Human Erythrocyte Protein 4.1 Is a Phosphatidylserine Binding Protein

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Abstract

The aminophospholipids phosphatidylethanolamine (PE) and phosphatidylserine (PS) are the major phospholipids contained in the cytoplasmic leaflet of the human erythrocyte (RBC) plasma membrane and are largely confined to that leaflet over the entire RBC lifespan. In particular, PS, which comprises ~ 13% of total RBC membrane phospholipids, is normally restricted entirely to the cytoplasmic leaflet. However, molecular mechanisms that regulate this asymmetric distribution of phospholipids are largely unknown. We examined elliptocytic RBCs that completely lacked protein 4.1 (HE[4.1°]), but contained normal amounts of all other peripheral membrane proteins, and found $\sim 10\%$ of total membrane PS was accessible in the exoplasmic leaflet of these membranes. Inside out vesicles (IOVs) derived from HE[4.1°] RBCs bound fewer PS liposomes than did IOVs derived from normal RBCs. Normal IOVs that were depleted of proteins 2.1 (ankyrin), 4.1, and 4.2 bound fewer PS liposomes similar to HE[4.1°] IOVs, and repletion with protein 4.1 restored PS liposome binding to control levels. Addition of purified protein 4.1 to PS liposomes resulted in saturable binding with the extent of binding being proportional to the liposome PS content. Our data suggests that human RBC protein 4.1 is a PS binding protein and may be involved in the molecular mechanisms that stabilize PS in the cytoplasmic leaflet of the human RBC plasma membrane.

Introduction

It is now well established that the phospholipids that comprise the human erythrocyte (RBC)¹ plasma membrane are arranged asymmetrically between the bilayer leaflets (1). This asymmetric organization results in the exoplasmic leaflet being enriched in the choline-containing phospholipids, phosphatidylcholine (PC) and sphingomyelin (SM), and the cytoplasmic leaflet being enriched in the aminophospholipids, phosphatidylethanolamine (PE), and phosphatidylserine (PS). Notably,

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1. Abbreviations used in this paper: IOV, inside out vesicle; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PLA₂, phospholipase A₂; PS, phosphatidyldylserine; SM, sphingomyelin; SMC, sphingomyelinase C; RBC, erythrocyte.

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PS is found only in the cytoplasmic leaflet. Recently, it has been shown that this arrangement is not static and that dynamic transbilayer movement (i.e., flip-flop) of some phospholipids does occur (2-4). That the asymmetric organization of phospholipids is maintained throughout the 120-d average lifespan of RBCs (5) implies that some mechanism must exist to stabilize specific phospholipids within bilayer leaflets. It has been suggested that spectrin is involved in this process since spectrin interacts with aminophospholipids in lipid monolayers (6), phospholipid vesicles (7), and in intact RBCs (8). Also, partial loss of membrane phospholipid asymmetry occurs upon depletion of RBC membranes of spectrin during preparation of inside out vesicles (IOVs) in vitro (9), and in spectrin-free microvesicles spontaneously released from aged RBC (10, 11).

The present work examines the contribution of protein 4.1 towards the maintenance of RBC membrane phospholipid asymmetry. Protein 4.1 is known to bind to the distal end of the spectrin tetramer (12), associate with the cytoplasmic domain of integral membrane glycophorins (13, 14), and interact with protease digested IOVs (15). We examined RBCs from a patient with homozygous hereditary elliptocytosis whose membranes completely lacked protein 4.1 (HE[4.1°])² but contained normal amounts of all other membrane cytoskeletal proteins. In addition to an abnormal cell morphology, the exoplasmic bilayer leaflet of these HE[4.1°] RBCs contained PS. We also evaluated the interaction of protein 4.1 with phospholipid model membranes (liposomes) and found that protein 4.1 interacts specifically and saturably with liposomes containing PS. Taken together, these studies suggest that interactions between protein 4.1 and PS occur in RBC membranes and may contribute to the maintenance of membrane PS asymmetry.

Methods

Materials. PS from bovine brain and PC from egg yolk were purchased from Avanti Polar Lipids, Inc., Birmingham, AL. Lipids migrated as single spots on TLC and were stored under nitrogen in sealed vials at -70°C until used. [³H]triolein and ¹25I-Bolton Hunter reagent were obtained from New England Nuclear, Boston, MA. Dialysis tubing (Spectrapor-2) was from Spectrum Medical, Los Angeles, CA. DTT was from Calbiochem-Behring Corp., La Jolla, CA. Sphingomyelinase C (SMC) from Staphylococcus aureus was the generous gift of Drs. B. Roelofsen and L. L. M. van Deenen, Utrecht, Netherlands. Phospholipase A2 (PLA2) from bee venom and Ficoll 400 were from Sigma Chemical Co., St. Louis, MO. All gel electrophoresis reagents were electrophoretic grade from Bio-Rad Laboratories, Richmond, CA. All other chemicals were reagent grade from standard sources.

Blood collection, RBC ghost, and membrane vesicle preparation. After obtaining informed consent, fresh blood samples from a patient with hereditary elliptocytosis whose RBCs were completely deficient in protein 4.1 (HE[4.1°]) and from healthy normal controls were col-

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^{2.} Nomenclature suggested by J. Palek and S. E. Lux to describe RBC membrane cytoskeletal defects (1983. Semin. Hematol. 20:189-224).

lected in acid citrate dextrose. RBC ghosts were prepared by the method of Dodge et al. (16), except that 0.4 mM diisopropylfluorophosphate was added at the lysis step. Spectrin-actin-depleted IOVs from normal and HE[4.1°] ghosts were prepared by the method of Steck and Kant (17). IOV membrane sidedness was determined by glyceraldehyde-3-phosphate dehydrogenase and acetylcholinesterase assays (18). Some of the normal IOVs were further depleted of ankyrin, protein 4.1, and protein 4.2 by the potassium iodide method of Bennett and Stenbuck (19). Protein 4.1 was reassociated with protein 4.1-depleted IOVs by the method of Shiffer and Goodman (20).

Treatment of RBCs with PLA₂ and SMC. Normal and HE[4.1°] RBCs were treated with PLA₂ for 1 h at 37°C followed by SMC for 1 h at 37°C, according to the procedure of Lubin et al. (21). Phospholipid degradation was terminated by washing RBCs three times with PBS containing 5 mM EDTA. The extent of cell hemolysis was determined by comparing the hemoglobin content in each sample supernate with hemolyzed controls. Lipid extraction was by the method of Rose and Oklander (22). Individual phospholipid classes were separated by the two-dimensional TLC technique of Roelofsen and Zwaal (23). Separated phospholipids were stained, extracted from the TLC plates, and quantitated by phosphorus determination, as described by Bartlett (24).

Preparation of liposomes. Unilamellar liposomes composed of varying molar ratios of PS and PC were prepared by the reverse phase evaporation technique of Szoka and Papahadjopoulos (25), with some modifications (26). In some experiments, the nontransferable phospholipid [³H]triolein was added in trace amounts (0.007% of total lipid) to allow quantitation of liposome binding. After the vesicles were formed, they were uniformly sized to 100 nm by extrusion through polycarbonate membranes (Bio-Rad Laboratories) (27, 28). Liposomes were stored under argon at 4°C and were used within 1 wk of their preparation. The concentration of phospholipid in the liposome preparations was determined by phosphorus assay (24).

Purification and radiolabeling of protein 4.1. Protein 4.1 was purified from RBCs by the method of Tyler et al. (29), with modifications by Cohen and Foley (30). For some experiments, protein 4.1 was denatured by heating at 65°C for 20 min. In our hands, heat-denatured protein 4.1 did not sediment at the concentrations used in these studies. Protein determinations were performed by the method of Bradford (31). SDS-PAGE was performed as described by Laemmli (32), and the protein bands were quantitated by densitometry of Coomassie Brilliant Blue stained gels. For some experiments protein 4.1 was radiolabeled with ¹²⁵I-Bolton Hunter reagent (33). Protein 4.1 was used when prepared, or in some cases, stored in siliclad-treated polycarbonate tubes at -70°C in PBS containing 20 mM potassium chloride, 1 mM disodium EDTA, 0.2 mM DTT, and 0.4 mM diisopropylfluorophosphate.

IOV binding to liposomes. IOVs (12 μ g phosphorus) were incubated at 37°C for 1 h with 40 nmol radiolabeled liposomes (PS/PC, 3:1 molar ratio) in a total volume of 0.2 ml buffer A (20 mM sodium Hepes, pH 7.4, containing 140 mM NaCl). After the incubation, the liposome-IOV complex was sedimented at 39,000 g for 30 min at 4°C and washed twice in buffer A. The amount of membrane-bound liposomes was determined by the radioactivity contained in the washed IOV pellets.

Protein 4.1 binding to liposomes. ¹²⁵I-Protein 4.1 was incubated with liposomes (50 μ mol lipid) in siliclad-coated polycarbonate tubes in a total volume of 0.25 ml buffer A at 37°C for 30 min. Liposome-bound protein was separated from unbound protein by flotation through a Ficoll 400 discontinuous density gradient as follows: a stock solution of 30% Ficoll 400 was added to the ¹²⁵I-protein 4.1-liposome incubation mixture to a final concentration of 3 ml 10% Ficoll. Siliclad-treated 13 \times 51 mm polyallomer ultracentrifuge tubes were layered (from bottom to top of tube) with 1 ml of 30% Ficoll 400, 3 ml of the 10% Ficoll-liposome-protein 4.1 incubation mixture, and 0.5 ml of buffer A. The gradient was centrifuged for 1 h at 25°C (model SW 50.1 swinging bucket rotor at 35,000 rpm in a model L8-80 ultracentrifuge; Beckman Instruments, Inc., Fullerton, CA). Protein 4.1 binding to liposomes was determined from the ¹²⁵I-protein 4.1 and

phosphorus (liposome) content of the top 0.8 ml (liposome containing portion) of the gradient. Samples were run in duplicate and the duplicates varied by no more than 5%.

Results

Bilayer distribution of membrane phospholipids in normal and HE[4.1°] RBC. RBC membranes from a patient with homozygous hereditary elliptocytis (HE[4.1°]) differed from normal RBCs in that they were completely deficient in protein 4.1, as shown by both SDS-PAGE and immunologic characterization (Fig. 1). When HE[4.1°] RBCs were subjected to analysis of membrane phospholipid sidedness, we found that $\sim 10\%$ of the total membrane content of PS was available for degradation using the combination of PLA2 plus SMC, whereas there was no degradation of PS in normal RBCs identically treated (Table I). Treatment of HE[4.1°] RBCs with PLA2 plus SMC at the levels used in these experiments did not produce significant cell hemolysis (6%). In the HE[4.1°] RBCs there was also a 9% decrease in the exoplasmic leaflet content of PC, presumably to offset the movement of PS into this leaflet and maintain an equimolar distribution of phospholipids in each bilayer leaflet.

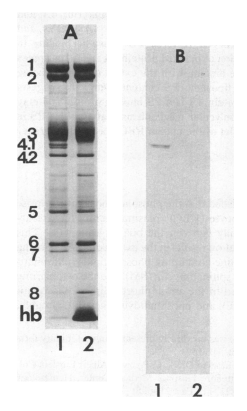


Figure 1. (A) Protein composition of normal and HE[4.1°] RBC ghost membranes. Proteins from normal (lane 1) and HE[4.1°] (lane 2) ghost membranes (20 μ g) were separated by SDS-PAGE (6-15% linear gradient). The nomenclature of the polypeptide bands is according to Steck (34). (B) Immunoblot of normal (lane 1) and HE[4.1°] (lane 2) ghost membranes. Proteins (20 μ g) were transferred to nitrocellulose paper that was incubated with monospecific rabbit anti-human RBC protein 4.1 IgG and the antigen-antibody complex visualized with goat anti-rabbit IgG horseradish peroxidase conjugate followed by color development with 4-chloro-1-naphthol (35).

Table I. Membrane Phospholipid Organization in Normal, Protein 4.1-deficient (HE[4.1°]), and Glycophorin C-deficient (Ge⁻) Human RBCs

RBC	Phospholipid degraded				
	PC	PE	PS	SM	Hemolysis
	%	%	%	%	%
Normal	78±6	23±4	0	63±6	3±2
HE[4.1°]	70±8*	21±3	10±3*	63±7	6±2
Ge ⁻	74±5	22±2	0	65±5	3±2

RBCs were treated with bee venom PLA₂ for 1 h at 37°C followed by *Staphylococcus aureus* SMC for 1 h at 37°C. Percentage phospholipid degradation was determined as described in Methods. Results are the mean±1 SD of 10 determinations.

Since these HE[4.1°] RBC membranes are also > 80% deficient of glycophorin C (14), it was possible that the loss of PS asymmetry in the HE[4.1°] membranes was related to the deficiency of this protein and not to the deficiency of protein 4.1. To examine this, we determined membrane phospholipid sidedness in RBCs lacking the Gerbich blood group antigen (Ge⁻). Ge⁻ RBCs differ from normal RBCs in that they completely lack glycophorin C, but contain normal amounts of all other cytoskeletal proteins (Fig. 2 and reference 36). We found, as previously reported by Kuypers et al. (39), that the bilayer distribution of membrane phospholipids was normal in Ge⁻ cells (Table I), indicating that the loss of PS asymmetry in the HE[4.1°] RBCs was not related to the deficiency of glycophorin C.

Binding of RBC membrane vesicles to PS liposomes. Incubation of spectrin-actin-depleted IOVs derived from normal RBCs with PS-containing liposomes (PS/PC, 3:1 molar ratio) resulted in liposome binding (Table II). Similar incubation of liposomes with IOVs derived from HE[4.1°] RBCs or with IOVs that were stripped of proteins 2.1 (ankyrin), 4.1, and 4.2 resulted in a 20% decrease in liposome binding. Reassociation of protein 4.1 with the stripped IOVs restored PS liposome binding to control values.

Effect of phospholipid species on protein 4.1-liposome binding. To further characterize the interaction of protein 4.1 with PS, we incubated liposomes containing varying molar ratios of PS and PC with radiolabeled protein 4.1. Liposomebound protein was separated from unbound protein by centrifugation of the liposome-protein 4.1 mixture through a Ficoll 400 discontinuous density gradient. In this method, liposomebound protein floats to the top of the gradient (Ficoll-buffer interface), whereas unbound protein remains in the gradient (Fig. 3). We found that the extent of protein 4.1 binding to liposomes was directly proportional to the liposome PS content (Fig. 4). Binding of protein 4.1 to 100% PC liposomes was ~ 15% of binding to 100% PS liposomes and may represent nonspecific protein binding. A binding isotherm performed with protein 4.1 and 100% PS liposomes demonstrated saturable binding; saturating at $\sim 100 \,\mu g$ protein 4.1/ μ mol PS (Fig. 5). Similar studies employing 100% PC liposomes showed that protein 4.1 binding was never > 5% of the binding to 100% PS

liposomes (data not shown). To ascertain whether radiolabeling protein 4.1 had denatured or altered the protein, thereby affecting its association with liposomes, we evaluated the specific radioactivity of protein 4.1 before and after incubation with PS liposomes. We found that the specific radioactivity of unbound and liposome-bound ¹²⁵I-protein 4.1 was essentially unchanged (unbound, 4,892 cpm/ μ g protein and bound, 4,861 cpm/ μ g protein). Furthermore, the extent of protein 4.1 liposome binding was identical using high specific radioactivity protein (150,000 cpm/ μ g) and low specific radioactivity protein (5,000 cpm/ μ g). These results demonstrate that iodinating the protein with the Bolton Hunter reagent does not affect its ability to bind PS liposomes, nor does it result in a selective population of protein 4.1 molecules that have unique PS binding characteristics.

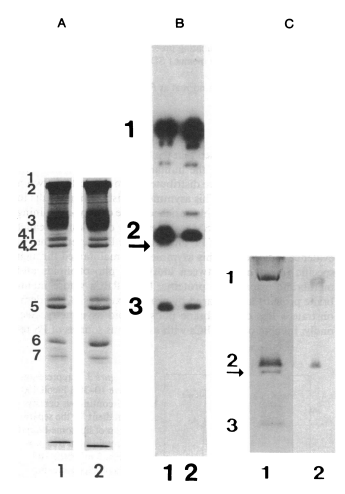


Figure 2. (A) Protein composition of normal and Ge⁻ RBC ghost membranes. Proteins from normal (lane I) and Ge⁻ (lane 2) ghost membranes (20 μ g) were separated by SDS-PAGE (9% acrylamide). The nomenclature of the polypeptide bands is according to Steck (34). (B) Fluorograph of gel shown in A using the Western immunoblotting technique (35). The blots were incubated with human antiglycohorin antibodies followed by 125 I-protein A and exposed to Kodak XAR-5 film for 2 d, as described by Lu et al. (37). The nomenclature of the glycophorins is according to Steck (34). The arrow indicates the position of glycophorin C. (C) Periodic acid-Schiff's base stain (38) of gel shown in A. The arrow indicates the position of glycophorin C.

^{*} Statistically different from normal at P < 0.01.

Table II. Liposome Binding to Normal, Protein 4.1-deficient (HE[4.1°]), Protein 4.1-depleted, and Protein 4.1-repleted IOVs

IOVs	IOV protein 4.1 content	IOV liposome binding	
	% of normal	nmol liposome/µg IOV	
Normal	100	4.7±0.1	
HE[4.1°]	0	3.9±0.4*	
Protein 4.1 depleted	18	3.9±0.2*	
Protein 4.1 repleted	110	4.6±0.6	

IOVs were incubated with radiolabeled liposomes (PS/PC, 3:1 molar ratio) for 1 h at 37°C and collected by centrifugation at 39,000 g for 30 min at 4°C. IOVs were then washed twice and the amount of bound liposomes was determined by the residual liposome radioactivity contained in the washed membrane pellet. IOVs were quantitated as micrograms membrane phosphorus. IOV protein 4.1 content was determined by quantitative densitometry of Coomassie Bluestained SDS-PAGE gels. Protein 4.1-depleted IOVs were prepared from normal IOVs by incubation with 1 M potassium iodide, which extracts ankyrin, protein 4.1, and protein 4.2. Protein 4.1-repleted IOVs were prepared by incubating protein 4.1-depleted IOVs with purified 4.1. Results are the mean±1 SD of 4 individual determinations.

Discussion

It is now firmly established that the four major classes of phospholipids that comprise the human RBC plasma membrane (PC, PE, PS, and SM) are distributed asymmetrically between the bilayer leaflets (1). This asymmetric distribution results in the exoplasmic leaflet being enriched in the choline-containing phospholipids PC and SM, whereas the cytoplasmic leaflet contains the majority of PE and all of the PS. Several lines of evidence indicate that this asymmetry is maintained through specific interactions between individual phospholipids and membrane cytoskeletal proteins. First, RBCs whose membrane proteins are damaged by protein oxidants (8, 40) demonstrate a loss of membrane phospholipid asymmetry. Secondly, incubations of RBCs with exogenous sources of PS re-

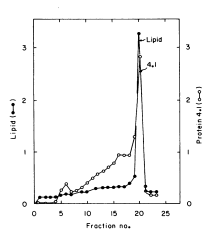


Figure 3. A representative 10-30% Ficoll 400 discontinuous density gradient for the separation of liposome-bound protein from free protein. 3 ml of the 10% Ficoll-liposome-protein incubation mixture and 0.5 ml of 5 mM sodium Hepes, pH 7.4, containing 140 mM NaCl were lavered over 1 ml 30% Ficoll and centrifuged for 1 h at 25°C in a rotor (SW 50.1, Beckman Instru-

ments, Inc.) at 35,000 rpm. Fractions were collected from the bottom of the centrifuge tube and (•) liposome content determined by phosphorus determination and (o) protein 4.1 content determined by ¹²⁵I-radioactivity.

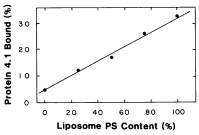


Figure 4. Protein 4.1 binding to liposomes. Radiolabeled protein 4.1 (5 μ g) was incubated with 50 nmol liposomes in a total volume of 0.25 ml for 30 min at 37°C. Liposome bound ¹²⁵I-protein 4.1 was separated from un-

bound protein by centrifugation through a discontinuous 10–30% Ficoll 400 gradient, as described in Fig. 3. Data points represent the average of duplicate determinations and were corrected for 2% nonspecific protein binding to liposomes evaluated using a heat-denatured (65°C for 20 min) protein 4.1 control. The line drawn was a best fit determined by linear regression analysis.

sults in the uptake of PS into the exoplasmic bilayer leaflet followed by translocation into the cytoplasmic leaflet; the rate and extent of the translocation being dependent on the PS acyl-chain composition and the metabolic state of the cell (4, 41). Once translocated into the cytoplasmic leaflet this exogenously added PS is maintained in that leaflet, as evidenced by the inability to quench spin-labeled PS with exogenously added reducing agents (4), and by the irreversible nature of stomatocytic shape changes induced by its incorporation into the cytoplasmic leaflet (41). Thirdly, RBC membrane phospholipid asymmetry is maintained throughout the 120-d average lifespan of the cells (5), despite the fact that dynamic bilayer translocation of phospholipids does occur in intact cells (3, 4).

The work presented in this paper was undertaken to examine the interactions between protein 4.1 and PS. A previous report by Sato and Ohnishi (15) demonstrated that RBC protein 4.1 interacts with PS liposomes and protease-digested IOV membranes. We hypothesized that such interactions could stabilize PS in the cytoplasmic leaflet of RBC plasma membranes. To test this hypothesis, we used RBCs that completely lacked protein 4.1 but contained normal amounts of all other membrane peripheral proteins (HE[4.1°]). The morphology of these cells was abnormal in that most cells were elliptocytic. Despite the presence of a normal amount of spectrin in these cells, we found that $\sim 10\%$ of the total membrane content of PS was accessible for hydrolysis by PLA₂ and SMC, whereas in normal RBCs no PS was hydrolysed by this treatment. The bilayer distribution of PE and SM was not affected in these

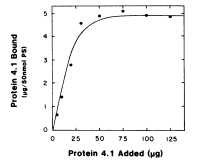


Figure 5. Binding isotherm of radiolabeled normal protein 4.1 to PS liposomes. Increasing amounts of 125 I-protein 4.1 (5–125 μ g) were incubated with 50 nmol liposomes in a total volume of 0.25 ml for 30 min at 37°C. Liposome-bound 125 I-protein 4.1 was separated from free protein by centrifugation through a discontinous Ficoll 400

density gradient and liposome and protein contents quantitated, as described in Fig. 3. Data points represent the average of duplicate determinations.

^{*} Statistically different from normal at P < 0.01.

cells although there was some redistribution of PC from the exoplasmic to the cytoplasmic leaflet, presumably to maintain an equimolar amount of total phospholipid in each bilayer leaflet. Note that differences in cell hemolysis caused by treatment with the phospholipases can affect the amount of phospholipids hydrolysed. However, our experience, and those of others using phospholipases to examine membrane phospholipid asymmetry, shows that below 10% hemolysis there is a 0.1% increase in PS degradation for every 1% increase in hemolysis. This increase is attributable to penetration of the enzymes into the cell interior. Therefore, at 6% hemolysis (measured for the HE[4.1°] RBCs) we would expect only 0.6% artefactual PS degradation, which could not account for the 10% degradation of PS measured in the HE[4.1°] cells. Also, note that in the HE[4.1°] RBCs only the transmembrane distribution of PS was found to be altered; in other pathologic RBCs that we have examined, in which the primary membrane defect is thought to involve spectrin, such as in sickle cells and spectrin-deficient cells, bilayer redistributions of both PE and PS were always observed together (5).

Membranes from HE[4.1°] RBCs are also > 80% deficient in glycophorin C (synonyms PAS-2, glycoconnectin), a minor sialoglycoprotein in the RBC membrane (14). To determine whether the absence of glycophorin C contributed to the availability of PS in the exoplasmic leaflet of the HE[4.1°] RBCs. we examined membrane phospholipid organization in RBCs lacking the Gerbich blood group (Ge⁻) antigen. Ge⁻ RBCs differ from normal RBCs in that they completely lack glycophorin C, but contain normal amounts of all other cytoskeletal proteins, including protein 4.1 (reference 36 and confirmed in our laboratory). We found that Ge⁻ RBCs did not have any PS in the membrane exoplasmic leaflet, confirming previous observations of Kuypers et al. (39). These observations strongly suggest that it was the absence of protein 4.1, and not a deficiency of glycophorin C, which was principally responsible for the abnormal increased availability of PS in the exoplasmic leaflet of the HE[4.1°] RBCs.

We next evaluated PS liposome binding capacity to IOVs derived from normal, protein 4.1-depleted, and HE[4.1°] RBC membranes. Normal IOVs are spectrin-actin-depleted but contain protein 4.1, the presence of which enhanced PS liposome binding as compared with either the protein 4.1-depleted or the HE[4.1°] IOVs. The differences were, however, relatively small (~ 20%) presumably because much of the protein 4.1 in IOVs is already bound to PS. Interestingly, IOVs from both protein 4.1-depleted and HE[4.1°] membranes, which contained different amounts of ankyrin and protein 4.2, had very similar PS liposome binding capacities, suggesting that neither ankyrin nor protein 4.2 significantly influences the capacity of IOVs to bind PS liposomes. Restoration of PS liposome binding to protein 4.1 repleted IOVs demonstrated the specificity of protein 4.1 in PS liposome binding.

When the interaction of protein 4.1 with different classes of phospholipids was characterized, we found that the extent of protein 4.1 binding was directly proportional to liposome PS content. The binding of protein 4.1 to PS liposomes was saturable, at saturation $\sim 100 \ \mu g$ protein 4.1 bound/ μmol PS.

Protein 4.1 is known to interact with other membrane proteins, including spectrin-actin (30), glycophorin (13), and band 3 (42). Furthermore, PS model membranes bind to spectrin in vitro (6, 7). Therefore, it is likely that both protein 4.1 and PS have multiple membrane attachment sites. Within this

context, the interactions between protein 4.1 and PS liposomes should not be taken as representing a true physiologic situation in that competition of protein 4.1 or PS for other membrane binding sites, as would almost certainly occur in intact RBC membranes, is not considered in these simplified isolated systems. Nevertheless, our data indicates that protein 4.1 is a PS binding protein since (i) its absence from HE[4.1°] membranes is associated with a bilayer redistribution of PS; (ii) the reassociation of protein 4.1 with protein 4.1-stripped membranes results in enhanced PS binding, and (iii) protein 4.1 binds saturably to PS liposomes where the extent of protein 4.1 binding is proportional to the membrane PS content.

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