Reconstitution of adhesive properties of human platelets in liposomes carrying both recombinant glycoproteins Ia/IIa and Ibα under flow conditions: specific synergy of receptor–ligand interactions

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Liposomes carrying both recombinant glycoprotein Ia/IIa (rGPla/Ila) and Ibα (rGPliba) (rGPla/Ila-Ibα-liposomes) instantaneously and irreversibly adhered to the collagen surface in the presence of soluble von Willebrand factor (VWF) at high shear rates, in marked contrast with translocation of liposomes carrying rGPliba alone on the VWF surface. In the absence of soluble VWF, the adhesion of rGPla/Ila-Ibα-liposomes to the collagen surface decreased with increasing shear rates, similar to liposomes carrying rGPla/Ila alone. While adhesion of liposomes with exofacial rGPla/Ila and rGPliba densities of 2.17 \times 10^{3} and 1.00 \times 10^{4} molecules per particle, respectively, was efficient at high shear rates, reduction in rGPliba density to 5.27 \times 10^{2} molecules per particle resulted in decreased adhesion even in the presence of soluble VWF. A 50% reduction in the exofacial rGPla/Ila density resulted in a marked decrease in the adhesive ability of the liposomes at all shear rates tested. The inhibitory effect of antibody against GPIaα (GUR83-35) on liposome adhesion was greater at higher shear rates. Further, the anti-GPIa antibody (Gi9) inhibited liposome adhesion more than GUR83-35 at all shear rates tested. These results suggest that the rGPla/Ila–collagen interaction dominates the adhesion of rGPla/Ila-Ibα-liposomes to the collagen surface at low shear rates, while the rGPla/Ila–collagen and rGPliba–VWF interaction complements each other, and they synergistically provide the needed functional integration required for liposome adhesion at high shear rates. This study thus has confirmed for the first time the proposed mechanisms of platelet adhesion to the collagen surface under flow conditions using the liposome system. (Blood. 2002;100:136-142)

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Introduction

The basic and important platelet functions for primary hemostasis are adhesion and aggregation, and this can be easily understood from the observations that patients with congenital platelet membrane defects such as Bernard-Soulier syndrome or Glanzmann thrombasthenia are deficient in platelet adhesion or aggregation and have severe bleeding tendencies. The contribution of specific platelet receptors or adhesion proteins to platelet adhesion and aggregation onto immobilized collagen under flow conditions is usually studied with monoclonal antibodies or inhibitors specific to particular platelet receptors or adhesive proteins or, also, with blood from patients with congenital bleeding disorders deficient in specific receptors or adhesive proteins. These analyses indicate that initial platelet adhesion depends on the interaction of glycoprotein (GP) Ib/IX/V complexes on platelets with von Willebrand factor (VWF) adsorbed on the collagen surface. This is a rapid but low-affinity interaction, suggesting that it serves to tether platelets, flowing at high speed in the bloodstream, to the collagen surface.1-4 The collagen receptors of the tethered platelets then bind strongly with the collagen surface, activating platelets to form aggregates. This was supported by observations that platelets deficient in one of the collagen receptors failed to adhere and form aggregates on subendothelium or the collagen surface under flow conditions.5-6 GPIa/Ila and GPVI are known to be involved in platelet adhesion under static conditions.6,7 GPIa/Ila (integrin α_{II}β_{3}), VLA2, CD49b/29) is a member of the integrin family of heterodimeric molecules that mediate both cell-to-cell adhesion and adhesion between cells and the extracellular matrix.8 GPIa/Ila is also a major collagen receptor in platelets.9-11 Although GPIa/Ila-mediated adhesion appears to be an essential primary step in collagen–platelet interactions, the functional integration of the distinct adhesion pathways involved in the initiation of platelet adhesion has not yet been defined. To address this issue, we prepared liposomes with covalently bound recombinant GPIa/Ila (rGPla/Ila)12 and/or recombinant fragments of GPIbα consisting of residues 1 to 302 (rGPliba)13 and evaluated their interaction with the collagen or VWF surface under flow conditions in the absence of other platelet components. Previously, we reported that liposomes carrying rGPliba (rGPliba-liposomes) reversibly interact with the VWF surface under flow conditions, depending on the shear rate and the densities of receptor and matrix, and the interaction is directly related to shear rate.14 The purpose of the present study was to examine how GPIa/Ila and GPIbα contribute to platelet adhesion to the collagen surface under flow conditions in an in vitro reconstituted system, using liposomes carrying both rGPla/Ila and rGPliba (rGPla/Ila-Ibα-liposomes). Our results suggest that rGPla/Ila and rGPliba reconstituted into liposomes retain hemostatic functions under flow conditions in vitro, and direct interaction of rGPla/Ila and GPIbα is required for efficient adhesion at high shear rates.
with the collagen surface dominates the adhesion of GPIa/IIa-Ibo-liposomes to the collagen surface at low shear rates. At high shear rates, tethering of liposomes through the interaction between rGPIbα and the VWF-adsorbed collagen surface reduces the velocity of liposomes, enabling binding of rGPIa/IIa to the collagen surface. This is the first study to prove the proposed mechanisms of platelet adhesion to the collagen surface involving 2 distinct receptor–ligand pairs with unique properties, GPIa/IIa–collagen and GPIbα–VWF, using reconstituted system, rGPIa/IIa-Ibo-liposomes.

Materials and methods

Materials

N-glutaryl-phosphatidylethanolamine (NGPE), egg phosphatidylcholine (EPC), and N-lysamine rhodamine B sulfonyl phosphatidylethanolamine (N-Rh-PE) were purchased from Avanti (Birmingham, AL). Cholesterol (CHO), 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDCI), n-octyl-β-glucopyranoside (OG), bovine serum albumin (BSA), HEPS, and MES were obtained from Sigma Chemical (St Louis, MO). N-hydroxysulfo succinimidic acid (NHS) was obtained from Pierce Chemical (Rockford, IL). Mouse monoclonal antibody against GPIbα (purified immunoglobulin G [IgG], GUR83-35, was made against crude glycocalicin fraction extracted from washed human platelets. A mouse anti-GPIa monoclonal antibody, G9, and a mouse anti-GPIa monoclonal antibody, Lia1/2, were purchased from Immunotech (Marseille, France). Sephadex G-25 and Sephadex G-75 were obtained from Pharmacia Biotech (Uppsala, Sweden). The phospholipid-test Wako was from Wako (Osaka, Japan). Nonidet P-40 was obtained from Nacalai Tesque (Kyoto, Japan). Expression and purification of rGPIbα containing the VWF binding site (residues 1 to 302) were performed as described by Murata et al.13 Specific binding of VWF to rGPIbα was assayed by measuring the 125I-VWF binding to rGPIbα.13 Preparation of the extracellular domain of rGPIa/IIa in which α2 and β1 chains were covalently bound by disulfide bond was performed according to the following method. Thus, DNA fragments encoding the extracellular domain of GPIa16 and GPIb17 were amplified by polymerase chain reaction using template complementary DNA obtained from human fibroblast cell line MRC-5 (ATCC CCL 171) and primers. Polymerase chain reaction products were subcloned into the pBlueScript polymerase chain reaction using template complementary DNA obtained from human fibroblast cell line MRC-5 (ATCC CCL 171) and primers. Polymerase chain reaction products were subcloned into the pBlueScript (Stratagene, La Jolla, CA), and then the fragments of resultant plasmid were introduced into the expression vector pCDSLra,18 respectively. One milligram each of the expression plasmids was mixed with 0.1 mg each of pSV2dhfr (Gibco, New York, NY) and pSV2neo (Gibco), and the mixture was introduced into dihydrofolate reductase–deficient CHO cells (ATCC CRL 9096) using lipofection reagent (Gibco). Then the cells were cultured in the nucelic acid–free modified Eagle medium containing 10% fetal bovine serum and 1 mg/mL neomycin (Gibco), and resistant cells were cloned by the limiting dilution method. The rGPIa/IIa-producing CHO clone was cultured using EX-CELL 301 media (JRH Biosciences, Lenexa, KS) without serum. The culture supernatant was collected and concentrated by ultrafiltration. The rGPIa/IIa was purified by collagen Sepharose affinity chromatography by the method described previously.19 The eluates by 20 mM Tris-HCl (pH 7.5) containing 10 mM ethylenediaminetetraacetic acid and 150 mM NaCl were further purified by gel filtration chromatography (TSKgel3000SW; TOSO, Tokyo, Japan). The purity of the obtained rGPIa/IIa was more than 95% by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and Coomassie blue staining. Apparent dissociation constant values for the binding of rGPIa/IIa to collagen were determined using an enzyme-linked immunosorbent assay (ELISA).12 Human VWF was purified from human plasma cryoprecipitate. Purification steps were performed according to the previously described method.20 The VWF preparations used in our experiments had a VWF concentration of 2 mg/mL, specific activity of 200 U/mL, ristocetin cofactor activity, and 210 U/mL VWF antigen. Polycarbonate was obtained from Mitsubishi Engineer-

Preparation of reconstituted blood

Blood drawn from a healthy volunteer was mixed with a 1:10 volume of acid-citrate-dextrose composed of 2.2% (wt/vol) sodium citrate, 0.8% (wt/vol) citric acid, and 2.2% (wt/vol) glucose (ACD). The blood was centrifuged at 100g for 15 minutes at room temperature, and the platelet-rich plasma on top of the erythrocytes was replaced with an equal volume of 0.9% NaCl solution containing 10% (vol/vol) ACD (10% ACD-saline). Red cells were resuspended and centrifuged at 220g for 10 minutes at room temperature, and the supernatant was replaced with 10% ACD-saline. Each procedure was repeated twice. For perfusion studies, the red cells were reconstituted to 37.5% of the hematocrit (Hct) using 0.9% NaCl solution. The residual platelet count was 1.25 × 10^11/L (12.5 × 10^10/L). The Hct and platelet concentrations were determined using an automated hematology analyzer (SYSMEX, Kobe, Japan).

Preparation of liposomes

Liposomes were prepared by the detergent-dialysis method,21,22 originally developed for the reconstitution of membrane proteins, using the detergent OG. The protein was first conjugated to GPIa/IIa in the presence of detergent. The conjugated protein was then mixed with the lipid-detergent mixture, and the incorporation of protein is achieved upon the removal of the detergent by dialysis. Thus, NHS (0.1 M in H2O) and EDCI (0.25 M in H2O) were added to NGPE solubilized with 2% (wt/vol) OG in 50 mM MES buffer, pH 5.5, and the mixture was incubated for 10 minutes at room temperature. NGPE with an NHS-activated carboxylic derivative was purified using a Sephadex G-25 column with 50 mM HEPS/0.1% (wt/vol) OG, pH 8.0, and was added to a solution of recombinant protein. The resultant solution was incubated for 12 hours at 4°C with gentle stirring. For rhodamine-labeled liposome preparation,23 a thin film of the lipid mixture containing EPC, CHO, and N-Rh-PE in a molar ratio of 2:1:0.024 was solubilized with OG in 50 mM HEPS/10 mM NaCl buffer, pH 7.4. The resultant solution was mixed vigorously with the NGPE-conjugated protein. The liposomes were then purified using a Sephadex G-75 column, CsCl density gradient centrifugation, and dialysis against 0.9% NaCl. Control liposomes were made with EPC, CHO, and N-Rh-PE in a molar ratio of 2:1:0.024 in the absence of the NGPE-conjugated protein. The liposomes were extruded repeatedly through double-stacked 1.0- and 0.8-µm pore-size polycarbonate membranes (Whatman/Nuclepore, Clifton, NJ) in a high-pressure extrusion cell (Lipex Biomembrane, Vancouver, British Columbia, Canada) as described before24 to produce a final mean diameter range of 800 to 900 nm. Liposomes with different protein-to-lipid ratios were obtained by altering the initial protein-to-lipid ratio. EPC and CHO were quantified using a phospholipid-test Wako and F-kit CHO, respectively. The exofacial densities of rGPIa/IIa and rGPIbα were determined using an ELISA with Integrin β1 EIA Kit (Takara Shuzo, Otsu, Japan) and Glycocalcin EIA Kit (Takara Shuzo, respectively). The exofacial density of rGPIa/IIa was also determined using an ELISA with anti-GPIa monoclonal antibody, Lia1/2, and horseradish peroxidase–conjugated functional anti-GPIa monoclonal antibody, HRP-G9. The amount of rGPIa/IIa or rGPIbα associated with the liposome bilayer was determined using the same method as described above in the presence of 1% (vol/vol) Nonidet P-40. The rGPIa/IIa and rGPIbα solutions were used as standards for measuring receptor density. Absorbance at 492 nm was measured with an Easy Reader EAR 340 (SLT-Lab Instruments, Grodig, Austria). The exofacial densities of rGPIa/IIa determined with Integrin β1 EIA Kit and Lia1/2/HRP-G9 system were very consistent within standard deviation. The liposome size was measured with a dynamic light scattering technique using a particle analyzer N4 PLUS (Beckman, Fullerton, CA). The particle numbers of liposomes were calculated based on particle size, EPC concentration, bilayer thickness (15.0 nm), and EPC specific gravity (1.0305).
Preparation of the immobilized collagen surface

Glass slides, 2.5-cm diameter and 0.5-mm thick, were spin-coated with 6% (wt/vol) polycarbonate solution in tetrachloroethane. The glass slides were then incubated with 30 μg/mL porcine tendon acid soluble type I collagen (Nitta Gelatin, Osaka, Japan) in phosphate-buffered saline overnight at 4°C followed by blocking with 1% (wt/vol) BSA in phosphate-buffered saline. After removing excess BSA with 3 sequential phosphate-buffered saline rinses, the glass slides were assembled in the chamber to measure the interaction of the liposomes with the immobilized collagen.

Measurements of the interaction of the liposomes with immobilized collagen

The interaction of rhodamine-labeled liposomes with immobilized collagen was studied using a recirculating chamber, mounted on an epifluorescence microscope, (ECLIPS TE300, Nikon, Tokyo, Japan), using the excitation and emission wavelengths of 550 and 590 nm, respectively. This allowed direct visualization in real time of the lipidosome interaction with the collagen surface, which was recorded with a videocassette recorder. The flow chamber consisted of upper lid, packing, and glass slide. The upper lid had a depression of 0.030 cm perpendicular to the blood flow and served as part of the roof of the flow chamber that was formed when the upper lid and the glass slide were joined with 4 screws. The packing, hollowed out of a square 1.5 x 1.5 cm, was put between the upper lid and the glass slide, making a flow chamber with a width, length, and depth of 1.5 x 1.5 cm by 0.030 cm. The wall shear rate ($\gamma_w$) is given by the Muggli equation$^{25}$: $\gamma_w = 1.03 \times 6Q/ab^2$, where $Q$ is the flow rate (cm$^3$/sec), and $a$ and $b$ are the chamber width and height (cm).

Perfusion studies were performed in the presence of liposomes at a final particle number of $2.5 \times 10^5$ μL$. Hct 37.5\%, platelet count $1.25 \times 10^9$ μL$^{-1}$, 2 mM Mg$^{2+}$, 10 μg/mL soluble VWF, and 37°C. Some experiments were performed in the absence of soluble VWF. Single-frame images of the liposomes interacting with the surface were obtained using the image processor ARGUS-50 (Hamamatsu Photonics, Hamamatsu, Japan). The percentages of surface coverage of liposomes were obtained using the image processor ARGUS-20 (Hamamatsu Photonics). For the inhibition experiments, the liposomes were incubated with 10 μg/mL mouse anti-GPIb monoclonal antibody, GUR83-35, 10 μg/mL mouse anti-GPIa monoclonal antibody, Gi9, or 10 μg/mL control mouse IgG for 5 minutes at 37°C before perfusion.

Results

Adhesion of rGPIa/IIa-liposomes to the collagen surface under flow conditions

In marked contrast with the translocation of GPIbα-liposomes on the VWF surface,$^{1,4}$ rGPIa/IIa-liposomes instantaneously and irreversibly adhered to the collagen surface. Each single frame shown in Figure 1 was obtained after 3 minutes of perfusion of rGPIa/IIa-liposomes with an exofacial rGPIa/IIa density of $2.22 \times 10^9$ molecules per particle on the collagen surface at different shear rates, as indicated. When exposed to shear rates of 600 s$^{-1}$ for 3 minutes, the percentages of surface coverage of rGPIa/IIa-liposomes were 23.0% ± 2.2% and 23.8% ± 2.0%, in the presence and absence of soluble VWF, respectively. At a shear rate of 2400
of soluble VWF (Figure 4A). When exposed to shear rates of 600 s⁻¹, the percentages of surface coverage of the liposomes were estimated to be 33.1% ± 2.3% and 23.1% ± 0.8%, in the presence and absence of soluble VWF, respectively. At a shear rate of 2400 s⁻¹, the surface coverage increased to 43.2% ± 3.8% in the presence of soluble VWF. In the absence of soluble VWF, however, the surface coverage decreased to 3.8% ± 0.7%, as observed with rGPIa/IIa-liposomes. The reduction of the exofacial density of

s⁻¹, the percentages of surface coverage remarkably decreased to 3.5% ± 0.6% and 3.0% ± 0.6%, in the presence and absence of soluble VWF, respectively. No interaction was observed between rGPIa/IIa-liposomes and the BSA surface at any shear rates tested regardless of whether or not soluble VWF was present (data not shown). The lipid adhesion was abolished by preincubation of the liposomes with the functional anti-GPIa monoclonal antibody, Gi9, or in the presence of free rGPIa/IIa (Figure 2). No effect of control mouse IgG on the lipid adhesion was observed (Figure 2). These results indicate that rGPIa/IIa-liposomes retain a receptor function against immobilized collagen, and the targeting of rGPIa/IIa-liposomes is specific to the collagen surface under flow conditions. Also, the adhesion of rGPIa/IIa-liposomes is more efficient in lower flow environments and is independent of VWF.

Adhesion of rGPIa/IIa-Iba-liposomes to the collagen surface under flow conditions

The adhesion of rGPIa/IIa-Iba-liposomes to the collagen surface was also instantaneous and irreversible. The images shown in Figure 3 are composites created by the superimposition of 30 successive frames, taken at 66-millisecond intervals. In the case of rGPIa/IIa-Iba-liposomes, the fluorescent dots of the liposomes stayed on the collagen surface, representing irreversible adhesion of the liposomes to the surface (Figure 3A,B). Short tracks formed by closely spaced fluorescent dots of rGPIa/IIa-liposomes extending in the direction of flow can be seen, demonstrating transient interaction of rGPIb-liposomes with VWF adsorbed on the collagen surface (Figure 3C). No interaction of control liposomes with the collagen surface was observed (Figure 3D).

Each single-frame image shown in Figure 4 was obtained after 3 minutes of perfusion of rGPIa/IIa-Iba-liposomes with different exofacial densities of rGPIb and rGPIa/Ila. The adhesion of rGPIa/Ila-Iba-liposomes with exofacial densities of rGPIb/Ila and rGPIb was 2.17 × 10⁷ and 1.00 × 10⁴ molecules per particle, respectively, was more efficient at high shear rates in the presence of soluble VWF (Figure 4A). When exposed to shear rates of 600 s⁻¹, the percentages of surface coverage of the liposomes were estimated to be 33.1% ± 2.3% and 23.1% ± 0.8%, in the presence and absence of soluble VWF, respectively. At a shear rate of 2400 s⁻¹, the surface coverage increased to 43.2% ± 3.8% in the presence of soluble VWF. In the absence of soluble VWF, however, the surface coverage decreased to 3.8% ± 0.7%, as observed with rGPIa/Ila-liposomes. The reduction of the exofacial density of

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rGPlbo by almost 50% (5.27 × 10^3 molecules per particle), while the exofacial density of rGPla/IIa was kept constant at approximately 2.00 × 10^3 molecules per particle, resulted in a decreased surface coverage at high shear rates (Figure 4B). The surface coverage decreased from 33.8% ± 0.9% to 19.2% ± 0.8% with an increasing shear rate from 600 to 2400 s⁻¹ in the presence of soluble VWF. In the absence of soluble VWF, the percentages of surface coverage decreased from 24.4% to 4.6% ± 0.8% with an increasing shear rate from 600 to 2400 s⁻¹ as observed with rGPla/IIa-Ib-liposomes with exofacial densities of rGPla/IIa and rGPlbo of 2.17 × 10^3 and 1.00 × 10^3 molecules per particle, respectively. These results suggest that the interaction of rGPlbo on the liposome surface with the collagen surface is negligible in the absence of soluble VWF. A 50% reduction in the exofacial rGPla/IIa density resulted in decreased adhesion by the liposomes at all shear rates tested (Figure 4C). The percentages of the surface coverage of the liposomes with exofacial densities of rGPla/IIa and rGPlbo of 0.96 × 10^3 and 1.08 × 10^3 molecules per particle, respectively, decreased from 16.9% ± 2.2% to 3.1% ± 0.9% with increasing shear rates from 600 to 2400 s⁻¹ in the presence of soluble VWF. In the absence of soluble VWF, the surface coverage decreased from 10.8% ± 3.9% to 0.8% ± 0.4%.

Surface coverage of the liposomes with different exofacial rGPlbo densities (1.00 × 10^3, 5.27 × 10^3, and 0 molecules per particle), and an equivalent exofacial rGPla/IIa density at approximately 2.00 × 10^3 molecules per particle, are shown in Figure 5A. It is clear that high densities of rGPlbo on the liposome surface and the presence of soluble VWF are required for efficient adhesion to the collagen surface at high shear rates. The surface coverage of the liposomes with different exofacial rGPla/IIa densities (2.17 × 10^3, 0.96 × 10^3, and 0 molecules per particle), while the exofacial rGPlbo density was kept constant at approximately 1.00 × 10^3 molecules per particle, is shown in Figure 5B. The liposomes carrying rGPlbo alone at an exofacial density of 1.16 × 10^3 molecules per particle never formed a stationary adhesion on the collagen surface, but stopped transiently in the millisecond range on the surface, demonstrating the tethering of the liposomes to VWF adsorbed on the collagen surface (Figure 5B, front row). The duration of contact with the surface was calculated to be less than 33 milliseconds. These results suggest that the rGPla/IIa–collagen interaction is important not only in lower flow environments, but also at high shear rates, and that rGPla/IIa and rGPlbo cooperatively contribute to the liposome adhesion, especially at high shear rates.

**Inhibitory effect of anti-rGPlbo or anti-rGPla antibody on the liposome adhesion to the collagen surface under flow conditions**

The inhibitory effects of antibodies are shown as the relative surface coverage, that is, the surface coverage in the presence of antibody relative to that in the absence of antibody (Figure 6). When the rGPlbo–VWF axis was blocked by the anti-rGPlbo antibody, GUR 83-35, the liposomes still adhered irreversibly to the collagen surface in a shear rate–dependent fashion. The relative surface coverage decreased from 66.2 ± 3.9% to 6.5% ± 2.6% with the shear rate increasing from 600 to 2400 s⁻¹ for the liposomes with exofacial densities of rGPla/IIa and rGPlbo of

![Figure 5. Percentages of surface coverage of rGPla/IIa-Ib-liposomes on the collagen surface.](image)

**Figure 5. Percentages of surface coverage of rGPla/IIa-Ib-liposomes on the collagen surface.** Dependence on shear rate, densities of rGPla/IIa and rGPlbo, and VWF. Values are the mean ± SD; n = 6. (A) Percentages of surface coverage of rGPla/IIa-Ib-liposomes with different exofacial densities of rGPlbo, as indicated. The exofacial density of rGPla/IIa was kept constant at 2.19 × 10^3 ± 0.02 × 10^3 molecules per particle. (B) Percentages of surface coverage of rGPla/IIa-Ib-liposomes with different exofacial densities of rGPlbo, as indicated. The exofacial density of rGPlbo was kept constant at 1.08 × 10^3 ± 0.08 × 10^3 molecules per particle. The percentages of surface coverage of rGPlbo-liposomes on the collagen surface in the presence and absence of soluble VWF (front row) were 0.07% ± 0.02% at all shear rates tested.

![Figure 6. Inhibitory effect of GUR83-35 or G9 on the adhesion of rGPla/IIa-Ib-liposomes to the collagen surface under flow conditions.](image)

**Figure 6. Inhibitory effect of GUR83-35 or G9 on the adhesion of rGPla/IIa-Ib-liposomes to the collagen surface under flow conditions.** Relative surface coverage of rGPla/IIa-Ib-liposomes with different exofacial densities of rGPla/IIa and rGPlbo, in the presence of specific antibody and 10 μg/mL soluble VWF, are shown. Values are the mean ± SD; n = 6. White bar indicates control mouse; black bar, GUR83-35 (+); grey bar, G9 (+). (A) The rGPla/IIa-Ib-liposomes with exofacial densities of rGPla/IIa and rGPlbo at 2.17 × 10^3 and 1.00 × 10^3 molecules per particle, respectively. (B) The rGPla/IIa-Ib-liposomes with exofacial densities of rGPla/IIa and rGPlbo at 2.19 × 10^3 and 5.27 × 10^3 molecules per particle, respectively. (C) The rGPla/IIa-liposomes with an exofacial density of rGPla/IIa at 2.22 × 10^3 molecules per particle.
2.17 × 10^3 and 1.00 × 10^4 molecules per particle, respectively (Figure 6A). The same trend was observed for liposomes with exofacial densities of rGPIa/IIa and rGPIb on 2.19 × 10^3 and 5.27 × 10^3 molecules per particle, respectively (Figure 6B), although the inhibitory effects were smaller than those in liposomes with a higher exofacial density of rGPIb at a shear rate of 2400 s⁻¹ (compare Figures 6A and 6B). No effect of GUR 83-35 was observed for the liposomes carrying rGPIa/IIa alone (Figure 6C). These results suggest that the inhibitory effect of GUR 83-35 is greater at higher shear rates and the extent of dependence of liposome adhesion on the rGPIb–VWF interaction is greater at higher shear rates. When the rGPIa/IIa-collagen axis was blocked by the anti-rGPIa antibody, Gi9, the liposome displacement on the surface was observed, as with the rGPIb-liposomes on the collagen surface. The inhibitory effect of Gi9 was always greater than that of GUR 83-35, especially at low shear rates. No effect of control mouse IgG on the liposome adhesion was observed. These observations indicate that both the rGPIa/IIa–collagen interaction and the tethering of the liposomes by the rGPIb–VWF interaction are required for liposome adhesion, and they synergistically contribute to stable adhesion of rGPIa/IIa-liposomes, especially at high shear rates.

**Discussion**

The recognition of exposed subendothelial collagen by blood platelets is a key early step in the formation of a hemostatic plug after vascular injury. Many different platelet surface and platelet surface–associated proteins have been proposed as mediators of platelet–collagen adhesion. Santoro has defined a Mg²⁺-dependent mechanism of platelet adhesion to collagen²⁶ apparently identical to that observed by Shadle and Barondes²⁶ and have isolated a platelet surface Mg²⁺-dependent heterodimeric collagen-binding complex composed of platelet membrane GPIa and GPIa/IIa.²⁷ When incorporated into liposomes, the purified complex mediated the Mg²⁺-dependent adhesion of the liposomes to collagen substrates at static conditions.²⁸,²⁹ The rGPIa/IIa used in this study has an activated form and the specific binding to collagen characterized by a dissociation constant of the same order of magnitude as that for the binding of collagen to GPIa/IIa on activated platelets.²² Also, rGPIb used in this study has an affinity of interaction with VWF characterized by a dissociation constant of the same order of magnitude as that reported previously for the binding of VWF to GPIb–IX on platelets.²³,²⁴,³¹

In the present study, liposomes carrying rGPIa/IIa and rGPIb were chosen as a model system used to examine the process of initiating the adhesion of platelets under flow conditions. Such proteoliposomes previously prepared by direct hydration followed by freeze–thawing³² are much too small (diameters of 200 nm or less) to be useful for fluorescence microscopy studies. In addition, liposomes with diameters less than 80 nm have a very high binding energy and thus will not undergo adhesion, with adhesion above this critical size, however, increasing with vesicle size.³³ We therefore prepared the liposomes carrying both rGPIa/IIa and rGPIb by detergent dialysis followed by extrusion through polycarbonate membranes to produce a final mean diameter range of 800 to 900 nm, suitable for adhesion studies with fluorescence microscope under flow conditions.

Our results suggest that 2 distinct substrates, collagen and VWF, are required in order to provide the biomechanical properties necessary to mediate stable liposome adhesion, especially at high shear rates. The rGPIa/IIa supports immediate arrest of flowing liposomes onto the collagen surface but works efficiently only at the lower shear rates, presumably because of a relatively slow rate of bond formation with immobilized collagen and a low resistance of the bond to tensile stress. In contrast, the interaction of rGPIb with immobilized VWF is inherently not sufficient to arrest the liposomes but results in a very marked decrease in velocity of flowing liposomes, relative to the hydrodynamic flow, when surface contact is established.¹⁴ Moreover, possibly because of a fast bond formation and the high resistance of the bond to tensile stress, this function is efficiently displayed even at higher shear rates. Thus, rGPIa/IIa is essential for the stability of liposome adhesion to the collagen surface. The interaction of rGPIb with VWF immobilized on the collagen surface, however, is required first to reduce the velocity of the liposomes contacting the surface under high flow conditions, thereby prolonging the time available for the bond formation of rGPIa/IIa with immobilized collagen. When the shear rate is low, the function of VWF is initially limited because of the reduction of the interactions between rGPIb and immobilized VWF,¹⁴ the soluble VWF and the collagen surface, or both, and the function of rGPIa/IIa as a collagen receptor is efficiently displayed.

Our findings now define a unique function for rGPIa/IIa, expressed by its ability to act in concert with the rGPIb–VWF interaction to promote stable adhesion of rGPIa/IIa-Ib-liposomes to the collagen surface. The 2 receptors, rGPIa/IIa and rGPIb, therefore, have complementary roles, and the corresponding adhesive substrates, collagen and VWF, are also complementary in the adhesion of rGPIa/IIa-Ib-liposomes.

These results contribute to the long-term purpose of our studies, which is to prepare liposome systems that improve primary hemostasis under thrombocytopenic conditions and that are promising agents for the prophylaxis and treatment of bleeding in patients with severe thrombocytopenia. The simplest type of artificial platelets might be particles carrying platelet membrane proteins and/or ligands of the proteins involved in platelet adhesion and aggregation. Based on this idea, some materials have been developed as platelet substitutes, such as erythrocytes with fibrinogen, or RGD peptides, covalently linked to their surfaces,³⁴,³⁵ liposomes bearing more than 15 kinds of platelet membrane proteins (eg, GPIb, GPIIb/IIIa, GPVI) isolated from the platelet membranes with deoxycholate,³⁶ and fibrinogen-coated albumin microparticles.³⁷,³⁸ Some of these are reactive with adhesive ligands or with normal platelets in vitro, or are effective in enhancing hemostatic function in thrombocytopenic or thrombocytopenic and fibrinogen-coated animals in vivo. Recently, it has been determined that rGPIa/IIa-liposomes have hemostatic activity in vivo.¹² However, no platelet substitute has yet been reported to be effective for hemostasis in large clinical studies so far.

In conclusion, we have developed an effective tool for studying adhesive interactions of platelets under flow conditions and proved that 2 distinct receptor–ligand pairs with unique properties, GPIa/IIa–collagen and GPIb–VWF, complement each other and synergistically provide the needed functional integration required for platelet adhesion under unfavorable shear forces. Furthermore, our results have demonstrated that the liposomes carrying rGPIa/IIa and/or rGPIb are the potential candidates for platelet substitutes. Development of effective platelet substitutes using liposome system is now underway.

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References


Reconstitution of adhesive properties of human platelets in liposomes carrying both recombinant glycoproteins Ia/IIa and Ib α under flow conditions: specific synergy of receptor–ligand interactions

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