Surface Functionalization of Single Superparamagnetic Iron Oxide Nanoparticles for Targeted Magnetic Resonance Imaging

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Magnetic resonance imaging (MRI), a non-invasive, non-radiative technique, is thought to lead to cellular or even molecular resolution if optimized targeted MR contrast agents are introduced. This would allow diagnosing progressive diseases in early stages. Here, it is shown that the high binding affinity of poly(ethylene glycol)-gallol (PEG-gallol) allows freeze drying and re-dispersion of 9 ± 2-nm iron oxide cores individually stabilized with ≈9-nm-thick stealth coatings, yielding particle stability for at least 20 months. Particle size, stability, and magnetic properties of PEGylated particles are compared to Feridex, a commercially available untargeted negative MR contrast agent. Biotin-PEG(3400)-gallol/methoxy-PEG(550)-gallol stabilized nanoparticles are further functionalized with biotinylated human anti-VCAM-1 antibodies using the biotin–neutravidin linkage. Binding kinetics and excellent specificity of these nanoparticles are demonstrated using quartz crystal microbalance with dissipation monitoring (QCM-D). These MR contrast agents can be functionalized with any biotinylated ligand at controlled ligand surface density, rendering them a versatile research tool.

1. Introduction

The good biocompatibility and low cost of iron oxide nanoparticles render them suitable for many biomedical applications[1,2] such as cell separation, ferrying cells, and contrast-enhanced MRI.[3] Much work has been dedicated to the synthesis of iron oxide nanoparticles because the synthesis determines particle composition, size distribution, and morphology.[4,5] These are parameters directly influencing physical properties[6,7] as well as clearance rate and clearance route of nanoparticles.[8,9]

Several complementary non-invasive techniques for biomedical imaging, for example, MRI, single-photon emission tomography (SPECT), positron emission tomography (PET), and computed tomography (CT), are used to elucidate the evolution and progression of many different diseases in humans and animals. One important advantage of MRI over SPECT, PET, and CT is its non-radiating nature. Moreover, MRI allows for imaging soft tissues in contrast to PET and SPECT, and it does so with higher contrast resolution than CT. The inherent low sensitivity of MRI compared to SPECT and PET can be improved by introducing MR contrast agents.
without other measures such as increasing field strength.\textsuperscript{[10]} Moreover, targeted MR contrast agents are thought to enable visualization of specific cell types or even molecules.\textsuperscript{[11–13]} Several positive and negative MR contrast agents are already FDA approved and routinely used in clinics. Commercially available negative MR contrast agents are mostly stabilized with biocompatible macromolecules such as dextran, which reduce toxicity and enhance particle stability. However, encapsulation of multiple iron oxide cores in one cluster by macromolecular dispersants leads to a hydrodynamic diameter many times larger than the core diameter and a broad size distribution.\textsuperscript{[14,15]}

Moreover, reversible adsorption of such polymers hampers efficient particle functionalization due to constant changes in macromolecule conformation, including desorption, which results in a poorly defined interface and prevents controlled presentation of ligands. Enhanced particle stability and a better control over the particle size distribution have been reported for dextran stabilized iron oxide nanoparticles where dextran was crosslinked after its adsorption.\textsuperscript{[13,20]} Nevertheless, the interfacial chemistry of such stabilized particles is difficult to control.

Alternatively, iron oxide nanoparticles have been stabilized with low-molecular-weight dispersants that have a well-defined, high-affinity anchor group such as dopamine.\textsuperscript{[17–25]} These dispersants increase particle stability and result in a substantially decreased hydrodynamic diameter due to a thinner, better defined, stabilizing layer surrounding individual particles. However, long-term stability under physiological conditions or long-term storage capability for iron oxide nanoparticles stabilized with dopamine-based dispersants have not yet been reported.

In this work, iron oxide nanoparticles were synthesized by aqueous precipitation.\textsuperscript{[24]} These nanoparticles were stabilized using spontaneous self-assembly from aqueous solution of either methoxy-PEG(550)-gallol (mPEG(550)-gallol), methoxy-PEG(5000)-6-hydroxy-dopamine (mPEG(5000)-gallol), or dopamine and mPEG(550)-gallol, or mixtures thereof. These anchor groups are less prone to oxidation than dopamine and DOPA.\textsuperscript{[20]} The high binding affinity of the gallol and trihydroxy-benzene units toward iron oxide allowed freeze drying and successful re-dispersion of individual particles. Particle size and magnetic properties were compared to Feridex, a commercially available iron oxide-based MR contrast agent. To prepare functionalized nanoparticles, iron oxide nanoparticles were first stabilized with a mixture of mPEG(550)-gallol and biotin-PEG(3400)-gallol. Biotinylated anti-human vascular cell adhesion molecule 1 (VCAM-1) antibodies were bound to neutravidin, a derivative of avidin, which shows lower non-specific adsorption.\textsuperscript{[27]} VCAM-1 is considered to be a well-suited target for MR contrast agents because it is up-regulated at the endothelial cell periphery of lesion prone sites and, thus, known to be an early marker of atherosclerosis.\textsuperscript{[26–30]} The average number of biotinylated dispersants, neutravidin, and biotinylated ligands bound to one iron oxide nanoparticle could be estimated due to the well-defined build-up of the nanoparticles.

The fast, and under physiological conditions easily established, layer-by-layer build-up of these individually stabilized MR contrast agents having a non-fouling PEG-shell with a controlled number of functional groups for presentation of modularly interchangeable active ligands has, to the best of our knowledge, not been demonstrated before.

## 2. Results

### 2.1. Nanoparticle Stabilization and Characterization

A comparison of the hydrodynamic diameter of PEG-gallol stabilized particles, schematically shown in Figure 1, and Feridex revealed a considerably smaller hydrodynamic diameter and narrower size distribution for the former particles. Moreover, PEG-gallol stabilized iron oxide nanoparticles are stable even after 20 months’ storage at room temperature in 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) containing 150 mM NaCl (Figure 2). Feridex consists of multiple iron oxide cores with an average diameter of 5 nm embedded in a dextran matrix,\textsuperscript{[30]} which results in a hydrodynamic diameter of 63 ± 21 nm. However, particles stabilized with a mixture of mPEG(550)-gallol and biotin-PEG(3400)-gallol have a hydrodynamic diameter of 27.0 ± 2 nm (Figure 2a), which is well in agreement with X-ray disk centrifuge (XDC) measurements that gave a hydrodynamic diameter of 25.7 ± 2.1 nm. Furthermore, Figure 2b indicates that particles are well separated even if dried on a carbon film. Moreover, freeze-dried PEG-gallol stabilized particles could also be re-dispersed in Dulbecco’s phosphate-buffered saline (PBS) and stored for more than 12 months without changes in the hydrodynamic diameter (Figure 2a).

Figure 3a exemplifies the high particle stability of biotin-PEG(3400)-gallol/mPEG(550)-gallol stabilized iron oxide nanoparticles even after storing them for more than 1 year in PBS at room temperature. The individually stabilized nanoparticles stay in solution even in the presence of an external magnetic field. This stands in strong contrast to unstabilized iron oxide nanoparticles, which agglomerate and sediment within seconds after applying a small external magnetic field as can be seen in Figure 3b (also see video in Supporting Information). While the net magnetic moment of
magnetite and maghemite. The BET surface area of...good agreement with what has been previously reported for...stabilization in HEPES (○), mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles stored as powder for 1 year, re-dispersed in PBS for 1 year (●) and Feridex (△). b) TEM image showing mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles dried onto an 8-nm-thick carbon film. The nanoparticles are well separated by the PEG shell.

Figure 2. a) Number-weighted DLS measurement of mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles kept as powder for 20 months before re-dispersion in HEPES (▲), mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles 20 months after stabilization in HEPES (●), mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles stored as powder for 1 year, re-dispersed in PBS for 1 year (●) and Feridex (△). b) TEM image showing mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized nanoparticles dried onto an 8-nm-thick carbon film. The nanoparticles are well separated by the PEG shell. (See also videos in the Supporting Information.)

The single-domain core of individually PEG-gallol stabilized iron oxide nanoparticles had a diameter of 9 ± 2 nm, as determined by transmission electron microscopy (TEM; Figure 4). High-resolution TEM (HRTEM) micrographs of single-domain iron oxide nanoparticles oriented in the (221) direction revealed an Fe–Fe distance of 3.0 ± 0.3 Å (Figure 4a) in good agreement with what has been previously reported for magnetite[32] and maghemite.[33] The BET surface area of uncoated powder was 88 m² g⁻¹. This area was lower than expected for the average particle diameter of 9 nm; however, during centrifugation of freeze-dried and re-dispersed particles, larger particles were removed and thus excluded from the core size evaluations of re-dispersed particles performed by TEM.

Thermogravimetric analysis coupled to a Fourier transform infrared spectrometer (TGA-FTIR) was performed to validate the presence and quantify the amount of PEG-gallol on the nanoparticle surface. The weight loss measured between 200 and 400 °C during TG analysis could be clearly assigned to the decomposition of PEG by FTIR spectroscopy, proving that PEG-gallol was adsorbed on the nanoparticles. The weight loss between 200 and 400 °C measured by TG analysis for mPEG(550)-gallol, mPEG(550)-gallol/biotin-PEG(3400)-gallol, and mPEG(5000)-gallol stabilized nanoparticles was 11, 15, and 50 wt%, respectively. Based on these values, approximately 235 molecules were adsorbed per particle (194 – 198 μmol·g⁻¹). This corresponds to an average contact area of 74–76 Å² per molecule for all dispersants, independent of PEG molecular weight. For mPEG(5000)-stabilized particles, the corresponding PEG density is 1.3 PEG(5000) molecules nm⁻², which is more than three times higher than the maximum PEG(5000) density reported (0.4 PEG(5000) nm⁻²) for mPEG(5000)-DOPA₃ adsorbed onto flat TiO₂ surfaces.[34] The molar ratio of biotin-PEG(3400)-gallol to mPEG(550)-gallol could thus be calculated by comparing the difference in mass loss measured with TGA between mPEG(550)-gallol and mPEG(550)-gallol/biotin-PEG(3400)-gallol stabilized particles assuming that the dispersant packing is independent on the PEG molecular weight. Based on these calculations, the dispersant shell surrounding the iron oxide consisted of 9 mol% biotin-PEG(3400)-gallol and 91 mol% mPEG(550)-gallol. The iron oxide cores had been stabilized by adding 10 mol% biotin-PEG(3400)-gallol and 90% mPEG(550)-gallol to the uncoated nanoparticle suspension, which is close to the percentage of these dispersants absorbed on the particles. Assuming particles were perfectly spherical and had a core diameter of 9 nm this equaled 20 biotin sites per particle.

Zeta potential measurements revealed a point of zero charge (PZC) of 6.7 for uncoated iron oxide nanoparticles, in good agreement with what other groups have reported.[35–37] The isoelectric point (IEP) of nanoparticles stabilized with...
mPEG(550)-gallol and dialyzed for 24 h against Millipore water, was shifted to 4.2 (Figure 5). The difference between the PZC of unstabilized and the IEP of mPEG(550)-gallol stabilized nanoparticles was assumed to result from the adsorption of gallol, as was reported by Studart et al. for Al₂O₃ nanoparticles. The protons from gallol likely dissociated prior to binding to iron oxide introducing negative charges to the surface and thus shifting the IEP toward more acidic values.

2.2. Magnetic Properties

The superparamagnetic properties of these single-domain iron oxide nanoparticles were confirmed with superconducting quantum interference device (SQUID) measurements (Figure 6a). The room-temperature saturation magnetizations of unstabilized and mPEG(550)-gallol stabilized particles was 58 and 50 emu⁻¹ g⁻¹, respectively, which were within the ranges of reported values. A reduction of the saturation magnetization upon iron oxide nanoparticle stabilization with different dispersants has also been described by other groups. Based on the observed decrease, the magnetization of the top of ~2 Å is likely to have been lost. R₂ and the ratio R₂/R₁ are parameters for contrast enhancement of conventional T₂-weighted MR images typically measured when negative contrast agents such as iron oxide nanoparticles are used. R₂ of iron oxide nanoparticles stabilized with mPEG(550)-gallol and biotin-PEG(3400)-gallol was higher (134 ± 91 mmolFe⁻¹ g⁻¹ s⁻¹) than R₂ of Feridex (102 ± 51 mmolFe⁻¹ g⁻¹ s⁻¹), measured at T=39 °C and B₀=11.7 T. Furthermore, R₁ was substantially lower for the former (1.4 ± 0.2 mmolFe⁻¹ g⁻¹ s⁻¹) compared to R₁ for Feridex (2.7 ± 0.2 mmolFe⁻¹ g⁻¹ s⁻¹) (Figure 6b and c) resulting in a more than two times higher R₂/R₁ ratio of PEG-gallol stabilized particles compared to Feridex.

2.3. Nanoparticle Functionalization

Neutravidin was bound to biotin-PEG(3400)-gallol/mPEG(550)-gallol stabilized particles (Figure 7). The number of neutravidins adsorbed per particle was maximized to prevent aggregation of different particles, which can bind to the same neutravidin since neutravidin has four biotin binding sites. The neutravidin saturation concentration of these biotinylated particles was determined with a) DLS (Figure 8a), b) fluorescence-activated cell sorting (FACS; Figure 8b), and c) QCM-D (Figure 8c) by adding different amounts of neutravidin to PBS containing 1 mg mL⁻¹ biotin-PEG(3400)-gallol/mPEG(550)-gallol stabilized iron oxide nanoparticles. The hydrodynamic diameter of neutravidin-coated particles increased with increasing neutravidin concentration up to a protein concentration of 7 μmol neutravidin g⁻¹ iron oxide yielding an increase in hydrodynamic diameter of 12 nm (Figure 8a). Streptavidin, another avidin derivative of similar molecular weight as neutravidin, has a thickness of ~5 nm. The observed
increase in hydrodynamic diameter of close to twice the neutravidin thickness thus agrees well with particles surrounded by a neutravidin monolayer. The increased fluorescence signal of nanoparticles with increasing addition of FITC-labeled neutravidin up to a concentration of 7 μmol g\textsuperscript{-1}iron oxide further supported the findings from DLS (Figure 8b). Lastly, SiO\textsubscript{2}-coated QCM-D sensors were used to form a neutravidin monolayer on top of a non-fouling supported lipid bilayer (SLB) containing biotinylated lipids. Particle binding to the neutravidin monolayer decreased as the surface coverage of neutravidin on the particles increased. No binding was observed at 7 μmol g\textsuperscript{-1}iron oxide (Figure 8c), which corresponds to eight neutravidin molecules per particle assuming an iron oxide density of 5.18 g cm\textsuperscript{-3}. If one considers that at least two biotin binding sites per neutravidin are accessible this would correspond to close to full saturation of the average 20 biotin binding sites per particle estimated from our TGA-FTIR results. Based on these results, it is unlikely that any free biotin site was still accessible after saturating the particle shell with neutravidin. This was supported by the observed plateau in both the DLS and FACS curves, indicating that there was no particle aggregation.

Real-time binding kinetics and binding specificity of functionalized particles were investigated using QCM-D. Recombinant human VCAM-1 chimera was immobilized via protein A onto SiO\textsubscript{2} QCM-D sensor crystals (schematically shown in Figure 9a). Neutravidin pre-coated nanoparticles were further functionalized with biotinylated human anti-VCAM-1 antibodies. Upon exposure of such functionalized nanoparticles to the sensor, fast, and specific binding was observed. Only minimal frequency and dissipation shifts (−1.5 Hz and 1.2 × 10\textsuperscript{-6}, respectively) were measured 2 h after particle injection for particles lacking the anti-VCAM-1 antibody. Corresponding shifts of −30 Hz and 18.7 × 10\textsuperscript{-6} for particles functionalized with biotinylated human anti-VCAM-1 antibody demonstrated that the binding was specific (Figure 9b). Neither neutravidin pre-coated particles bearing biotinylated anti-E-selectin antibodies nor PEGylated iron oxide particles or bovine serum albumin (BSA) adsorbed

Figure 7. Negatively stained TEM image of a single, mPEG(550)-galoll/biotin-PEG(3400)-galoll-stabilized Fe\textsubscript{3}O\textsubscript{4} particle further functionalized with a saturated neutravidin layer (white shapes surrounding particle).

Figure 8. The neutravidin saturation concentration of biotin-PEG(3400)-galoll/mPEG(550)-galoll-stabilized iron oxide nanoparticles was determined by a) DLS where an increase in hydrodynamic diameter of particles bearing increasing amounts of neutravidin was measured up to the neutravidin saturation concentration, b) FACS measurements revealing an increase in fluorescence of particles coated with increasing amounts of FITC-labeled neutravidin up to the neutravidin saturation concentration, and c) QCM-D measurements demonstrating particle binding toward a neutravidin monolayer. Binding decreased to 0 with increasing amount of neutravidin bound to the biotinylated particles. The neutravidin monolayer was immobilized onto SiO\textsubscript{2}-coated sensors through non-fouling SLBs containing biotinylated lipids. All three techniques showed that maximally ≈ 7 μmol neutravidin g\textsuperscript{-1}iron oxide can be bound to the biotin-bearing nanoparticles, corresponding to eight neutravidin molecules per particle.
protein A on SiO₂-coated QCM-D crystals (functionalized particles toward E-cadherins which had previously been immobilized through functionalized particles.

These experiments demonstrated high binding specificity of anti-VCAM-1 antibody (-)

c) Binding of PEG-gallol stabilized neutravidin-coated particles bearing no antibodies (Δf = -30 Hz) (○) to recombinant VCAM-1 chimeras was considerably lower compared to anti-VCAM-1 antibodies functionalized particles (Δf = -3 Hz) (●). No significant binding of anti-VCAM-1 antibody functionalized particles toward E-cadherins which had previously been immobilized through protein A on SiO₂-coated QCM-D crystals (Δf = 0 Hz) (○) or human serum immobilized on Au-coated QCM-D crystals (Δf = -0 Hz) prior to particle injection (○) was measured. Taken together, these experiments demonstrated high binding specificity of anti-VCAM-1 antibody functionalized particles.

Figure 9. a) Schematic of the immobilization of recombinant human VCAM-1 chimera onto SiO₂ QCM-D crystals via protein A and the layer-by-layer build-up of functionalized Fe₃O₄ nanoparticles. These nanoparticles were stabilized with a mixture of mPEG(550)-gallol and biotin-PEG(3400)-gallol, further coated with 7 μmol neutravidin g⁻¹ oxide, and subsequently functionalized with 14 μmol human E-selectin antibodies.

b) Whereas non-specific binding of neutravidin-coated nanoparticles (SPIONs) lacking anti-VCAM-1 antibodies was negligible as shown in QCM-D experiments by small changes in frequency (△) and dissipation (△), there was strong and fast binding of SPIONs functionalized with anti-VCAM-1 antibodies to recombinant VCAM-1 chimeras, indicated by a large frequency (∼30 Hz) (●) and dissipation (1.3 × 10⁻⁶) (◆) shift after 2 h exposure.

c) Binding of PEG-gallol stabilized neutravidin-coated particles bearing no antibodies (Δf = -2 Hz) (△) and particles functionalized with human E-selectin antibodies (Δf = -3 Hz) (◆) to VCAM-1 chimeras was considerably lower compared to anti-VCAM-1 antibodies functionalized particles (Δf = -30 Hz) (●). No significant binding of anti-VCAM-1 antibody functionalized particles toward E-cadherins which had previously been immobilized through protein A on SiO₂-coated QCM-D crystals (Δf = 0 Hz) (○) or human serum immobilized on Au-coated QCM-D crystals (Δf = 0 Hz) prior to particle injection (○) was measured. Taken together, these experiments demonstrated high binding specificity of anti-VCAM-1 antibody functionalized particles.

3. Discussion

The high binding affinity of PEG-gallol toward iron oxide particles was demonstrated in that particles functionalized with anti-VCAM-1 antibodies did not bind to cadherins nor to human serum (Figure 9c and S1 in Supporting Information). However, binding of functionalized particles was highly dependent on the amount of anti-VCAM-1 antibodies added to neutravidin precoated nanoparticles (see Figure S2 in Supporting Information). Further control experiments of the binding of biotinylated anti-VCAM-1 antibodies with and without neutravidin to recombinant VCAM-1 chimera and binding of functionalized particles toward protein A are also given in the Supporting Information (Figure S1).

4. Conclusions

We have shown that a layer-by-layer build-up of successive shells of superparamagnetic iron oxide nanoparticles individually stabilized with a mixture of mPEG(550)-gallol and PEG(3400)-gallol leads to high particle stability under the additional possibility to freeze-dry and re-disperse individually PEG-gallol stabilized iron oxide nanoparticles. This simplifies particle handling and allows for long-term particle storage, which is of high commercial value. In contrast to what Carpenter and co-workers [45] and Xie et al. [46] have reported for core/shell Fe₃O₂ and Fe₃O₄ nanoparticles stabilized with PEG(600)-dopamine and PEG(2000)-trichloro-s-triazine-dopamine, respectively, no degradation or corrosion of iron oxide nanoparticles upon stabilization was observed with PEG-gallol.

The much lower standard deviation of the hydrodynamic diameter of mPEG-gallol-stabilized nanoparticles compared to Feridex (7% versus 33%) is a result of the high binding affinity of gallol and trihydroxy-benzene, which leads to a well-defined surface chemistry and no aggregation. The weight percent of dispersants adsorbed on nanoparticles related to the BET surface area results in a dispersant contact area of 75 Å². This corresponds to a dispersant packing density of 30% if one refers to the reported theoretical cross-sectional anchor group area perpendicular to the benzyl ring of 22.3 Å². [46] The PEG layer of iron oxide nanoparticles stabilized with biotin-PEG(3400)-gallol/mPEG(550)-gallol is approximately 9 nm thick, if the iron oxide core diameter determined by TEM is compared to the hydrodynamic diameter measured with DLS and XDC. With a single iron oxide core diameter defined only by the gallol anchor, the particle size distribution is determined solely by the size distribution of the core and the choice of PEG molecular weight.
physiological conditions. Particles could be freeze-dried and re-dispersed without noticeable corrosion or aggregation. Furthermore, they could be functionalized by the addition of a controlled number of neutravidin and biotinylated antibodies. Nanoparticles functionalized with human anti-VCAM-1 antibodies showed high binding affinity and specificity toward recombinant human VCAM-1 chimeras, whereas non-specific binding of neutravidin-coated nanoparticles was negligible, suggesting suitability for in vivo imaging of atherosclerosis.

Importantly, any biotinylated ligand can be used to functionalize these MR contrast agents, which allows for targeting of any receptor that has a known ligand. Furthermore, comparison of immobilized ligand binding efficiencies can easily be accomplished by binding different biotinylated ligands to the neutravidin-decorated nanoparticles and performing in vitro binding studies. The particles can be produced with variable and controlled ligand densities. This and their ease to prepare and handle render them particularly well suited for research purposes. Finally, the family of catechol-derived anchors has been shown to strongly bind to a large variety of substrate types in particular metal oxides.

Thus, the surface modification concept presented in this work is a promising platform for the specific functionalization of a wide range of micro- and nanoparticle types.

5. Experimental Section

Materials: Biotin-PEG(3400)-NHS was purchased from Nektar, 3,4,5-Trihydroxyphenethylamine hydrochloride from Acros Organics, L-α-Phosphatidylcholine (egg-PC) and 1,2-Dipalmitoyl-sn-Glycero-3-Phosphoethanolamine-N-(Cap Biotinyl) (biodo-PE) from Avanti Polar Lipids, protein A, neutravidin, BSA and PBS (containing 2.67 mM KCl, 1.47 mM KH2PO4, 137.93 mM NaCl and 8.5 mM Na2HPO4) from Invtrogen, Feridex from Berlex, human serum Percinorm U from Roche Systems, recombinant human VCAM-1 chimera and biotinylated human anti-E-selectin antibodies from R&D systems, biotinylated anti-human CD106 (VCAM-1) from Lubio science and all the other chemicals from Fluka. Fc-tagged human E-cadherin was a kind gift from Prof. Deborah E. Leckband.

Dispersant synthesis: mPEG(550)-gallol synthesized according to Studart et al. was a kind gift from André Studart. mPEG(5000)-gallol and biotin-PEG(3400)-gallol were synthesized following the same protocol as for mPEG-dopamine but using 6-hydroxy-dopamine instead of dopamine and mPEG(5000)-NHS respectively biotin-PEG(3400)-NHS. The average molecular weight of the dispersants was estimated based on matrix assisted laser desorption/ionization time of flight (MALDI-ToF) and nuclear magnetic resonance (NMR) spectra.

Particle synthesis: Iron oxide nanoparticles were synthesized by aqueous precipitation according to the protocol of Massart. Particles were neutralized by washing them with 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) solution containing 150 mM NaCl. 24 μmol of PEG(550)-gallol or PEG(5000)-gallol dissolved in 25 mL Millipore water (18.2 MΩ) was added to a typical batch size resulting from 198.8 mg (1.2 mmol) FeCl2 and 540.6 mg (2.3 mmol) FeCl3. Particles were dispersed in this solution by pulsed sonication (5 min, P = 105 W cm−2, pulse frequency = 1 s on, 1 s off, UP260s, Hierschler GmbH). To stabilize particles with a mixture of biotin-PEG(3400)-gallol and mPEG(550)-gallol, 25 mL Millipore-water containing 2.4 μmol of the former dispersant was added to uncoated particles before they were sonicated, followed by the addition of 21.6 μmol mPEG(550)-gallol. Particles were sonicated a second time according to the above described procedure. Excessive dispersants were removed by ultracentrifugation using a filter with a cut-off of 30 kDa (vivaspin, Epson). The filter cake was re-suspended in Millipore water before the suspension was freeze-dried. Particles were re-suspended in PBS, HEPES, or Millipore water, sonicated for 30 min, and centrifuged for 10 min at 25 000 g (Eppendorf centrifuge 5417R) to eliminate agglomerates. The well-dispersed particles in the supernatant were collected and used for further studies.

Vesicle preparation: Egg-PC and biotin–PE were dissolved in chloroform. 0.5–2 wt% biotin–PE was added to egg–PC prior to drying this mixture to thin films under a steady flow of N2 gas. The lipids were re-suspended in PBS at a concentration of 5 mg mL−1. After allowing lipids to swell for 1 h, they were sonicated for 30 min using a bath sonicator (Aquasonics 75T, WVR Scientific products). Such-prepared vesicles were stored at 5°C under N2 for maximally 3 weeks.

Particle characterization: TEM imaging was performed on a Philips CM12 microscope operated at 100 kV and on a HRTEM Philips CM30 operated at 300 kV. Particles were suspended in Millipore-water at 60 μg mL−1 before they were air-dried on 8-nm thick carbon-coated 400-mesh Cu grids. Based on these TEM micrographs, the iron oxide core diameter was measured manually for 240 particles.

DLS experiments were conducted on a Brookhaven instrument 90 Plus particle size analyzer at a 90° angle using 100 μg mL−1 PBS-based particle suspensions. Data were analyzed using a log normal fit. XDC experiments of 16 vol% HEPES-based particle suspensions were conducted on a BI-XDC machine (Brookhaven instruments limited) operated at 6000 rpm for 1 h. N2 adsorption–desorption isotherms were measured at room temperature on a NOVA 1000 series Quantachrome machine. Before N2 adsorption, freeze-dried, uncoated samples (NB. dried nanoparticles are respiratory and thus potentially harmful) were degassed in vacuum at 160°C for 2 h. The specific surface area was calculated using the Brunauer–Emmett–Teller method (BET).

Zeta potential: The zeta potential (DT-1200 dispersion technology, Inc.) was measured on 1 vol% Millipore water-based suspensions. These suspensions had previously been dialyzed against Millipore-water for 24 h. For titrations 1 w KOH and HNO3 were used respectively.

Magnetic properties: SQUID measurements using vacuum grease paste containing 0.03–3.17 wt% freeze-dried iron oxide nanoparticles were performed on a magnetic property measurement system at room temperature. The magnetization was calculated assuming the iron oxide core was Fe3O4. A Bruker Avance 500 Wide Bore spectrometer (500 MHz for protons) fitted with a gradient amplifier (Bruker) for imaging using a fast spin-echo sequence in a 30-mm birdcage transmitter/receiver coil operated at 11.7 T was used for MRI. The slice thickness was 2 mm, field of view = 3.2 cm and the matrix = 64×64 for T1 and
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T₂-weighted images. For T₁ weighted images Tₑ = 6.3 ms, while T₂-weighted images were performed using a Carr-Purcell-Meiboom-Gill sequence\[49\] with Tₑ = 3000 ms and Tₚ = 10 ms. MRI measurements were performed at 39 °C. Relaxivities were calculated based on linear regressions through all data points shown in Figure 6b and c.

Dispersant quantification: The amount of admixed dispersant was quantified using TGA (TGA 500 Q series, TA Instruments) run from 25 to 400 °C at a heating rate of 10 °C min⁻¹ in N₂ atmosphere. Coupled to the TGA was an FTIR spectrometer (Thermo Nicolet Nexus 470 FT-IR) with which spectra of the volatile fractions coming off during TG analysis were measured every 30 s.

Neutravidin quantification: FACScanibur instrument (BD Biosciences). FITC-labeled neutravidin was dissolved in PBS prior to the addition of 10 μg mL⁻¹ biotin-bearing particles. QCM-D studies\[50\] were carried out on a Q-Sense E4 equipment (Q-Sense AB). Adding the PBS-based vesicle solutions in situ to SiO₂-coated quartz crystals (Q-Sense AB) resulted in the formation of biotinylated SLBs. Neutravidin was immobilized on the biotinylated SLBs prior to injection of biotinylated particles pre-coated with different amounts of neutravidin. The change in resonant frequency, Δf, and energy dissipation, ΔD, was measured in real time during the adsorption. The adsorbed mass can be approximated by m = −17.7 × Δf/n (ng cm⁻²) for the 5 MHz crystal used according to the Sauerbrey equation,\[52\] where n is the overtone number.

QCM-D Binding Studies: 23 μg neutravidin was added to 500 μL HEPES containing 150 mM NaCl and 50 μg iron oxide nanoparticles stabilized with mPEG(550)-gallol and biotin-PEG(3400)-gallol (corresponding to 7 μmol neutravidin 560 μl⁻¹ iron oxide). Unless stated otherwise, 7.5 μg biotinylated anti-human VCAM-1 antibodies were added to neutravidin pre-coated particles. QCM-D studies were performed at room temperature in HEPES containing 150 mM NaCl (pH 7.4) under static conditions. A 400 ng cm⁻² protein A was adsorbed on SiO₂-coated QCM-D crystals from 50 μg mL⁻¹ solution followed by the addition of 5 μg mL⁻¹ recombinant human VCAM-1 chimera before 100 μg mL⁻¹ nanoparticles were injected. Between every step the system was rinsed with buffer. For control experiments, recombinant human VCAM-1 chimera was exchanged by Fc-tagged E-cadherins on the sensor substrate. Alternatively, biotinylated anti-human VCAM-1 antibodies were replaced by anti-human E-selectin antibodies on the nanoparticles. The mass concentrations were always kept constant. A concentration of 100 μg mL⁻¹ was used for the control experiment with BSA. Human serum was immobilised on Au-coated QCM-D crystals before functionalized particles, prepared according to the same procedure as described above, were injected into the QCM-D chamber.

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[19] Even though 6-hydroxy-dopamine is not a direct derivative of gallic acid because it has no acidic group on the phenol ring, we call it
gallol for simplicity. It has three hydroxy groups directly linked to the phenol as is the case for gallic acid.


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