasma membrane is stiffened by a pH shift to pH 6, which can be induced by tumor tissue acidosis (Fig. 3) (Tannock and Rotin, 1989). In experiments on plasma membrane vesicles, multiscale simulation and phagocytosis, we consistently find reduced CD47 signaling (Fig. 4A). Clearly, enhanced phagocytosis at pH 6 is not caused exclusively by the diminished CD47 (Kd) binding affinity. However, the model system presented here can be used to address the contribution to total cell signaling from binding affinities of individual signaling components in absence of the cytoskeleton, while presenting a physiological membrane-bound environment. Finally, we speculate that reduction in CD47 signaling by membrane rigidification selects for the elevated levels of CD47 in many tumors. Reduction of CD47 signaling would otherwise tend to enhance phagocytosis.

Multiscale simulation and experimental approaches identify a few intuitive parameters, which describe binding, unbinding and cooperative effects for a wide range of membrane-bound complexes. Binding and signaling in cell-cell interactions are not only determined by the relative affinity of the complexes but also indirectly by changes such as the membrane fluctuation spectrum. Our method of reconstituted adhesion should have many applications where affinities of cell signaling complexes are investigated. Ultimately, convolution of membrane fluctuations with binding of adhesion molecules provides an elegant way to control activation and clustering for cell signaling.

MATERIALS AND METHODS
Generation of CD47-GFP HEK cell line and cell culture
HEK 293T cells were transfected with a lentivector encoding CD47-GFP and puromycin resistance transgenes. Lentiviral transduction was carried out over 72 h with a MOI of 5 and followed by puromycin selection (3 μg/ml from Gibco by Life Technologies) for 3 weeks. Cells were then cultured under standard condition in DMEM (without Phenol Red; Millipore-Sigma), 10% FBS and 1% penicillin-streptomycin and split every 2-3 days.

GPMV Isolation
Before experiments, CD47-GFP HEK cells were plated under identical conditions in T-25 culture flasks and cultured for 3 days until cells were

---

Fig. 4. Phagocytosis and CD47 levels change with acidosis. (A) Phagocytic uptake assay of opsonized A549 (red in insets) cells by THP-1 macrophages (green). Yellow indicates overlapping cells, and full overlap indicates completed phagocytosis. Shift to pH 6 increases such engulfment. Left inset shows phase-contrast images overlaid with fluorescence. Scale bars: 10 μm. Data from two independent experiments (n=90 cells for pH 7, n=47 for pH 6). (B) CD47 levels increase on viable A549 cells under acidosis conditions. Experiment was performed with three technical replicates. Error bars are ±s.e.m. Differences between distributions are significant (P<0.05, t-test).
90% confluent. GPMVs were isolated according to a published protocol (Sezgin et al., 2012). HEK cells were incubated with 2 mM DTT, 25 mM paraformaldehyde at 37°C for 1 h. For experiments in which the pH was varied, the GPMV buffer was adjusted to the desired pH on the day of the experiment.

**Purification and absorption of SIRPs on glass slides**
The soluble extracellular domain of SIRPα was purified as previously described (Seifert et al., 1999). The resulting protein was tagged with GST and we refer to the resulting SIRPαEx-GST as to ‘SIRPs’. SIRPα at a concentration of 34 nM was then incubated on clean glass slides (2× rinsing with ethanol and ultra-pure water or a 5 min exposure to oxygen plasma) with room temperature. Unbound SIRPα was washed away and uncovered glass surface was blocked by incubation with 1% BSA, 0.05% Tween-20 for 1 h at room temperature. The coated glass slides were then washed in GPMV buffer at the desired pH and immediately covered with GPMV suspension.

**Confocal imaging, FRAP experiments and image analysis**
Confocal images were obtained on a Leica SP8 system, dyes were excited using the 488 and 638 laser lines and fluorescence was collected (495-600 nm and 650-750 nm). To avoid cross-talk between the channels, two-color images were always obtained sequentially. FRAP experiments were performed on the same system. A circle of 1 µm in diameter was bleached using the 488 laser for two consecutive frames and recovery was observed under normal imaging conditions for 60 frames and 0.2 frames per second. The data were corrected for bleaching by dividing the intensity obtained in the bleached spot by the intensity of an unbleached area about 10-15 µm away from the bleaching spot. The background level was subtracted as estimated from the first frame after bleaching. To estimate the diffusion of free CD47, GPMVs were bleached on the upper pole and recovery curves were fitted using the standard model for free diffusion (Soumpasis, 1983). For the bound membrane segment fluorescence recovery, a reaction diffusion model was used (Sprague et al., 2004):

\[
\frac{\partial[L]}{\partial t} = D_{frev} \nabla^2[L] - k_{on}[L][R] + k_{off}[RL],
\]

\[
\frac{\partial[R]}{\partial t} = -k_{on}[L][R] + k_{off}[RL],
\]

\[
\frac{\partial[RL]}{\partial t} = k_{on}[L][R] - k_{off}[RL].
\]

Here \([L]\), \([R]\) and \([RL]\) are the concentrations of free CD47-GFP, surface-immobilized SIRPα molecules and bound complexes respectively. \(\nabla^2\) has the usual meaning of the Laplace operator and \(D_{frev}\), \(k_{on}\) and \(k_{off}\) are parameters of the model. The data was subsequently fitted to the model using the routine MATLAB invlap (Karl J. Hollenbeck, 1998) and nlinfit (MATLAB, 2014a). Note that within the bleached spot there are only about 30-80 CD47 molecules; hence in the first seconds of recovery only a few GFP molecules contribute to the signal. The data is inherently noisy and we found that the recovered absolute values for \([R]\) and \([L]\) depend significantly on the starting values of the fitting routine. However, the ratio of these rates is robust; in other words \([R]\) and \([L]\) are obtained with higher degree of confidence. The error in fitting the individual fluorescent recovery trajectories was estimated following the method Müller et al. (2009). The value of the best fit of \([R]\) and \([L]\) is varied around the best estimate (in the least-square sense) until the deviation is found to be outside the bounds set by the χ² distribution, effectively giving the 95% confidence intervals.

Fluorescence intensities were quantified either by a line profile on the vesicle equator (to obtain the correlation between GFP and anti-CD47), a line profile on the x-z contour in the case of adhering vesicles, both lines were 1 µm wide, as shown in Fig. S1. We used the peak of the resulting intensity distribution as an indicator of dye concentration in the membrane. Using this method, we could quantify both the fluorescence intensity in the bound and unbound membrane segments, \(I_{bound}\) and \(I_{free}\) respectively. We found that the value \([RL] = G(I_{bound} - I_{free})\) is the most robust way to quantify CD47-SIRPα complex densities using fluorescence confocal microscopy, presumably because small variation or defects in GFP tagging are eliminated this way. Here, \(G\) is the proportionally factor to convert fluorescent intensities to number density per membrane area and is determined as described below. Plots of \([RL]\) vs \([L]\) were in general too scattered for detailed analysis and we used these values only as a consistency check as shown in Fig. 3C. To assess the CD47 density between RBC and GPMVs we compared the maximum intensity projection at the upper surface of labelled RBC to GPMVs as obtained by high-resolution confocal microscopy (63×1.2 water immersion lens). To convert fluorescence intensities to CD47 concentration, we incubated red blood cells (Fig. S3B), which naturally express CD47 (about 250 molecules/µm²; Tsai et al., 2010), together with the GPMVs and fluorescent CD47 antibodies. The average anti-CD47 fluorescence of RBC and co-incubated GPMVs was determined from \(x\) y confocal sections of circular shape (≈1 µm²) at poles and the average number density per area of CD47 molecules on the GPMV population was determined to be about 100 CD47/µm². The factor \(G\) was subsequently assessed from the average CD47 coverage and the average GFP fluorescence intensity in the GPMV population, as determined from line profiles in \(x\) y confocal scans on free GPMVs obtained as described above.

**Immunocytochemistry**
CD47 on GPMVs was labelled by incubation of B6H12 Alexa Fluor 647 mouse anti-human antibody (BD Biosciences, 561249; 1:100) for 1 h at room temperature (RT) according to the manufacturer’s protocol. RBCs were isolated from a healthy male individual, washed in PBS and incubated in the same conditions as the GPMVs with the antibody. In some experiments, SIRPα was labelled on the glass surface by incubating with rabbit anti-GST Alexa Fluor 488 (Invitrogen) at 20 µg/ml for 1 h at RT and subsequent washing.

**Phagocytosis assay**
A549 Cells were opsonized with a rabbit anti-human RBC antibody (Rockland, 109-4139 polyclonal; Alvey et al., 2017) at final concentration of 250 µg/3 ml for 1 h at RT. Cells were washed and labelled with the dye PKH26 (Sigma-Aldrich) according to the manufacturer’s protocol. THP1 cells were left to adhere in standard cell 6-well culture plates in RPMI/PMA medium (Millipore-Sigma; initial concentration 4×10⁵ cells per well) for 48 h. A549 cells were then added to the wells and incubated at either pH 7.4 or pH 6. After 75 min, wells were digested with trypsin so that ‘uneaten’ A549 cells were removed by trypsinization. THP-1 were labelled by incubation with CD11b APC antibody (Biolegend, 101211, clone M1/70; Alvey et al., 2017) at 2 ng/ml for 1 h at RT. Finally, cells were fixed by incubation with 4% paraformaldehyde. Imaging was performed using a 10× objective on Leica SP8 system as described below. Images were processed by a Gaussian filter, edges of cells were detected and thresholded to obtain binary images of cells (ImageJ v.1.50b). Overlay of the different channels was then calculated per cell.

**3D binding affinity assay using RBCs**
Fresh RBCs were washed and incubated with SIRPα (final concentration 34 nM) for 1 h at RT in PBS adjusted to pH 7.4 or pH 6. GST was labelled as described before. RBCs were pelleted and unbound protein was washed away. RBCs were then imaged and the mean fluorescence in the center of a RBC was measured in a circle of 6 µm diameter.

**Quantification of CD47 expression during long-term acidosis**
A549 cells were seeded and grown until confluent for 3 days in full F12 medium (supplemented with 25 mM HEPES) in the absence of carbon dioxide. Cells were cultured either at pH 7.4 or pH 6. Cells were then detached non-enzymatically, washed and resuspended in PBS supplemented with 5% FBS. CD47 was labeled by incubation for 1 h at RT using 3 µl Alexa Fluor 647 human CD47 antibody (BD Biosciences, 561249, clone B6H12; Alvey et al., 2017), 0.5 µl Hoechst 33342 (Thermo Fisher, H3570) and 1 µl of 7-amino-actinomycin D (Sigma, A9400-1MG) per 300 µl cell suspension. Cells were then washed, resuspended in Flow buffer (5% FBS in PBS), and transferred to flow cytometry tubes. Cell suspension was analyzed at a flow rate of <1000 cells per second for 1 min. For analysis, aggregates were removed from data using Hoechst intensity and forward scatter signal (FSC).
Dead cells, identified as 7ADD1 and low FSC, were also removed from analysis. Quantification of CD47 intensity was recorded as the geometric mean of a given population of cells.

Fluctuation spectroscopy
We measured the bending rigidity of the GPMVs by fluctuation analysis of the thermally induced motion of the membrane. Details of the method are published elsewhere (Gracia et al., 2010). Experiments were performed on an Axiovert 135 microscope (Zeiss, Germany) using a 40× objective in phase contrast mode. Imaging was done with a fast digital camera HG-100K (Redlake, San Diego, CA) using a mercury lamp HBO W/2 as a light source. We acquired a total of 4000 snapshots per vesicle with exposure time of 200 µs. Only vesicles with clearly visible fluctuations and no visible defects were considered for analysis.

Molecular modeling and coarse-grained simulations of the CD47-SIRPα complex
A structural model of the full-length protein CD47 in complex with the SIRPα-GST construct was built using PDB structures 2JJS, 2WNG and 1GTA for the CD47 ectodomain, the SIRPα ectodomain and the GST domain, respectively. The transmembrane domain of CD47 (comprising residues 117 to 275) was modeled using EVfold (Hopf et al., 2012) and positioned in the membrane environment using the PPM server (Lomize et al., 2006). The ectodomains of CD47 and SIRPα as well as the GST domain were next positioned beneath the membrane (see Fig. 2B) using VMD (Humphrey et al., 1996). The interface between CD47 and SIRPα was taken to be as in the PDB structure 2JJS. The 33-residue peptide segment connecting SIRPα with GST, the 5-residue linker between the two domains of CD47, and the 7-residue loop in the C-terminal Ig-like domain of SIRPα (comprising residues 288 to 294) were built into the structural model using MODELLER (Fiser et al., 2000). The resulting structural model was taken as the initial conformation for molecular simulations. Equilibrium conformations of a single CD47-SIRPα complex were simulated using the coarse-grained model of Kim and Hummer (2008). The model was adapted to the CD47-SIRPα complex in the following way. Four protein domains were represented as rigid bodies: the transmembrane domain of CD47, the ectodomain of CD47, the ectodomain of SIRPα and the GST domain. The interactions between the individual domains were treated at the residue level and described as in the Kim–Hummer model. A bias potential was applied to 45 inter-domain contacts of the ectodomains of CD47 and SIRPα to ensure the correct binding interface as observed in the crystal structure of the CD47-SIRPα ectodomain complex (PDB code 2JJS). The inter-domain contacts were determined using an overlap criterion (Różycki et al., 2014). The bias potential was taken to have a form of the Lennard–Jones potential. The depth of the Lennard–Jones potential was set to 0.67 kT to prevent dissociation of the CD47-SIRPα complex. The GST domain was positioned on a flat substrate with the linker connecting GST to SIRPα oriented away from the substrate and the transmembrane domain of CD47 was embedded in a membrane (see Fig. 2B). The membrane was assumed to be flat and parallel to the substrate surface. Both the 33-residue peptide segment connecting SIRPα with GST and the 5-residue linker joining the two domains of CD47 were represented as chains of amino acid beads with appropriate bending, stretching and torsion potentials (Kim and Hummer, 2008). The interactions between the flexible linkers and the rigid domains as well as the interactions between the membrane and the proteins were simulated as in the original model of Kim–Hummer. A harmonic potential was applied between Cys 15 (located in the extracellular domain of CD47) and Cys 245 (positioned in the transmembrane domain of CD47) to mimic the disulfide bond. Monte Carlo (MC) simulations
The basic MC moves of the CD47 and SIRPα ectodomains were random rotations and translations in three dimensions. For the transmembrane domain of CD47, the basic MC moves were translations parallel to the membrane surface and rotations around the axis perpendicular to the membrane surface. For the linker peptides, crank-shaft moves are employed to enhance sampling in addition to local MC moves on each amino acid bead. The separation between the substrate surface and the membrane was varied in additional MC moves. In the MC simulations, 10⁵ configurations taken every 10³ MC sweeps were generated for analysis. The simulations were performed at room temperature (T=300 K). The ectodomains of CD47 and SIRPα were observed to remain in the bound state throughout the simulation runs. The basic MC moves in our simulations of large adhering membrane segments were variations of the local separations one of all membrane patches, and hopping of the harmonic bonds between neighboring patches (Weikl and Lipowsky, 2006). The bonds are taken to be mobile because positions of CD47-SIRPα complexes in the experiments vary due to binding and unbinding events. Simulations were performed with up to 8×10⁵ MC steps per membrane patch.

Distance of membrane to substrate in molecular simulations
The separation between the membrane and substrate was identified by measuring the distance in the z-direction between the C-terminal residue of the short inter-domain linker in CD47 and the plane at which the GST domain was placed. The histogram of the separations obtained from the MC simulations is shown in Fig. S2. The key quantities obtained from this histogram are the mean value $\bar{z}$=17.2 nm, which is the preferred membrane-substrate separation of the complex, and the standard deviation $\sigma$=1.2 nm, which reflects the flexibility of the complex. Visual inspection of the simulation structures reveals that variations in the separation between membrane and substrate are related mainly to the flexibility of the long linker between SIRPα and GST. Other joints between the rigid domains appear to be relatively stiff. First, in CD47, the transmembrane domain is connected to the ectodomain by a short linker and by the disulfide bond between Cys15 and Cys245. These two connections, together with steric effects between the protein and the membrane, effectively restrain the orientation of the ectodomain relative to the transmembrane domain. Second, the relative orientation of the ectodomains of CD47 and SIRPα is restrained by the geometry of their interface. In fact, only small fluctuations around the orientation given by the crystal structure were observed in the simulations.

Acknowledgements
Some data and Materials and Methods in this paper form part of the PhD thesis of Jan Steinkühler, Technischen Universität Berlin, 2016.

Competing interests
The authors declare no competing or financial interests.

Author contributions

Funding
Financial support for J.S. from the Deutsche Forschungsgemeinschaft (DFG) (IRTG-1524) is gratefully acknowledged. B.R. was supported by the National Science Centre, Poland (2016/21/B/NZ1/00006). This work is part of the MaxSynBio consortium which was jointly funded by the Federal Ministry of Education and Research of Germany and the Max Planck Society. Financial support for C.A. and D.E.D. is from the National Institutes of Health (National Cancer Institute U54-CA193417, National Heart Lung and Blood Institute R01-HL124106, R21-HL128187), and a National Science Foundation Materials Science and Engineering Center grant to the University of Pennsylvania. Deposited in PMC for release after 12 months.

Supplementary information
Supplementary information available online at http://jcs.biologists.org/lookup/doi/10.1242/jcs.216770.supplemental

References
