A Multiscale Study of Phosphorylcholine Driven Cellular Phenotypic Targeting

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INTRODUCTION

Selectivity and specificity are among the most desirable qualities for a drug. The former defines the ability of a drug to target only a particular cell population, and the latter ensures that it has an impact on that cell population. Though small molecules account for most therapeutics in use, their poor solubility in water, inability to cross cellular membranes, and promiscuous interactions (leading to adverse side effects) have placed the focus in recent years on drug delivery systems.\(^1\) Fuelled by many advances in nanotechnology and biotechnology, the past decades have witnessed rapid growth in the research and development of drug delivery devices in the form of polymeric multivalent carriers achieving phenotypic targeting.\(^2\) The physicochemical properties of nanocarriers are easy to tune, and a high degree of selectivity can be achieved by decorating their surface with ligands. Nanocarriers’ high selectivity increases their ability to cross biological barriers that small molecules cannot overcome, opening the door to target biological macromolecules inside the cells,\(^5\) including cells within the central nervous system.\(^6\)

The higher the ligand affinity, the lower the ligand concentration required to saturate its receptor. We can enhance affinity by creating a carrier containing multiple ligands targeting the same receptor in the surface (multivalent scaffolds),\(^7\) therefore increasing the drug carrier affinity or, in this case, its avidity.\(^8\) Nature exploits the collective binding effect of multivalent objects and avidity in most biological processes.\(^9\) Multivalent interactions in biological systems enhance weak individual interactions and change the proximity of the proteins in the cell (clustering), inducing signal transduction.

Although high affinity is a desirable quality, the targeted receptors are expressed in both tumor and healthy cells in diseases such as cancer. Therefore, high-affinity ligands will bind to any cell that expresses the targeted receptors, thus leading to unwanted interactions that, in some cases, outweigh the clinical benefits. However, in 2007, Carlson et al. showed that multivalent targeting was more selective when multivalent low-affinity ligands were used,\(^10\) a concept that was later mathematized by Martinez-Veracoechea and Frenkel, in what they called superselectivity theory (SST).\(^11\) SST shows that the combination of multiple low-affinity ligands creates on–off

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association profiles, where the multivalent scaffold saturates the receptors only above a given cutoff receptor density, and it does not bind below that density. However, multivalent systems are strongly affected by nonspecific binding of the ligands to untargeted receptors due to the weak affinity of the single ligands. Moreover, using a multiplexed-multivalent strategy, e.g., including multiple ligands that target different receptors, we can target a specific cell phenotype and increase the selectivity of the carrier toward a particular cell population. Our group has recently shown that we can still use high-affinity ligands and engineer the drug carrier surface, lowering the overall affinity of the carrier by including a repulsive element that shields the ligands.

The scavenger receptor class B member 1 (SRB1) and scavenger receptor class B member 3 (CD36) can be targeted using PMPC-decorated polymersomes (PMPC Psomes). The high-affinity interactions of PMPC Psomes to SRB1 and CD36 are due to the phosphorylcholine groups (PC) present in the PMPC chains, which induces their internalization via endocytosis in cells. Moreover, we have shown that the affinity of PMPC for SRB1 allows Psomes to target M. tuberculosis and S. aureus infected macrophages as well cancer cells. SRB1 has been associated with CD81, a four-pass transmembrane protein belonging to the tetraspanin family, in the entry mechanism of Plasmodium sporozoites into hepatocytes. Tetraspanins play diverse roles in the immune systems and cancer, and they have been described as a receptor for cholesterol. Still, there is a need to understand exactly the role of these receptors and how they interact with PMPC Psomes.

Rational drug design relies on single-ligand affinities to describe the interaction of a drug with the cell receptor or target, but in the case of big objects like viruses, small proteins, or nanoparticles, one must consider the repulsive effect and steric hindrance of the cell glycocalyx in the molecular recognition process of the receptors expressed in the cell membrane. Most cells are covered by a complex polysaccharide matrix comprising proteins and complex sugar chains (glycosaminoglycans and glycans), forming the glycocalyx. Post-translation modifications can occur at specific sites on protein backbones at N-linked or O-linked residues by the addition of glycans, altering the physical environment of the cell-surface receptors and modifying nanocarrier affinities.

This work shows that by taking advantage of very promiscuous binding motifs or ligands, we can selectively target precise cell populations in vivo. We use a multiscale approach,
starting from the all-atom molecular modeling of the ligand/receptors (including the receptor glycosylation) involved in the uptake affinity. We then build up a statistical model based on the description of the interactions between the nanocarrier and the targeted cell phenotype (receptor density and glycocalyx). Our model describes the \textit{in vitro} and \textit{in vivo} superselective targeting of monocytes using phosphorylcholine-based polymersomes.

\section*{RESULTS AND DISCUSSION}

Receptors Involved in the PMPC Psomes Uptake, \textit{In Vitro}. We investigated the role of SRB1, CD36, and CD81 on the cellular uptake of PMPC Psomes (PMPC\textsubscript{25}-PDPA\textsubscript{70}). We considered three cell types: human primary dermal fibroblasts (HDF), oral carcinoma FaDu cell line, and the human monocytic cell line THP-1. First, we confirmed that all the cell types express the receptors of interest by Western blot. All three cell types highly express CD81 and CD36, while SRB1 expression fluctuates among the cell lines, being less expressed in HDF than in FaDu and THP-1 cells (Figure 1a). The cellular distribution of the receptors is represented in immunofluorescence micrographs (Figure 1b). The uptake kinetics of the PMPC Psomes in all cells is shown in Figure 1c. The cellular uptake of PMPC Psomes was measured by flow cytometry and fitted as $A(t) = A(\infty) \times (1 - e^{-t/\tau})$, where $A(t)$, with a single exponential association model, is the relaxation time and $A(\infty)$ the uptake at equilibrium. Although

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{PMPC free chain binding to SRB1, CD36, and CD81 receptors. Autodock vina estimated relative binding affinities are shown as swarm plots at increasing number of PMPC chains ($N_{PC}$) for SRB1, CD36, and CD81 (a). All-atom depiction of the PMPC free chain (for $N_{PC}$ with highest affinity) binding site, shown as VDW surface in magenta. For CD36 and SRB1, glycans are shown as licorice and their VDW surface is also overlay. The secondary structure of the three receptors is shown as a cartoon in silver. Cellular membrane is indicated in gray. In the CD81 case, two binding modes are depicted (b). PMPC free chain binding site inside CD36 and SRB1 (c). PMPC free chain docking pose versus MD stable pose onto CD81 (d).}
\end{figure}
PMPC Psome uptake plateaus within 2 h in all three cell lines, it indicates a strong interaction of the Psomes with the receptors, higher expression of SRB1 correlates with faster uptake (Figure 1c). Our simple association model shows different relaxation times for the PMPC Psome cellular uptake: \( \tau_{\text{THP-1}} = 7.9 \pm 2.6 \) min (THP-1 cells), \( \tau_{\text{FaDu}} = 19.2 \pm 2.3 \) min (FaDu cells), and \( \tau_{\text{HDF}} = 54.4 \pm 4.1 \) min (HDF cells). We also determined the role of these receptors in the cellular uptake of PMPC Psomes using specific antibodies to selectively block each receptor following 1 h incubation with fluorescent-labeled PMPC Psomes (Figure 1d). As expected, the Psome uptake was significantly impaired by blocking CD81 in all cell types, while SRB1 blocking inhibited the Psome uptake in THP-1 and FaDu but not in HDF cells. On the other hand, CD36 blocking did not affect any of the three cell types, and only when both CD36 and SRB1 were blocked, we observed an uptake inhibition in all cell types including the HDF (Figure 1d). This reduction in Psome uptake in HDF resulted from the synergistic effect of blocking CD36 and SRB1. With the levels of the receptors in HDF compared to other cell lines kept in mind, this result suggests that the cellular uptake in HDF is preferentially through CD36 and CD81. In contrast, carcinoma cells prefer the Psome cellular uptake through SRB1 and CD81 receptors. Moreover, the multiple receptor binding is associated with the clustering of receptors

Figure 3. PMPC binding induces CD81 opening and membrane curvature. Conformational landscape explored by the PMPC free chain (\( N_{\text{PC}} = 4 \)) on the surface of CD81. The free-energy landscape is projected onto the distance of the center of mass of the PMPC free chain and the surface of CD81 and the polar angle (described in Figure 2d) (a). Time evolution of the angles describing the allosteric movement induced by PMPC onto the helix TM4 of CD81, colored according to simulation time. The TM4-EC2 angle is highlighted in red and the TM4 kink in lavender (b). Initial membrane curvature represented by the phosphorus atoms of the lipid heads (c). All-atom depiction of CD81 (as white cartoon) and the PMPC chain (magenta surface) along the MD simulation. The TM4 is colored according to simulation time. The angles describing the allosteric movement are indicated in snapshot 1 (TM4-EC2) and 7 (TM4-kink) (d). Final membrane curvature (e).
and the hijacking of the endocytic machinery.\textsuperscript{20,21} To understand the interaction between the three receptors during PMPC Psome cellular uptake, we performed a proximity ligation assay (PLA) on FaDu cells.\textsuperscript{22} After 1 h of incubation with Psoomes, we observed an increase of the PLA signal in all combinations of receptors (Figure 1e). In physiological conditions, the three receptors are widely distributed on the surface of the cells and modestly associated with one another. However, we observed a striking increase in the clustering of SRB1 with both CD36 and CD81 after incubation with PMPC Psoomes. This observation was confirmed using an ad hoc pro algorithm or PLA image analysis,\textsuperscript{23} highlighting the significant clustering of SRB1 with both CD36 and CD81 receptors (Figure 1e).

Thus, the data in Figure 1 demonstrate the role of the SRB1, CD36, and CD81 for PMPC Psoome uptake. Moreover, the Psoome uptake appears to depend on receptor expression in cells. Indeed, the low levels of SRB1 in HDFs are very likely balanced through the CD36 receptor, which shares several ligands with SRB1.\textsuperscript{24,25} The uptake kinetics suggests a critical role of SRB1 with a correlation between the uptake rate and the cellular expression in FaDu and THP-1. Finally, the PLA data shows that SRB1 clusters with CD36 and CD81 during the uptake process, suggesting a possible role for the latter.

**Modeling the PMPC Binding to SRB1, CD36, and CD81**

**In Silico.** We performed an in silico characterization of the PMPC interaction with the three receptors involved in the PMPC Psoome cellular uptake in vitro shown in Figure 1. For each receptor, we built an all-atom model (Figure S1), including all the predicted glycosylations (see Methods) for CD36 and SRB1 (Figure S1, Figure 2e). We assessed the relative interaction affinity of different free PMPC chains (with varying degrees of polymerization, \( N_{PC} \)) with the three receptors using docking techniques (see Methods). We computed 3000 binding models, 100 models per receptor at an increasing number of PC units. We report in a swarm plot the relative binding affinity as a function of \( N_{PC} \) for SRB1, CD36, and CD81 (Figure 2a). As expected, the binding affinity of PMPC in SRB1 and CD36 is higher than in CD81. Although the predicted overall architecture for CD36 and SRB1\textsuperscript{26} is identical, their amino acid nature composition (Figure S2a) and glycosylation pattern (Figure 2b, Figure S2) differ between the two, influencing their interaction with the PMPC free chain. The oxidized phosphatidylcholine (PC) binding site is well-known in both SRB1 and CD36.\textsuperscript{27–29} The affinity of PMPC toward both receptors follows the same trend, but chains with a low polymerization degree (\( N_{PC} = 1 \) to \( N_{PC} = 5 \)) occupy different regions in CD36 and SRB1 (Figure S3). In CD36, PMPC (\( N_{PC} = 1 \)) binds in a deep pocket with the PC motif forming hydrogen bonds with an arginine (R62) and the side chain of a tyrosine (Y78) (Figure S3), while in SRB1, both amino acids are replaced by a phenylalanine (Figure S2), and the interaction is not favorable. Hence, PMPC (\( N_{PC} = 1 \)) binds in the upper part of the receptor (Figure S3), where its PC motif forms a hydrogen bond with an arginine (R296) (Figure S3). With increasing \( N_{PC} \), the PMPC chain binding site migrates toward the PC binding region described in the literature.\textsuperscript{30,31} In the maximum affinity configuration in both receptors (Figure 2b), PMPC occupies the maximum volume inside the receptor (Figures 2b and S3). In CD81, the PMPC free chain reaches the maximum number of possible interactions with the surface of the receptor at \( N_{PC} = 4 \), and it plateaus at \( N_{PC} = 8 \). As shown in Figure 2b, only the top surface of CD81 is exposed to the solvent and available for interaction. We confirmed the stability of the maximum affinity pose for all three receptors using molecular dynamics (see Experimental Methods: Molecular Modeling section). The docking pose (Figure 2bc) is stable for both CD36 and SRB1 during the microsecond long run (Figure S4). The PC units in SRB1 form only 3 to 4 hydrogen bonds with the receptor during MD (Figure S4bc), while in the docking pose it exhibits 6 contacts (Figure 2c). The full PMPC (\( N_{PC} = 5 \)) chain is very stable inside CD36, forming, on average, 1 to 7 hydrogen bonds with the residues in the binding pocket (Figure 2c, Figure S4bc). Only one PC unit remains fixed (RMDS < 4 Å, Figure S4a,b) inside CD36/SRB1 during the simulation, while the rest of the PC units are exposed to the solvent and have higher mobility. In CD36, the binding pocket is positively charged with several lysines and arginines available for interaction (Figure 2c), while in SRB1, the interactions with positively charged residues are replaced by residues with polar chains like serine or asparagine (Figure S2). In CD81, the docking pose in which PMPC binds to helix A is not stable, and the PMPC chain leaves the binding site (Figure 2d) within 10 ns. The PMPC chain binds and unbinds multiple times throughout the simulation. Nonetheless, the site highlighted in Figure 2d, where PMPC binds to helices C and D, remains stable during more than 400 ns.

We computed the free energy landscape from the 1.5 µs trajectory (Figure 3a); the barrier for binding/unbinding is very low, \(-4\) kcal/mol. Interestingly, the unbinding of PMPC from the EC2 induces an allosteric movement of the transmembrane helix 4 (TM4) (Figure 3d), which has only been described recently\textsuperscript{18} in the absence of cholesterol in the binding site. The observed opening reported enabling the export of cholesterol\textsuperscript{18} involves the detachment of helix B and involves the recruitment of a partner, CD19.\textsuperscript{18,34} In our case, the rebinding of the PMPC chain to the EC1 loop induces the opening of the EC2 that displaces helices C and D, increasing the TM4-EC2 angle (Figure 3d), but the receptor is still closed. Rebinding into helices C and D induces a kink in TM4 (Figure 3de), that has not been previously described in the literature. This rearrangement or kink of TM4, which resembles the TM6 kink in G protein receptor activation,\textsuperscript{35} induces a negative curvature in the lipid membrane (Figure 3e) as opposed to the initial flat configuration (Figure 3c), which is compatible with the hypothesis of the CD81 acting as a membrane reshaper, as previously hypothesized for tetraspanin CD9.\textsuperscript{35} To confirm that the free PMPC chain induces the allosteric movement described in Figure 3, we ran a simulation without PMPC and analyzed the conformational plasticity of the receptor (Figure S5). Both the TM4-EC2 angle and the TM4 kink remain stable during the simulation (Figure S5a). Despite the interaction of the intramembrane helices with the lipids, the extensive allosteric rearrangements observed in the presence of PMPC are not present. The TM4-EC2 angle explores two conformations, \( 130° \) and \( 145° \), though the second has a lower population (Figure S5b), which is compatible with the previously investigated closed structured,\textsuperscript{34,35} while in the presence of PMPC, the observed values for this angle are above \( 155° \). Interestingly, the simulation without PMPC revealed an alternative kink in TM4 (Figure S5c) that induces a positive curvature in the membrane due to the interaction with the lipids, as already hypothesized for tetraspanins.\textsuperscript{31}

**Partition Function of the Binding of the Single PMPC Chain.** Despite the particularities of the interaction pattern for each PMPC chain, we observed that the binding affinity changes non-monotonically with \( N_{PC} \) for all three receptors. As one PC
unit is bound to its natural site, the other units are forced to interact with the juxtaposing residues giving rise to a cooperative effect, where the number of binding sites, \( \lambda \), increases with the \( N_{PC} \). We can write, in a first approximation, that \( \lambda \approx 1 + b(N_{PC} - 1) \) with \( b \) being an arbitrary constant. Hence as the single PMPC chain binds to SRB1, CD36, or CD81, each interaction of all \( \lambda \) binding sites are within reach of each PC unit, giving rise effectively to a radial topology. If we assume that \( N_{PC} \gg \lambda \) and write the partition function between the free PMPC chain and the \( \zeta \) receptor as

\[
d_{PC/\zeta}^{free} = \sum_{j=1}^{\lambda} \left[ (\lambda_j) \right]^{N_{PC} - 1} e^{-(E_{g,N} + U_{PC,N})/kT} \]

where \( k \) is the Boltzmann constant, \( T \) is the absolute temperature, \( E_g \) is the binding energy between the PC unit and its relative site on the receptor surface, and \( U_{PC,N} \) is an energy term that takes into account any steric effects emerging from the chain binding at \( N_{PC} > 1 \) and can be approximated as \( U_{PC,N} \approx u_\zeta(N_{PC} - 1) \) with \( u_\zeta \) being the steric repulsion between the nonbound PC units chain and the receptor. From eq 1 we can thus derive the binding energy of the single free PMPC brush to the receptor as

\[
E_{PC/\zeta}^{free} = -kT \ln \left[ (1 + N_{PC} e^{-(E_g + u_\zeta(N_{PC} - 1))/kT})^{-1} \right]
\]

We used eq 2 to fit the average binding affinities for each value calculated via docking, as shown in Figure 2a. Our analytical model describes the binding behavior of PMPC in CD36 and SRB1. The fitting parameters are similar for both CD36 and SRB1, and we observe how for a single chain above \( N_{PC} = 15 \) the configurations are no longer attractive; the steric contribution takes over, and the overall affinities are positive. In Figure S3, we illustrate this behavior by superimposing a PMPC15 brush onto the PMPC15 binding mode with the highest affinity for both receptors.

**Multivalent and Multiplexed Binding.** The data above show a good agreement between the estimated avidity and a multivalent binding with linear topology, i.e., a single chain binding to the receptor. However, when assembled into Psomes, the PMPC chains are generally packed to the vesicle surface, each occupying an area per molecule \( \sigma_0 \) forming an archetypal Alexander De Gennes polymer brush. Each PMPC polymer can be schematized as a multivalent chain comprising \( N_{PC} \) units of PC (Figure 4a) and thus with end-to-end to distance \( H \approx a_{PC}N_{PC} \) with \( a_{PC} \) being the PMPC monomeric length. When the PMPC Psomes approach the cell surface, the chains binding to SRB1, CD36, and CD81 receptors generate steric repulsion (Figure 4a). Indeed, the binding of the first PC unit is very different from the binding of the PC units buried within the polymer brush. Each \( \zeta \) receptor must insert into the PMPC brush, displacing the chains and giving rise to a steric repulsive potential \( U_{\zeta}(z) \) that increases with the insertion distance along the PMPC chain and normal to the Psome surface, and we define it as \( z \). We can approximate the insertion distance as an integer \( z \in [1, \cdots, N_{PC}] \), with \( z = 1 \) corresponding to the outer layer of the brush, and \( z = N_{PC} \) being the inner layer of the brush at the hydrophilic/hydrophobic Psome interface. We can use eq 2 to derive the binding energy for the chain and within the brush as

\[
E_{PC/\zeta}^{free} = \sum_{\zeta=1}^{N_{PC}} \left[ E_{PC/\zeta}^{free}(z) + U_{\zeta}(z) \right]
\]

\[
\approx kT \sum_{\zeta=1}^{N_{PC}} \left[ -\ln \left( 1 + z e^{-(E_g + u_\zeta(z-1))/kT} \right)^{1+b(z-1)} \right]
\]

\[
-kT \ln \left( 1 + \frac{N_{PC}}{ \frac{N_{PC} - z + 1}{N_{PC}} + \frac{N_{PC} - z}{R} } \right)^{\frac{\gamma}{4}} \left[ \sigma_0 \left( 1 + \frac{N_{PC} - z}{R} \right) \right]^{\frac{\gamma}{2}}
\]

The new steric potential \( U_{\zeta}(z) \) is the consequence of the receptor inserting into the brush and can be derived by adapting the Halperin model as we have previously shown for Psomes. \( V_\zeta \) is the receptor volume, \( R \) is the Psome radius, \( \sigma_0 \) the area per chain, and \( \gamma \) is a geometrical parameter that represents the packing of the chains on a curved surface. For \( \frac{a_{PC}N_{PC}}{R} \leq (\sqrt{3} - 1) \), \( \gamma = \left( \frac{a_{PC}N_{PC}}{R} + 1 \right)^2 \) unless \( \gamma = 3 \). Finally, to account for the cases when \( U_{\zeta}(z) \) overcomes the attraction forces we define the binding energy \( \epsilon_{PC/\zeta} \) per single PMPC chain to the receptor as
Figure 5. Influence of glycans on Psome binding. Lectin binding assay to assess the amount of glycans on FaDu, HDF, and THP-1 cells (a). 3D heat maps showing the Psome surface coverage in the three cell lines analyzed, FaDu, HDF, and THP1, as a function of the Psome radius, R, and the PMPC degree of polymerization, \( N_{PC} \). The different panels correspond to different glycan compositions: from top to bottom, the polymerization degree of the branches is increased (b). 100 snapshots taken at 10 ns intervals for glycans depicted as sticks are superimposed onto the van der Waals surface representation of the receptor. The volume occupied by the glycans is highlighted as a semitransparent surface. 100 snapshots of the bound PMPC free chain (\( N_{PC} = 4 \)) are also depicted as magenta sticks in the binding site. Front and top views of the receptor-free chain complex are provided for SRB1 and CD36. In the bottom panel, a schematic representation of the DNeup5Aca2−6DGalpβ1−4DGlcNAcb1−2DManpα1−6[DNeup5Aca2−6DGalpβ1−4DGlcNAcb1−2DManpα1−3]DManpβ1−4DGlcNAcb1−4DGlcpNAcb1 complex glycans used in the MD simulations together with its all-atom representation and VDW surface (c). Scattered plot of the percentage of the receptor solvent accessible surface area (SASA) hidden by glycans as a function of the glycan radius of gyration for CD36 (blue) and SRB1 (orange). In the mirror axis, the receptor SASA is reported as a solid line (left). The probability density function (PDF) of the receptor volume and the receptor plus glycans extracted from a 1 \( \mu \)s molecular dynamics simulation is reported for CD36 (blue) and SRB1 (orange) (right) (d).

\[
\epsilon_{PC/\zeta} = \begin{cases} 
E_B & \text{for } E_{PC/\zeta} \geq E_B \\
E_{PC/\zeta} & \text{otherwise}
\end{cases}
\]  

(4)

From eq 4, we can now write the partition function for the Psome binding to the receptor \( \zeta \) as

\[
d_{\text{Psome}/\zeta} = \sum_{\lambda=1}^{\min(N_{PMPC},N_{\zeta})} \left( \frac{N_{PMPC}}{\lambda} \right) N_{\zeta}^{\lambda} \lambda! e^{-\epsilon_{PC/\zeta}/kT}
\]  

(5)

where \( N_{PMPC} = 4\pi R^2 / \sigma_0 \) is the number of PMPC chains per Psome surface and \( N_{\zeta} \) is the number of \( \zeta \) receptors expressed on the given cell. Even small Psomes comprise thousands of PMPC chains, and we can always assume \( N_{PMPC} \gg N_{\zeta} \) simplifying eq 4 as

\[
d_{\text{Psome}/\zeta} = \left( 1 + \frac{4\pi R^2}{\sigma_0} e^{-\epsilon_{PC/\zeta}/kT} \right)^{N_{\zeta}} - 1
\]  

(6)
Finally, most cells have their surface coated with a complex mixture of glycan chains expressed by proteoglycans and glycoproteins, forming the so-called glycocalyx.\textsuperscript{19} Such a barrier can be as thick as tens of nanometers, and it creates a steric protection that nanoparticles need to overcome before they reach the membrane and infect the cell. The basic proteoglycan unit consists of a "core protein" with one or more covalently attached glycosaminoglycan (GAG) chains. The resulting polymer brush formed by the many GAG chains will repel the polymersomes approaching the cell surface via a steric potential:

\[ U_{\text{GAG}} = \frac{4}{3} \pi R^3 kT \left( 1 - \frac{\min(h_j)}{d_{\text{GAG}}} \right)^{9/4} \]  \hspace{1cm} (7)

The three receptors here considered are considerably smaller than the GAG chains; hence \( \frac{\min(h_j)}{d_{\text{GAG}}} \approx 0 \) and eq 7 becomes

\[ U_{\text{GAG}} = \frac{4}{3} \pi R^3 kT \]  \hspace{1cm} (8)

We can write the total partition function as

\[ Q_{\text{Psome}} = \prod_{\zeta=CD81,CD36,SRB1} \left( 1 + \frac{4 \pi R^3}{d_{\text{GAG}}} e^{-\frac{\sigma}{\sigma_{\text{GAG}}}} \right)^{N_{\zeta}} e^{-\frac{U_{\text{GAG}}}{kT}} \]  \hspace{1cm} (9)

and from eq 7 we derive the total energy of binding of single PMPC Psomes to a given cell expressing SRB1, CD36, and CD81 as

\[ E_{\text{Psome}} = -kT \sum_{\zeta=CD81,CD36,SRB1} \ln \left( 1 + \frac{4 \pi R^3}{d_{\text{GAG}}} e^{-\frac{\sigma}{\sigma_{\text{GAG}}}} \right)^{N_{\zeta}} \]  \hspace{1cm} (10)

We can approximate the Psome binding to cells as a Langmuir–Hill isotherm\textsuperscript{13,38,39} and derive the fraction of bound particle, \( \theta \), as

\[ \theta = \frac{a Q_{\text{Psome}}}{1 + a Q_{\text{Psome}}} \]  \hspace{1cm} (11)

where \( a \approx \frac{\pi}{3} N_{\zeta} [P] \left[ 3(R + a_{\text{GAG}} N_{\text{PC}}) - R^3 \right] \) is the Psome activity within the binding volume with \([P]\) being the Psome bulk concentration and \( N_{\zeta} \) the Avogadro number. Note that the angle brackets \( \langle ... \rangle \) designate an average over all the possible receptor number \( N_{\zeta} \) distributions weighted by their Poisson probability.

We exposed FaDu cells to PMPC Psomes formed with PMPC–PDPA copolymers with different polymerization degrees to explore the effect of \( N_{\text{PC}} \) on cell binding. The uptake kinetics, reported in Figure S6a, show almost no differences for \( N_{\text{PC}} = 12, 19, \) and 25, and a considerably decreased uptake occurs for \( N_{\text{PC}} = 6 \). We estimated the corresponding Langmuir–Hill isotherm fraction of bound Psomes\textsuperscript{39} from the ratio between the four kinetics and the \( N_{\text{PC}} = 25 \) reported in Figure S6b. We normalized the fluorescence per cell and represent the experimental fraction of bound Psomes, \( \theta_f \), in Figure 4b. For each receptor, we define its relative expression measured by Western blot (Figure 1a) as \( \phi_i = \phi_i N_{\zeta} \) with the parameter \( \phi \) being a constant with dimensions \( [\mu \text{m}^{-3}] \) and dependent on the cell type alone. The Psomes were produced with average radius, \( R = 40 \) nm, and the uptake experiments were performed with bulk concentration \( [P] = 3.5 \times 10^{-10} \) M. The area per chain, \( a_0 = 0.17 \) nm\(^2\) and PC monomer unit length \( a_0 = 0.257 \) nm, while the receptor volumes can be estimated from the structural data shown in Figures 2 and S2. Finally, the binding energy of the single PC unit, \( E_{\text{GAG}} \), as well as the two semiempirical parameters, \( \sigma_{\text{GAG}} \) and \( \sigma_{\text{PC}} \) were calculated from fitting of the docking data (Figure 2a). We can thus fit eq 9 for the experimental fraction of bound Psomes, \( \theta_f \) using the corresponding \( \phi_f \). In Figure 4b, we report the fitted experimental fraction of bound Psomes (and experimental data points) and the total binding energy calculated using eq 8 which shows a clear minimum at \( N_{\text{PC}} \approx 40 \). This indicates that by increasing the number of PC units we can increase the total avidity up to a certain limit. For \( N_{\text{PC}} > 40 \), the steric repulsive potential dominates the overall interaction limiting the binding to the first PC unit on the Psome surface.

**Influence of the Receptor Glycosylation on the Psome Binding.** The PC units that cannot bind any more once the receptor binding site is filled are not the only source of steric repulsion: steric repulsion due to the cell glycocalyx must also be taken into account. We characterize the differences in cell glycocalyx through lectin-binding assay (Figure 5a). Lectin has a high binding affinity for glycoprotein N-acetylglucosamines and has been used to identify the glyocalyx composition.\textsuperscript{40} We show that the HDF cell line saturates at lower lectin concentrations than FaDu and THP-1, indicating the presence of more sugars in the glycocalyx. In Figure 5b we use the experimental values from Figure 1a to fix the \( N_{\text{CD81}}, N_{\text{CD36}} \), and \( N_{\text{SRB1}} \) for the three different cell lines, to estimate the effect of the Psome radius and the degree of polymerization \( N_{\text{PC}} \) on the binding energy. Moreover, in Figures 2b and 5c, we can see that both CD36 and SRB1 are highly glycosylated, with 10 glycosylation sites predicted for CD36 and 9 for SRB1, but only N102 is conserved among the two (Figures S1, S2c).

As shown in Figure 5c, the pattern and regions hidden by the glycans are vast and different between the two receptors. The complexity of the glycans present on the surface of the receptor influences both the solvent-accessible surface area (SASA) of the receptor and its volume. In Figure 5d (left), we calculated the percentage of the receptor SASA hidden by glycans as a function of the glycan radius of gyration which dramatically decreases with increasing glycan complexity until the point in which if all glycosylation sites have been modified with very complex long and branched glycans, the surface of the receptor is completely hidden. Interestingly, and due to the position of the glycans, this effect is bigger in CD36 than in SRB1. In Figure 5d (right), we show the probability density function of the receptor and the total system (receptor + glycans) volume during 1 \( \mu \)s of molecular dynamics. Glycans introduce an effective volume \( V_\zeta \) on the receptor that varies with glycan complexity. The variations of \( V_\zeta \) significantly affect the binding energy of the Psome to the cell. Especially, the number of branches or antenna of the glycans can completely switch off the interaction with the receptors by hiding it. The degree of polymerization of the glycan affects the morphology of the Psomes that can effectively target the cells, both particle radius and the number of PC units. Glycosylation is one of the most important post-translational modifications of proteins, and it varies during the cell cycle and with the onset of disease. Recently, it has been shown how glycosylation affects viral virulence, not only due to the shedding of the virus and hence helping it to escape the recognition by
antibodies but also due to its influence in the binding of the cell receptors.41

The heat maps in Figure 5b show the Psome surface coverage for different N-glycan degrees of polymerization. In HDF cells, only small Psomes (R < 20 nm) are predicted to bind, while bigger Psomes up to ∼60 nm can bind to FaDu and THP-1 cells. By increasing the degree of polymerization of the glycans, Psomes with longer chains are preferred. The binding energy peaks above with N_{PC} = 15–20 for most Psome sizes.

**Superselective Targeting of Monocytes, In Vivo.** To validate our model prediction in vivo, we injected PMPC Psomes with a degree of polymerization N_{PC} = 25 and radius R = 30 nm (Figure S7). We injected intravenously (i.v.) the PMPC Psomes in mice and observed the cellular uptake in different cells present in the bloodstream by flow cytometry. Even though SRB1 and CD36 are mainly expressed in monocytes (Ly6C+ cells), they can also be found in lymphocytes and granulocytes,15,42 while CD81 is not commonly expressed on red blood cells, granulocytes, or platelets, unlike monocytes and lymphocytes.43 We observed a remarkable selectivity of PMPC Psomes toward monocytes (Ly6C+ cells) after 5 min of i.v. injection, while in lymphocytes, granulocytes, and erythrocytes, less than 10% of Psomes were taken up (Figure 6a,b). There are a few hundred monocytes per μL, 5 orders of magnitude lower than erythrocytes (Figure 6c), showing that PMPC Psomes selectively target monocytes.

Moreover, CD36, SRB1, and CD81 levels were down-regulated after 1 h of i.v. injection of PMPC Psomes due to the quick uptake in classical monocytes (Ly6C high). After 24 h, the receptor levels were restored to the same levels as the untreated cells (Figure 6d), while in nonclassical monocytes (Ly6C low), only the SRB1 expression was reduced after 1 h in nonclassical monocytes, and after 24 h, the levels went back to basal levels.

PMPC Psomes enter the cells through the scavenger and tetraspanin receptors especially in monocytes, confirming the selectivity of PMPC Psomes to target monocytes. Moreover, we have presented a statistical model describing the particle-cell interaction, leveraging all-atom simulations, and including a key component of the cell: the
glycocalyx. Differences in cell-glycocalyx translate into differences in the particle design features, such as particle radius and single-chain polymerization degree, leading to successful binding to the cell. We showed that we can leverage our model to optimize Psomes and selectively target a cell population in the bloodstream: monocytes, which account for 2–8% of the blood cells. Thus, the intrinsic avidity of PMPC Psomes toward immune cells, especially monocytes, can be a helpful therapeutic approach in cancer immunotherapy.

**EXPERIMENTAL METHODS**

**Polymersomes Assembly and Characterization.** PMPC–PDPA copolymer was synthesized either by atom transfer radical polymerization (ATRP) or by reversible addition–fragmentation chain transfer polymerization according to a previously published protocol, whereas rhodamine 6G-, Cyamine 3-, and Cy-5 labeled PMPC–PDPA copolymers were always synthesized by ATRP. PMPC–PDPA and rhodamine-labeled PMPC–PDPA assembly was carried out under sterile conditions using the pH-switch method as previously described. Briefly, 20 mg of copolymer was dissolved in a 2:1 mixture of chloroform:methanol (Fisher Scientific), followed by its evaporation in a vacuum oven at 60 °C. This results in the deposition of a thin polymeric film on the walls of the vial that was then dissolved with PBS (100 nM) at pH 2.0 for a final 10 mg/mL solution. Self-assembled structures were formed by dropwise addition of 1 M NaOH to the polymer solution, hence increasing the pH over the PDPA pKₐ (∼6.2) to a final pH of 7.4. This dispersion was then sonicated for 30 min (Sonicator Instrument Corporation) and kept under continuous stirring (200 rpm) for 2 days at room temperature. In order to isolate the vesicular structures from micelles, the dispersion was injected through a hollow fiber with 50 nm pores, using a KrosFlo Research III Tangential Flow Filtration System (Spectrum Laboratories, Inc.). Additional Rho-, Cy3-, and Cy5-labeled PMPC–PDPA Psomes used for imaging purposes were produced by the film rehydration method as previously described. Here, 10% (w/w) of either Rho- or Cy3- or Cy5-labeled PMPC–PDPA was dissolved together with 25 mg of copolymer in a 2:1 mixture of chloroform:methanol. The solvent is then evaporated in a vacuum oven, and the resulting polymeric thin film was rehydrated with PBS (100 nM) at pH 7.4 to a final concentration of 5 mg/mL. Psomes were formed after this solution was kept under continuous shear stress using magnetic stirring (200 rpm, RT15 power, IKA-Werke GmbH & Co.) for 16 weeks. Finally, Psome dispersions were purified via gel permeation chromatography using a size-exclusion column containing Sepharose 4B and PBS at pH 7.4. Afterward, Psome dispersions were characterized in terms of polymer concentration by HPLC, vesicle size, and morphology by DLS and TEM, respectively (details in Supporting Information). Until further in vitro and in vivo experiments, all the Psome dispersions were stored at 4 °C.

**Cells.** Squamous carcinoma cell line (FaDu) was cultured in complete Eagle’s minimum essential medium (Sigma) supplemented with 10% (v/v) heat-inactivated fetal bovine serum (Sigma-Aldrich) and 1% (v/v) penicillin–streptomycin (Sigma-Aldrich). Human leukemia monocytes (THP-1) were cultured and maintained in RPMI1-1640 supplemented with 10% (v/v) heat-inactivated fetal bovine serum (Sigma-Aldrich) and 1% (v/v) penicillin–streptomycin (Sigma-Aldrich). These two cell lines were obtained from American Type culture collection (ATCC). Human dermal fibroblast (HDF) cell line was purchased by Sigma and cultured in complete Fibroblast Basal Medium (Lonza) with 1% (v/v) penicillin–streptomycin.

**Western Blot Analysis.** Cell lysates were prepared using lysis buffer (RIPA buffer, Sigma) with protease inhibitors (Sigma) and collected using a cell scraper and kept on ice for 30 min. Then, the lysates were centrifuged at 13 000 rpm for 15 min, and supernatants were collected. Bradford assay (BioRad) was performed to assess the protein concentrations, and 4× Laemmli sample buffer was added to the cell lysates and boiled for 5 min at 95 °C. 30 μg of protein was loaded per lane in a polyacrylamide gel and run for 120 min at 120 V. The PVDF membranes (BioRad) were used to transfer the proteins from the gel for 90 min at 4 °C and 100 V. The membranes were previously stained with ponceau S and then blocked with 5% nonfat milk in TBST for 1 h and incubated with anti-SRB1 (#ab6942), anti-CD36 and anti-CD81 (Abcam), and GAPDH (Cell signalling) overnight at 4 °C. For detection on the LI-COR Odyssey (LI-COR, Germany), a goat anti-rabbit IgG, Dylight 800 4xPEG, and goat anti-mouse IgG, Dylight 7004xPEG (Invitrogen) were used. All densitometry analyses were performed by using ImageJ.

**Immunofluorescence.** Cells were seeded in IBIDI chamber at a cell density of 5 × 10⁴ for 24 h. Then, cells were washed with PBS, fixed in 4% (w/v) PFA for 10 min, permeabilized with 0.1% (w/v) Triton X-100 in PBS for 5 min, and blocked with 5% (w/v) BSA in PBS for 1 h at RT. Cells were then incubated with primary antibodies for SRB1 (1:1000, #NB400-104, Novus), CD36 (1:1000, #NB400-145, Novus), and CD81 (1:100, #sc-166029, Santa Cruz) diluted in 1% (w/v) BSA in PBS overnight at 4 °C in a humidity chamber. Thereafter, cells were washed four times with PBS and incubated with the corresponding secondary antibody for 1 h at RT (Dylight donkey anti-mouse 647#406629, Dylight donkey anti-rabbit 667#406410). Nuclei were counterstained by Hoechst 33342 (Tocris #5117) for 10 min. Cells were imaged at 63× using Leica TCS SP8 confocal microscope (Leica Microsystems) and analyzed with ImageJ software (ver. 2.0).

**FACS Analysis.** Polymersome uptake was analyzed by flow cytometry in different cell lines. In brief, cells were plated at 2.5 × 10⁵ in a 6 well plate for 24 h and then incubated with RhofPMPC-PDPA polymersomes for different incubation periods (10 min to 2 h). After PBS washing and centrifugation, cell pellets were resuspended in 2% (v/v) PFA in PBS and analyzed by FACS Fortessa (BD). Acquired data were analyzed using flowJo software.

**Proximity Ligation Assay.** PLA assay uses specific antibodies for two proteins of interest that are recognized by secondary antibodies conjugated with DNA primers. Upon proximity-mediated hybridization, these secondary antibodies produce a fluorescent signal that can be imaged and quantified. However, hybridization can only happen if the proteins of interest are ∼20 nm; hence any fluorescent signal is the result of a close proximity of the two proteins of interest. FaDu cells were plated in IBIDI u-slide (#80826, IBIDI) and incubated for 1 h with Cy3-PMPC-PDPA Psomes. After that, cells were washed with PBS and fixed with 3.7% PFA for 10 min. Scavenger receptors (SRB1 and CD36) and CD81 were labeled by primary antibodies (anti-SRB1, Novus Biologica; anti-CD36, Abcam; anti-CD81, Santa Cruz). The proximity ligation assay was performed using Duolink in situ kit (Sigma) according to the manufacturer’s instructions. Images were randomly collected by confocal microscopy. PLA data were quantitatively analyzed using a Python script based on Trackpy, modified for...
identification of particles with high polydispersity in the direction of objective translation ("z"). A 3-pixel median filter was applied to remove salt and pepper noise, and a low-pass Gaussian filter was applied to remove large-scale features present due to channel crosstalk and optical aberrations. Local maxima were identified and linked into single particles by hierarchical clustering using the Nearest Point Algorithm as implemented in scipy. Data were reported as a number of detectable PLA events ("dots") per nucleus.

**Lectin Binding by FACs.** The DC2.4, THP1-macrophages, FaDu, and HDF cells were incubated with different concentrations of fluorescein isothiocyanate (FITC)-labeled lectin from *Lycopersicon esculentum* (Sigma L0401) for 30 min at 37 °C. After that, cells were washed with PBS twice and centrifuged for 5 min at 1200 rpm and the cell pellet resuspended in 2% PFA/PBS. The specific binding of lectin in different cells was assessed by FACs.

**Molecular Modeling.** CD36 (gene CD36, UniProtKB - P16671 (CD36_HUMAN)) and SRB1 (gene SCARB1, UniProtKB - Q8WTV0 (SCRB1_HUMAN)) structures were constructed using the 3.0 Å resolution crystal structure of the lysosomal domain of limp-2 (PDBid 4F7B) as a template in the Robetta web server (https://rosetta.bakerlab.org). For CD81 (gene CD81, UniProtKB - P60033 (CD81_HUMAN)), we used the available 2.96 Å resolution crystal structure, PDBid STCX, obtained from the PDB databank PDBids 4F7B and STCX. SRB1 and CD36 were glycosylated using the glycoprotein server Glycam, high-manose complex glycans (DNeup5Aca2−6DGlcNAcb1−2DManpa1−6[DNeup5Aca2−6DGlcNAcb1−2DManpa1−3]-[DManpa1−4DGlcpNAcb1−4DGlcpNAcb1−OME]). The poly-(2-methacryloyloxyethyl phosphorylcholine) monomer partial atomic charges were evaluated according to the RESP approach:49 the molecule was first optimized at the HF/6-31G(d) level, up to a convergence in energy of 10−5 au, using the Gaussian09 package.56 Atomic RESP charges were derived from the electrostatic potential using the antechamber module of the AMBER package as well as GAFF parameters.51,52 Different polymerization degree PMPC molecules were constructed and minimized using Amber20 and AmberTools20.53 For all three receptors and different polymerization degree PMPC molecules, we performed docking experiments with a fixed grid of 40 × 40 × 40 centered in the center of mass of the receptor, except for CD81 for which only the solvent-exposed region was considered. Docking calculations were performed with Autodock-vina.1.1.248 with default parameters. One hundred models were generated for each receptor/PMPC-N (N = 1, 2, 3, 4, 5, 6, 8, 10, 15, 25), 3000 models in total. For the best pose, 2D-interaction maps were calculated using LigPlot +.55 The docking maximum affinity poses were simulated using molecular dynamics. CD36 and SRB1 in complex with PMPC (NPM = 5) were simulated in a 150 mM KCl solution using charmm36m56 in GROMACS2019.3.57 CD81 was simulated in a plasma membrane containing cholesterol, phosphoglycerides (PS, PE, PC, PI), sphingomyelin (SM), and glycolipids (GM) in a 150 mM KCl solution using charmm3655 and GROMACS2019.3.57 The membrane systems were created using the membrane builder of CHARMM-GUI.58 Molecular modeling figures, root-mean-square deviations, and angles were measured/created using the visual molecular dynamics suite.59

**In Vivo (Mice) Biodistribution of Polymersomes in Bloodstream.** Three-month-old male C57/BL6 mice were intravenously (i.v.) injected via the tail vein with 10 mg/kg rhodamine labeled PMPC-PDPA polymersomes (mice = 5 per group). Control mice were i.v. injected with saline. The volume of solution injected was 8% of the total blood volume (TBV). TBV was calculated as 58.5 mL of blood per kg of body weight. At 0.16, 0.5, 1, 2, 4, 6, 24, and 168 h post-injection, the mice were terminally anaesthetized, and blood samples were collected through cardiac puncture. The plasma concentration of Psomes was measured after centrifugation of the whole blood at different time intervals. To determine the interactions of fluorescein-labeled Psomes with different types of mouse blood cells, such as lymphocytes, monocytes, granulocytes, and red blood cells, we separated the different fractions using an untouched neutrophil isolation kit, Murine Peripheral Blood Neutrophil Isolation – Easysep kit and Easyplate magnet.

All procedures involving animals were approved by and conformed to the guidelines of the Institutional Animal Care Committee of The University of Sheffield, University College London, and University of Ghent. We have taken great efforts to reduce the number of animals used in these studies and also taken effort to reduce animal suffering from pain and discomfort.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscentsci.2c00146.

Additional figures supporting the molecular modeling. Experimental data on Psome uptake kinetics on FaDu cells. Additional experimental details on polymersome characterization, including transmission electron microscopy images and dynamic light scattering (DLS). Photographs of experimental setup (PDF)

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S.A.G. designed and performed the in silico research, analyzed the data, and wrote and edited the manuscript. D.M. performed part of the in vitro data and analysis; wrote and edited the manuscript. M.A.O. performed some of the in vitro experiments. V.M.G. produced the polymersomes. E.S. performed part of the in vitro experiments. J.F. performed the P.L.A. analysis. C.C. and A.D.C. synthesized the PDPA–PMPC polymer. L.R. designed research and performed some in vitro and in vivo experiments. G.B. designed and conceived the research, and the statistical model, analyzed the data, and wrote and edited the manuscript.

Notes
The authors declare no competing financial interest.

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REFERENCES
(22) Fredriksson, S.; Gullberg, M.; Jarvis, J.; Olsson, C.; Pietras, K.; Gustafsdottir, S. M.; Ostman, A.; Landegren, U. Protein detection using


