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Identification of High Affinity Folate Binding Proteins in Human Erythrocyte Membranes

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Abstract

Mature human erythrocyte membranes contained specific, high affinity ($K_a$ 3.3 x 10^-11 M) folate binding moieties. Folate binding was pH, time- and temperature-dependent, saturable, and was much greater for pteroylmonoglutamate and 5-methyltetrahydrofolate than 5-formyltetrahydrofolate and amethopterin. On detergent solubilization of membranes, two peaks of specific folate binding with $M_r$ $\geq$ 200,000 and 160,000 were identified on Sephacryl S-200 gel filtration chromatography in Triton X-100, and this corresponded to two similar peaks of immunoprecipitated material when solubilized iodinated membranes were probed with anti-human placental folate receptor antiserum. Age-dependent separation of erythrocytes by Stractan density gradients revealed a sevenfold greater folate binding capacity in membranes purified from younger compared with aged erythrocytes. Since this difference was not reflected in proportionately higher immunoreactive folate binding protein, (as determined by a specific radioimmunoassay for these proteins), or differences in affinity in younger than aged cells, these findings indicate that erythrocyte folate binding proteins become progressively nonfunctional at the onset of red cell aging.

Introduction

The intracellular role of folates as donors of one-carbon groups in enzyme reactions that lead to DNA synthesis has been well documented, and an adequate supply of folates must be assured each cell since folate deficiency leads to disordered DNA synthesis and consequent cell death (1). Therefore, specific mechanisms for cellular folate uptake and its retention could exist physiologically. This is most convincingly suggested in the developing human fetus, which concentrates and assures itself adequate folate stores even in the presence of maternal folate deficiency (1, 2). This suggests that a folate concentrating mechanism is mediated by the placenta, the organ that transfers nutrients between the maternal and fetal circulation. Based on these clinical observations, we identified, purified, and characterized a human placental particulate membrane-associated folate binding protein that had a high affinity for 5-methyltetrahydrofolate in the range for binding folates in serum (3). Although the hypothesis that these placental folate binding proteins are involved in transplacental transport of folates in humans cannot be further tested in vivo, taken together, the clinical observations and experimental studies are supporting (albeit indirect) evidence that these proteins may act as physiologically relevant specific folate receptors in human placenta (4). At the cellular level, it has long been speculated that other particulate folate binding proteins (P-FBPs) in a variety of human tissues (5-7) have some important role in folate metabolism, either as intracellular folate storage proteins or as carriers for the transmembrane transport of folates. However, only in a human nasopharyngeal carcinoma cell line in culture, has there been direct evidence for a specific transport role for such externally oriented high affinity P-FBPs in human cells (8). To identify whether folate receptors are unique to malignant human cells, it is necessary to investigate their presence in other normal human cells that have been shown to transport folate intracellularly. Earlier, antiserum to purified placental folate receptors (PFR) identified moieties with shared antigenic determinants in mature human erythrocytes (3). Since human erythrocytes can take up folates (9-11) and retain them intracellularly (1), these cells seemed a logical choice to investigate for the presence of P-FBPs. Such studies have not been performed with these or other homogeneous plasma membranes from normal human cells. A novel stable folate analogue, (a histamine derivative of folic acid), with a specific activity 40-50-fold higher than any previously available radiolabeled folate, allowed for the identification of proteins on erythrocyte membranes that have folate binding characteristics that are similar to that reported in human KB cells (12). Solubilization of radioligand-bound as well as radioiodinated membranes released folate binding proteins and iodinated proteins, respectively, that were specifically immunoprecipitated by anti-PFR antiserum. Further quantitation of these solubilized moieties by a specific, sensitive radioimmunoassay (RIA) to the PFR and correlation with the particulate membrane ligand binding assay allowed for a study of the age-related changes of folate binding proteins in young and old erythrocytes.

Methods

A histamine derivative of folic acid, 1-carboxy-3-(N-(2-(4-imidazo-1yl)ethyl)carbamoyl)-1-pteroylaminopropane, was synthesized by Du Pont/New England Nuclear, Boston, MA, and iodinated by the method of Hunter and Greenwood (13) using chloramine T and carrier free 125I. The 125I-labeled derivative 125I-labeled PteGlu, histamine

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1. Abbreviations used in this paper: BSG, buffered saline glucose, LDH, lactate dehydrogenase; PEP, phosphoenolpyruvate; P-FBP, particulate folate binding proteins; PFR, placental folate receptor; PK, pyruvate kinase; PMSF, phenylmethylsulfonyl fluoride; PteGlu, pteroylmonoglutamic acid (folic acid); 2-ME, 2-mercaptoethanol; TEA, triethanolamine; TLC, thin-layer chromatography.
Determination of the purity and stability of $^{35}$S-labeled PteGlu (histamine derivative). Since a novel radiolabeled folate compound was used, it was necessary to determine its purity and stability under various experimental conditions used to identify folate binding sites on erythrocyte membranes. Analysis for purity was determined by high performance liquid chromatography (HPLC) and thin-layer chromatography (TLC). Both techniques have been used to establish the purity of and purify folates (8). The HPLC apparatus (Waters Associates, Milford, MA) consisted of two model 510 pumps connected to a U6K injector (2 ml capacity), a (Lambda-Max model 481 LC) spectrophotometer (set at 0.005 AUFS) connected to a model 1200 linear recorder (set at 30 cm/h) and a 3.9 mm $\times$ 30 cm Bondapak C$_{18}$ stainless steel column. The column was run isocratically in 84% buffer A (5 mM Pic A, 10 mM KH$_2$PO$_4$; pH 5.5) and 16% buffer B (acetonitrile). The column flow rate was 1 ml/min and 0.5-ml fractions were collected and counted for radioactivity. Samples were routinely suspended in 250-ml columns equilibration buffer before injection. Thin-layer chromatography analysis was performed by application of radiolabeled folate samples (5 $\mu$l) to Avicel microcrystalline cellulose plates (20 $\times$ 20 cm, 250 $\mu$m, Analtech, Newark, DE) developed in N-propanol: N$_2$H$_4$OH 7:3 at room temperature. The plates were eluted radioactively and subjected to autoradiography using Kodak X-omat AR film and Cronex Lightning Plus intensifying screens at -70°C. Samples subjected to HPLC and TLC analyses were as follows: (a) commercially supplied $^{35}$S-labeled PteGlu (histamine derivative); (b) an aliquot of the peak fraction from sample a after HPLC; (c) sample b that was incubated for 2 h at 37°C. (d) To determine if the peak fraction from HPLC (which corresponded to maximum absorbance at 280 nm as well as the maximum radioactivity) was authentic $^{35}$S-labeled PteGlu (histamine derivative), its specific binding to erythrocyte membranes was compared to an equivalent amount of unfraccionated $^{35}$S-labeled PteGlu (histamine derivative); the radioactivity that was eluted by low pH (5.5) washes after binding of unfraccionated or HPLC purified $^{35}$S-labeled PteGlu (histamine derivative) to erythrocyte membranes (see below), was also analyzed by TLC to determine if the peak eluted radioactively and corresponded to authentic $^{35}$S-labeled PteGlu (histamine derivative).

Preparation of purified erythrocyte membranes. Erythrocyte membranes were prepared from whole blood collected in heparin (33 U/ml) obtained from normal human volunteers (whose reticuloocyte count was not > 1%) and 99% of leukocytes were removed as previously described (3). After four centrifuge-wash cycles at 1,000 g for 10 min at 4°C with 10 vol 0.01 M potassium phosphate (KPO$_4$), pH 7.5 containing 0.15 M NaCl (phosphate-buffered saline, PBS) per wash cycle and aspiration of the supernatant with each wash, the pellets of packed cells were lysed by the method of Lieber and Steck (14). The lysis buffer contained 5 mM sodium phosphate, pH 8, 10 $\mu$M magnesium sulfate (MgSO$_4$), and 0.1% Triton X-100 (200-fold fresh PMSF stock solution 200 $\mu$g/ml in dimethylsulfoxide), and the ratio of packed cells to lysis buffer was 1:40. The mixture was centrifuged at 30,000 g in a centrifuge (Beckman J2-21 M, Beckman Instruments, Inc., Palo Alto, CA) at 4°C for 10 min. After aspiration of the supernatant, the centrifuge-wash cycle was repeated for four times. The erythrocyte membranes were then resuspended in a volume of lysis buffer that was equal to the original volume of packed cells. Each milliliter of washed erythrocyte membranes contained an average of 2 mg protein as measured by the method of Lowry et al. (15); there was < 0.1 mg of hemoglobin associated with each milliliter of membranes as determined by the cyanmethemoglobin method using Drakkins Reagents and a stable human hemoglobin standard (16). Glyceroldehyde-3-phosphate dehydrogenase accessibility assays (14, 17) were routinely performed to ensure that the membranes were not sealed either as right-side-out ghosts or inverted vesicles and confirmed > 90% accessibility to the enzyme.

Folate binding assay. Unless specified, all incubations were for 1 h at 37°C. To determine the saturability and specificity of erythrocyte membrane folate binding, 1.3 ml of a suspension of erythrocyte membranes were mixed with increasing concentrations of $^{35}$S-labeled PteGlu (histamine derivative) diluted in 2 ml lysis buffer. After incubation, 3 ml of ice-cold lysis buffer was added and the sample was centrifuged in Biovials (Beckman Instruments) at 30,000 g for 7 min at 4°C. The supernatant was aspirated and the pellet was washed four times (10 vol/wash cycle) and counted for bound radioactivity in a Beckman gamma 5500 counter at 78% efficiency. For each level of radiolabeled $^{35}$S-labeled PteGlu (histamine derivative) used, simultaneous assays were performed containing a 1,000-fold excess of unlabeled PteGlu (in addition to $^{35}$S-labeled PteGlu (histamine derivative)) to determine the extent of nonspecific folate binding.

To determine the optimum pH for ligand binding, vials containing 1 ml of erythrocyte membranes were centrifuged to sediment membranes. After aspiration of the supernatant, pellets were suspended with 25 fmo/labeled $^{35}$S-labeled PteGlu (histamine derivative) in a final volume of 1 ml containing 10 amol sodium phosphate at pH ranging between 4.5 and 9.25. (The pH was adjusted in stock solutions by adding 0.1 M dibasic sodium phosphate to 0.1 M monobasic sodium phosphate). After incubation, the pellets were washed with the same pH buffer four times. After determination of the amount of membrane-bound radioligand at varying pH's, the pellets were washed with 10 vol of lysis buffer (pH 8) for two cycles, (to uniformly neutralize the pH in membranes that were previously washed with various pH buffers) and mixed with 25 fmo/labeled PteGlu (histamine derivative) in 1 ml lysis buffer (pH 8). After incubation, the pellets were washed with lysis buffer (pH 8) and counted for bound radioactivity. All folate binding assays were routinely performed in triplicate and the data are expressed as a mean of three experiments. There was < 5% deviation of folate binding from the mean for each set of assays. Erythrocyte membranes that were washed with lysis buffer pH 5.5 (to dissociate bound endogenous folate) were also assayed for the presence of endogenous (bound) folate as previously described (8). The upper limits of sensitivity of this assay was 50 fmo/lolate (range 50 fmo/l-5 pmol) using $^{35}$S-labeled PteGlu (histamine derivative) as the radioligand. Membranes were prepared for this analysis as follows: 10 ml of membranes that were first washed with lysis buffer pH 5.5 as described above, were then resuspended in lysis buffer, pH 8, and sedimented in a Beckman L8-55M ultracentrifuge (SW 55 rotor) at 55,000 rpm for 1 h at 4°C. The supernatants were aspirated and the pellets resuspended in a final volume of 3 ml in lysis buffer containing 2% NP-40 (vol/vol), 1% Triton X-100 (vol/vol) and 2 mg PMSF. After incubation on a shaking platform for 24 h at 4°C, the mixture was centrifuged at 55,000 rpm for 30 min, and the supernatant filtered through a 0.22- $\mu$m
filter (Millex-GV, Millipore Corp., Bedford, MA) to retain insoluble particulate material. The filtrate was incubated with 250 fmol 125I-labeled PteGlu (histamine derivative) for 1 h at 37°C, cooled to 4°C, and applied to a 2.5 × 30 cm Sephacryl S-200 gel filtration column that was equilibrated and eluted with 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100 at 4°C. The column was calibrated with [methyl-14C]-methyleneated marker proteins (New England Nuclear) as previously described (18). Fractions of 2.75 ml were collected and counted for radioactivity. Recovery of the applied radioactivity in eluted fractions was > 85%. A duplicate sample was prepared as above except that during incubation with 125I-labeled PteGlu (histamine derivative) a 1,000-fold excess of unlabeled PteGlu was also added to the mixture to determine nonspecific 125I-labeled PteGlu (histamine derivative) binding.

Antigenic identity of erythrocyte FBP's. To determine if the solubilized erythrocyte FBP's were antigenically related to PFR (3), human PFR was purified to apparent homogeneity, based on a single band on sodium dodecyl sulfate polyacrylamide gel electrophoresis, and antiserum to the purified protein was raised in rabbits as previously described (3). 1 ml of erythrocyte membranes was iodinated by the method of Markwell and Fox (19) using 400 μg Igodo-gen and 1 μCi 125I Na. After transfer of the quenched reaction mixture to a 15-ml Corex tube, the iodinated membranes were washed with 16 centrifuge-wash cycles with lysis buffer (30 vol per wash cycle). Incorporation of radioiodine into erythrocyte membranes ranged between 55 and 70% in five different experiments. The sample was solubilized with 2% Triton X-100 and 2% NP-40 and subjected to gel filtration chromatography, as described above. 10 μl of crude rabbit antiserum raised against the purified human PFR was added to 1.350 ml of each gel filtration fraction. The mixture was incubated on a shaking platform for 16 h at 4°C, following which 100 μl IgG Sorb was added to each tube. After incubation for 30 min at 4°C, the samples were centrifuged at 30,000 g for 60 min at 4°C. The supernatants were aspirated and the pellets were washed with 30 vol of lysis buffer for three cycles and counted for radioactivity. Simultaneous studies were performed on the second aliquot (1.350 ml) of each eluted fraction using 10 μl crude rabbit preimmune serum. Specific immunoprecipitation in each fraction was defined as the difference between total and nonspecific counts precipitated by immune anti-human PFR serum and by preimmune serum, respectively.

To determine if the moiety immunoprecipitated by anti-PFR serum was a folate binding protein, tubes containing 1 ml of ghosts that were pretreated to remove endogenous folate, as described above, were incubated with 35 pM 125I-labeled PteGlu (histamine derivative) in lysis buffer, pH 8, for 1 h at 37°C. After removal of unbound radioligand by four centrifuge-wash cycles, the pellets were counted for radioactivity. Each tube was subsequently solubilized with 1% deoxycholate (vol/vol) and 2% Triton X-100 (vol/vol) in a final volume of 3 ml at 4°C for 18 h on a shaking platform. The samples were then centrifuged at 30,000 g for 30 min at 4°C to sediment insoluble material. The supernatants (2.9 ml) from each tube were subsequently incubated for 18 h at 4°C on a shaking platform with increasing concentrations of crude anti-human FFR serum diluted into a final volume of 100 μl with crude rabbit preimmune serum. 1 ml of 20% polyethylene glycol (wt/vol) was then added to each tube and the mixture incubated for 10 min at 4°C. After centrifugation at 30,000 g, for 1 h at 4°C, the supernatants were aspirated and the pellets were counted for radioactivity. Simultaneous control studies were performed as described above except that increasing concentrations of preimmune serum were substituted for immune serum. Specific immunoprecipitation was determined as defined above.

Radioimmunoassay for placental folate receptors. PteGlu-Sephrose was prepared as previously described (3). 10 μl of PFR was incubated with a 1:15 dilution of PteGlu-Sephrose in a final volume of 50 μl in 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100 on a shaking platform at 4°C for 18 h. The PteGlu-Sephrose with bound PFR was washed five times with 1.5 ml 0.01 M KPO₄, pH 7.5, (1.5 ml per wash cycle) in a microcentrifuge (model 59A, Fisher Scientific, Pittsburgh, PA) at top speed for 2 min at 22°C. The beads were resuspended in 1 ml of 0.01 M KPO₄, pH 7.5 and iodinated with 1 μCi 125I Na and 5 μg lodo-gen in a 20-ml glass scintillation vial, as described (19). After incubation with the reaction mixture for 20 min at 22°C on a shaking platform, the contents were applied to a 0.5 × 10 cm Econo-Column (Bio-Rad Laboratories, Richmond, CA). The retained beads were successively washed with 1 liter of 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100 and 25 ml of 0.001 M KPO₄, pH 7.5, containing 1% Triton X-100. The iodinated PFR that remained bound to PteGlu-Sepharose was finally eluted with three successive batches of 500 μl 0.2 M glacial acetic acid containing 1% Triton X-100 and the eluate was collected in its entirety. After 1 M KPO₄, pH 7.5 was added to the eluate to adjust the final concentration to 100 mM KPO₄, the pH was adjusted to 7.5 by the addition of 1 N NaOH. The sample was subsequently analyzed by Sephacryl S-200 gel filtration (8), sucrose density gradient ultracentrifugation in the presence of detergent (18) (data not shown) and sodium dodecyl sulfate polyacrylamide gel electrophoresis of a detritonized sample (3) (Fig. 8 A). Since all three techniques revealed a single major iodinated species corresponding to a Mᵣ of 40,000, the 125I-placental folate receptor (125I-PFR) was therefore used as the radiolabeled antigen in the radioimmunoassay. Assuming a 50% recovery of non-denatured 125I-PFR that remained bound to PteGlu-Sepharose, the specific activity of the iodinated receptor was 1.6 μCi/μg protein. A final reaction mixture for the standard curve contained 5 μl KPO₄, pH 7.5, 1% Triton X-100 (vol/vol), 0.2 μl of crude rabbit antiserum to the placental folate receptor diluted in 20 μl rabbit preimmune serum, increasing concentrations of purified placental folate receptor (stock solution 1 mg/ml in 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100) diluted in 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100. After incubation for 18 h at 4°C on a shaking platform, 10,000 cpm 125I-PFR was added and incubation was carried out for another 18 h at 4°C. IgG Sorb (200 μl) was then added and the mixture incubated for another 2 h after which the IgG Sorb was sedimented in the microcentrifuge for 2 min at top speed. After aspiration of the supernatant, the pellet containing 125I-antigen-antibody complexed to IgG Sorb was counted for radioactivity. Each level of unlabeled PFR in the standard curve was performed in triplicate and the results expressed as the mean of the three assays. There was < 5% deviation from the mean at each point on the standard curve. For unknown samples subjected to evaluation for crossreacting material to the PFR, the sample solubilized in 2% Triton X-100 and 1% deoxycholate was added in place of unlabeled PFR. Pilot studies (not shown) revealed that 2% Triton X-100 and 1% deoxycholate did not affect the standard curve with respect to altering its sensitivity when compared with a simultaneous RIA performed in 1% Triton X-100. For unknown samples, a dose response (in triplicate) was routinely assessed to ensure that the assayed material was reasonably measurable in the mid-range of the standard curve.

Density gradient and age dependent separation of erythrocytes. Stractan (arabinogalactan) was dissolved in distilled water (wt/wt) and passed through an Amberlite MB-4B column (2.5 × 30 cm) twice to remove impurities and clarify the solution, as described earlier (20). This process reduced the osmolarity to ~ 100 mosM/liter. BSA (bovine serum albumin) powder (3 g) was added to the solution, which was further diluted to 291±1 mosM/liter by the addition of buffered saline glucose (BSG) (20). The osmolarity and density of the solution were determined by an (model 2001, Precision Systems, Framingham, MA) osmometer and a refractometer (Abbe 3L, Fisher Scientific), respectively. Sucrose and Stractan solutions had the same osmolarity and density (data not shown). Stock solutions were further diluted with BSG to obtain the final concentrations given below. The osmolarity of each diluted sample was maintained at 291 mosM/liter, and each solution was prepared fresh before the experiments and with desired densities determined by refractometry.

2-ml Stractan solutions (23–19%) were successively layered on each other in a 15-ml Beckman polyallomer tube, and 1.5 ml of freshly processed packed cells were layered on the top of the density gradient and separated using an SW 40 rotor for 55 min at 52,000 g at 4°C in a Beckman L8-55M ultracentrifuge. After centrifugation, each layer was
carefully removed (beginning from the top) manually, or by using an 18-gauge needle attached to a peristaltic pump (both methods gave similar results).

$^{125}$I-Labeled PteGlu (histamine derivative) binding to erythrocyte membranes was determined, as described above, after erythrocytes from each of seven Strataclean density-gradient-separated layers from 18 individual gradient tubes were pooled, washed three times with PBS, and then adjusted to a similar volume (2 x 10^7 cells per fraction) before lysis in 15-ml Corex tubes. Radioligand binding and pyruvate kinase assays were performed within 4 h of venipuncture. These membranes were washed with lysis buffer, pH 5.5, to expose cryptic folate binding sites, as described above; this ensured that the observed radioligand binding was to both exposed and unmasked (cryptic) folate binding sites. Enzymatic measurements were determined at 25°C at 340 nm, in a Perkin-Elmer Lambda 7 UV/VIS spectrophotometer (Perkin-Elmer Corp., Norwalk, CT). Each assay was performed in triplicate on a diluted hemolysate obtained through saponin treatment (21) of each layer of density-separated packed red cells and diluted with PBS. The assay system contained TEA buffer (85.6 mM, pH 7.6), PEP 0.521 mM (dissolved in 0.005 M MgSO4/0.2 M KCl), ADP 6.7 mM (neutralized with 1 M KOH), and LDH 9.5 U/ml. The reaction was initiated by the addition of 50 µl of hemolysate (1:20) and the decrease in absorbance was recorded over 5 min at 22°C. The slope of each recording was calculated and activity was expressed in units per milligram of hemoglobin using an extinction coefficient of NADH = 6.22 (21).

Red cell and leukocyte counts were determined using a Coulter counter (model M 430, Coulter Electronics, Hialeah, FL). Manual counting of reticulocytes were routinely determined among 10,000 cells after staining with brilliant cresyl blue (22). Erythrocyte membranes from red cells separated on the basis of age were subsequently solubilized with 2% Triton X-100 and 1% deoxycholate, filtered through 0.2-µm Milliflex-GV filters and analyzed for protein content (23). These samples were also subjected to the RIA to determine cross-reactive material to the PFR as described above. Routinely, a dose response for each sample was assessed and the mean of three independent results (nanograms of cross-reacting material per milligram protein) was determined. A normal volunteer was subjected to phlebotomy of 300 ml whole blood per week for 2 mo while on a normal diet. This resulted in an increase in reticulocyte count to 2.6%. This individual's blood was used for study of erythrocyte FBPs after age-dependent separation of cells.

**Results**

Purity and stability of $^{125}$I-labeled PteGlu (histamine derivative). Preliminary studies using commercially available or synthesized $[^3H]$PteGlu and 5-methyl-$[^3H]$tetrahydrofolate were unable to identify specific folate binding to erythrocyte membranes due to the small number of these exposed folate binding moieties on these cells. Only with the availability of a folate derivative with ~ 40–50-fold higher specific activity, was it possible to demonstrate P-FBPs on erythrocyte membranes. The structure of $^{125}$I-labeled PteGlu (histamine derivative) is shown in Fig. 1. The location of $^{125}$I is on the histamine ring, although its exact position has not been determined. Although this is an unnatural form of radiolabeled folate, its affinity for other FBPs from human placenta and milk was comparable to that obtained using $[^3H]$PteGlu or $[^4C]$PteGlu in pilot studies. Studies were therefore undertaken to establish the purity of $^{125}$I-labeled PteGlu (histamine derivative) as well as its stability under the experimental conditions used in these studies. This was accomplished by conventional analytical procedures with HPLC and TLC. Fig. 2 shows the HPLC elution profile of commercially supplied $^{125}$I-labeled PteGlu (histamine derivative). There was a single peak of radioactivity that corresponded to the maximum absorbance (A_{280}) of $^{125}$I-labeled PteGlu (histamine derivative), indicating its purity; when the peak fraction was reinjected into the column, the elution profile was identical to that of the commercially available $^{125}$I-labeled PteGlu (histamine derivative). Furthermore, there was no change in the elution profile or additional peaks of radioactivity after incubation of the HPLC-purified radiolabeled material for 2 h at 37°C. These findings confirmed that $^{125}$I-labeled PteGlu (histamine derivative) was greater than 99% pure and stable over the entire period of experiments detailed below. To verify purity by another method, both unfractionated (commercially available) and HPLC-purified $^{125}$I-labeled PteGlu (histamine derivative) were analyzed by TLC. As shown in Fig. 3, both preparations were identical in their migration on microcrystalline cellulose. In the unfractionated sample however, there was a faint additional band (which did not correspond to free $^{125}$I and is poorly reproduced in the figure) comprising < 1% of the major species; this was not found in the HPLC fractionated sample. To determine if the major form of $^{125}$I-labeled PteGlu (histamine derivative) from the unfractionated preparation exhibited comparable binding to erythrocyte membranes as HPLC-purified $^{125}$I-labeled PteGlu (histamine derivative), a dose-response curve with increasing concentrations of the two preparations were deter-

**Figure 1.** Structure of a derivative of folate acid for iodination, 1-carboxy-3-(N-(2-(4-imidazolyl)ethyl)carbamoyl)-1-pyrolaminopropene ($^{125}$I-labeled PteGlu (histamine derivative)).

**Figure 2.** HPLC analysis of $^{125}$I-labeled PteGlu (histamine derivative). (A) 50 fmol (10 µl) of unfractionated (commercially available) radiolabeled folate was applied to a 3.9 mm x 30 cm C_{18} µBondapak column that was isocratically equilibrated and eluted at a flow rate of 1 ml/min with 5 mM PiCA, 10 mM KH_{2}PO_{4}/H_{3}PO_{4}, pH 5.5 (buffer A), and acetoniurile (buffer B) at 84/16%, respectively. Absorbance at 280 nm (0.05 AUFS) was monitored and 0.5 ml fractions were collected and counted for radioactivity (c). (B) The peak fraction from (A) was reapplied to the column before (a) and after (b) incubation for 2 h at 37°C and fractions were counted for radioactivity.

714  A. C. Antony, R. S. Kincaide, R. S. Verma, and S. R. Krishnan
sponded to purity (histamine derivative). 5 μl of sample was applied to microcrystalline cellulose plates that were developed in N-propanol/NH₄OH 7:3 and plates were subjected to autoradiography. (A) Commercially available radiolabeled material. (B) HPLC-purified (peak fraction) ¹²⁵I-labeled PteGlu (histamine derivative). (C) Commercially available radiolabeled material that was specifically bound to erythrocyte membranes and eluted with lysis buffer, pH 5.5. (D) HPLC-purified ¹²⁵I-labeled PteGlu (histamine derivative) that was specifically bound to erythrocyte membranes and eluted with lysis buffer, pH 5.5.

Table I. Binding of Commercially Available and HPLC-purified ¹²⁵I-Labeled PteGlu (Histamine Derivative) to Erythrocyte Membranes

<table>
<thead>
<tr>
<th>Concentration of ¹²⁵I-labeled PteGlu</th>
<th>Specific ¹²⁵I-labeled PteGlu (histamine derivative) bound</th>
<th>Commercially available radioligand</th>
<th>HPLC-purified radioligand</th>
</tr>
</thead>
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<tr>
<td>pM</td>
<td>fmol</td>
<td>fmol</td>
<td>fmol</td>
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<td>2.0</td>
<td>1.8</td>
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</table>

* Erythrocyte membranes, 0.5 ml, which were prewashed to remove endogenous bound folate as described in Methods, were incubated with various concentrations of commercially available and HPLC-purified ¹²⁵I-labeled PteGlu (histamine derivative) for 1 h at 37°C in a final volume of 1 ml lysis buffer, pH 7.5, in the absence (total binding) and presence of a 1,000-fold excess unlabeled PteGlu (non-specific binding). Unbound radioactivity was removed by three centrifuge-wash cycles with 10 vol of lysis buffer, pH 7.5, and the pellets were counted for radioactivity. Specifically bound ¹²⁵I-labeled PteGlu (histamine derivative) was the difference between total and non-specific binding and constituted > 90% of total binding.

Each data point represents the mean of three values that had < 5% variation from the mean.

fractions (HPLC) and major species (> 99%) of radiolabeled material (TLC); (b) this commercially available ¹²⁵I-labeled PteGlu (histamine derivative) bound erythrocyte membranes with a high degree of specificity and to a comparable extent as HPLC-purified radioactive material; and (c) following specific binding to erythrocyte membranes by commercially available and HPLC-purified ¹²⁵I-labeled PteGlu (histamine derivative) and subsequent elution of bound radioactivity, there was no detectable change in the relative migration of eluted radioactivity on TLC when compared with HPLC-purified radioligand, indicating that authentic ¹²⁵I-labeled PteGlu (histamine derivative) was responsible for binding to erythrocyte membranes.

Erythrocyte membrane folate binding characteristics. As shown in Fig. 4 A, washed purified human erythrocyte membranes specifically bound ¹²⁵I-labeled PteGlu (histamine derivative), and non-specific binding accounted for < 5% of total folate binding. Specific folate binding, defined as the difference between radioligand binding in the presence (non-specific) and absence (total) of a 1,000-fold excess unlabeled PteGlu, was saturable, while non-specific folate binding was linear and not saturable. The binding isotherms indicated a single high affinity folate binding site with a dissociation constant of 3.3 × 10⁻¹¹ M (Fig. 4 B). Folate binding linearly increased as a function of erythrocyte membrane concentration (Fig. 4 C), and was time and temperature dependent; ligand binding equilibrium was reached by 1 h at 37°C and 6 h at 4°C (Fig. 4 D). The pH optimum for ligand binding was 7.5 (Fig. 4 E, closed bars). To determine if the observed decrease in folate binding at lower pH was due to alteration in ligand binding capacity or to destruction of folate binding sites (as a result of washing membranes with low pH buffers), the pH of membranes were neutralized and folate binding studies were repeated. Under these conditions there was up to a fourfold increase in folate binding.
bound to samples prewashed with low pH buffers when com-
pared to samples maintained at pH 8.0. (Fig. 4 E, open bars).
This increase in folate binding was not due to an increase in
affinity, which was identical to that shown in Fig. 4B ($K_d = 3.3$
$\times 10^{-11}$ M) and was likely due to unmasking of folate binding
sites that were occupied by endogenous folate (cryptic folate
binding sites). This possibility was confirmed by washing
membranes containing bound radiolabeled folate with pH 5.5
buffer, which led to $\sim 90\%$ release of the bound radioligand
(data not shown). These studies support the conclusion that
prewashing membranes with low pH buffers resulted in endog-
igenous folate removal and unmasking of cryptic folate binding
sites. The equivalent affinity between exposed and total (ex-
posed and unmasked cryptic) folate binding sites also sug-
gested that exposed and cryptic folate binding sites have func-
tional similarities. It is not known if a significant number of
erthrocyte membrane folate binding sites are occupied by
endogenous folates in vivo.

Since a derivative of folate [$^{125}$I-labeled PteGlu (histamine
derivative)] was used as the radioligand in this study, it was
important to determine if its affinity for particulate erythro-
cyte membrane FBPs was similar to authentic pteroylmonog-
lutamate. The results of the competitive inhibition studies
using unlabeled PteGlu (Fig. 5) suggested that erythrocyte P-
FBPs had similar affinity for the commercially available io-
dinated histamine derivative of PteGlu as for unlabeled
PteGlu. This was indicated by the ability of equimolar concen-
trations of unlabeled PteGlu in reducing $^{125}$I-labeled
PteGlu (histamine derivative) binding to erythrocyte mem-
branes by 50%. The results with various folate analogues indi-
cated higher affinities for PteGlu and 5-methyltetrahydrofolate
than 5-formyltetrahydrofolate and amethopterin. Under these

Figure 4. Erythrocyte membrane folate binding characteristics. (A)
saturability and specificity; Erythrocyte membranes (1.3 ml) were in-
cubated with increasing concentrations of $^{125}$I-labeled PteGlu (hista-
mine derivative) in the absence (c) and presence (a) of a 1,000-fold
excess of unlabeled PteGlu in a final volume of 2 ml lysis buffer con-
sisting of 10 mmol sodium phosphate, pH 8, and 20 mmol magnesi-
um sulfate, 40 $\mu$g PMSF, and membranes washed in lysis buffer
were counted for radioactivity. (B) affinity: The dissociation constant
of folate interaction with membranes. (C) Substrate dependence: 50
fmol $^{125}$I-labeled PteGlu (histamine derivative) were incubated with
increasing concentrations of membranes in a final volume of 1 ml
lysis buffer. (D) Time and temperature dependence: membranes,
(2.75 ml), were incubated at 4°C (a) or 37°C (b) with 25 pM $^{125}$I-la-
beled PteGlu (histamine derivative) in 3 ml lysis buffer and at var-
ious times the membrane pellets assayed for bound radioactivity. (E)
olarity: membranes (1 ml), were incubated with 25 fmol
$^{125}$I-labeled PteGlu (histamine derivative) in lysis buffers of varying
pH. After washing membranes with the same pH buffer that they
were incubated with radioligand, the pellets were counted for radio-
activity (solid bars). These membranes were subsequently washed
with lysis buffer, pH 8, and reincubated with 25 fmol $^{125}$I-labeled
PteGlu (histamine derivative); the results of specific radioligand up-
take under these conditions (open bars). Unless specified, all incuba-
tions were for 1 h at 37°C and unbound folate removed by centrifuge-
wash cycles as described under Methods.
experimental conditions, it would suggest that d,1-5-methyltetrahydrofolate had an ~ 5-10-fold less affinity for binding erythrocyte membranes compared with PteGlu. When this is related to human serum folate concentrations (5-40 nM) and the dissociation constant of folate by these membranes, it appears that this affinity is in the range for binding folates in serum. Similar findings have been reported with particulate folate binding proteins in human placenta, milk, and KB cells and other mammalian cells (3, 5-7, 12, 18, 24-29), further supporting the possibility that these proteins may have some physiological relevance in binding serum folates.

**Solubilization and identification of membrane FBP s.** We found that 2% NP-40 and 2% Triton X-100 led to the solubilization of specific folate binding moieties that eluted on Sephacryl S-200 in the void volume (M ≈ 200,000) as well as at M = 160,000 (Fig. 6 A). The third radioactive peak at V/Vo = 3.5 represented the elution position of free 125I-labeled PteGlu (histamine derivative). Duplicate studies of membranes solubilized and incubated with similar 125I-labeled PteGlu (histamine derivative) concentrations but also containing a 1,000-fold excess of unlabeled PteGlu resulted in virtually complete quenching of the folate binding peak corresponding to M = 160,000, similar to those results found among FBPs identified in human placenta, milk, and KB cells (3, 8, 18). Although there was a significantly higher amount of nonspecific binding by the minor peak eluting in the void volume, there was nevertheless a peak of specific folate binding in this fraction that could be due to incomplete solubilization of this moiety out of erythrocyte membranes. The greater degree of nonspecific binding, although possibly unique to this FBP in comparison to FBPs from other human tissues (3, 8, 18) may also be explained by the coelution of other ≥ 200,000 M, proteins that have nonspecific folate binding properties. No FBP was identified in the M = 40,000 range, which was the elution position for the major solubilized membrane FBP from human placenta (3); these were similar results, however, to those found with human KB cells that had a single 160,000 M, FBP (8).

Studies were further extended to determine if the eluted FBPs after Sephacryl S-200 gel filtration were recognized by antibody to the purified human placental folate receptor (3). Although the 125I-labeled PteGlu (histamine derivative)-bound FBPs eluting from the gel filtration column could be specifically immunoprecipitated with purified IgG from immune serum raised against the FPR using either 5% polyethylene glycol or proteins A bearing S. aureus cells, the quantity of specifically precipitated FBPs was consistently lower in comparison to the total radioactivity in each FBP peak (data not shown). The reason for this was not clear but could be related

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**Figure 5.** Competitive inhibition of 125I-labeled PteGlu (histamine derivative) binding to erythrocyte membranes by unlabeled folates. Membranes (1 ml) were suspended with 50 fmol 125I-labeled PteGlu (histamine derivative) in the absence (maximum bound) and presence of increasing concentrations of unlabeled PteGlu (a), d,1-5-methyltetrahydrofolate (c), 5-formyltetrahydrofolate (w) or amethopterin (a) in a final volume of 2 ml of lysis, pH 8, buffer containing 200 µmol 2-ME. After incubation for 1 h at 37°C, the membranes were washed and assessed for bound radioactivity.

**Figure 6.** Gel filtration chromatography and identification of erythrocyte membrane FBPs. (A) 20 ml of erythrocyte membranes were treated to remove endogenous folate (see Methods), concentrated, and solubilized with 2% Triton X-100 and 2% NP-40 for 24 h at 4°C. The supernatant (3 ml) after ultracentrifugation was filtered, and incubated with 250 fmol 125I-labeled PteGlu (histamine derivative) (8.6 × 10^3 cpm) in the absence (a) and presence (a) of 250 pmol unlabeled PteGlu and applied over a 2.5 × 30 cm Sephacryl S-200 column that was equilibrated and eluted with 0.01 M KPO4, pH 7.5, containing 1% Triton X-100. Fractions of 2.75 ml were collected and counted for radioactivity. The recovery of total applied radioactivity in eluted fractions was 90%. (B) 1 ml of erythrocyte membranes were iodinated, solubilized, and subjected to gel filtration as above. Each eluted fraction was divided into two 1.350-ml aliquots and incubated with either 10 µl crude rabbit anti-human placental folate receptor immune serum or preimmune serum for 16 h at 4°C. After adding IgG Sorb, (100 µl), and incubation for 30 min at 4°C, the mixture was centrifuged to sediment IgG Sorb bound antibody complexed to iodinated proteins. The data in each eluted fraction is the difference between the precipitated radioactivity obtained using immune serum and preimmune serum. Both qualitatively and quantitatively similar results were obtained on five different studies on blood from the same individual.
to a higher proportion of blocking compared to immunoprecipitating antibodies in the polyclonal antiserum raised against the purified protein; these blocking antibodies could potentially displace bound $^{125}$I-labeled PteGlu (histamine derivative) from the FBPs. Therefore, after iodination of erythrocyte membranes followed by solubilization and Sepharcl S-200 gel filtration chromatography, the eluted fractions were probed with rabbit immune and preimmune serum. The results (Fig. 6 B), identified two peaks of specifically immunoprecipitated material in the void volume and at $M_r$ 160,000 corresponding to the elution position of $^{125}$I-labeled PteGlu (histamine derivative)-bound FBPs (Fig. 6 A). Although solubilized iodinated membranes contained a moiety that was specifically immunoprecipitated by anti-FPR serum, findings consistent with earlier immunofluorescent studies (3), it was nevertheless necessary to determine if this species was a folate binding protein. We found that 90% of the $^{125}$I-labeled PteGlu (histamine derivative) previously bound to membranes was released into the supernatant after treatment of membranes with 2% Triton X-100 and 1% sodium deoxycholate. When the supernatant was reacted with increasing concentrations of anti-FPR anti-

Figure 7. Immunoprecipitation of solubilized erythrocyte folate binding protein by anti-human placental folate receptor antiserum.

Specific cpm precipitation

ANTI-HUMAN PLACENTAL FOLATE RECEPTOR ANTISERUM

In summary, there was a linear increase in specific precipitation of $^{125}$I-labeled PteGlu (histamine derivative)-bound protein (Fig. 7) which, at $100$ uL antiserum, constituted 38% of the total radioactivity. The inability to precipitate 100% of protein bound radioligand could be due to (a) competitive displacement of radioligand by unlabeled folates present in rabbit antiserum, or (b) partial homology of antigenic determinants between the erythrocyte FBP and PFR. The fact that the crude antiserum contained a higher concentration of folate (100 nM) compared with 37.5 nM folate present in preimmune serum could explain the former possibility. However, since preimmune serum precipitated < 5% of the total radioligand-bound protein at all levels compared to antiserum, the results nevertheless indicate that the erythrocyte FBP shared epitopes with the PFR. Taken together, the ligand binding data as well as the immunological studies suggest that the two erythrocyte membrane folate binding proteins were antigenically related to the FBPs from human milk, placenta, and KB cells, which share antigenic determinants that are recognized by the antiserum used in these studies (3, 8, 18–19). The purified FBPs with apparent $M_r$ of 160,000 on gel filtration in Triton X-100 from human milk (18) and KB cells (8, 24, 30) are hydrophobic detergent binding proteins, findings that are consistent with their association with membranes; it is therefore possible that the erythrocyte 160,000 $M_r$ FBPs may have similar characteristics. Although the $\geq 200,000$ $M_r$ FBP may be unique to erythrocyte membranes, a more trivial explanation may be the incomplete solubilization from erythrocyte membranes by the non-ionic detergents used in this study. Further characterization of detergent binding and the $M_r$ of these proteins by native and denaturing gel electrophoresis has been attempted; preliminary studies indicate, however, that further purification of these proteins is required due to the small amounts of these P-FBPs in erythrocytes (see below).

Identification of nonfunctional FBPs. Scatchard analysis from the folate binding isotherms indicated that in spite of removal of endogenous folate by "acid stripping" washes, there was less than one functional (in folate binding) molecule of FBP per erythrocyte. These results were not consistent with previous immunofluorescence studies on erythrocytes (3) that revealed both linear and beaded fluorescence in virtually every cell (albeit in varying degrees) when probed with anti-human PFR serum. A gross underestimation of functional folate binding protein could be possible if a large amount of endogenous folate remained bound even after acid washing of membranes. However, when purified membranes (from 10 ml of erythrocytes) were assayed for the presence of endogenous folate by boiling for 30 min in the presence of reducing agents (8, 18), no folate was released (detected) in the supernatant. To determine whether there was folate that remained bound to membrane FBPs even after the boiling process, two (indirect) tests were employed to investigate if FBPs were denatured and lost the ability to bind folate. In the first instance, membranes failed to bind $^{125}$I-labeled PteGlu (histamine derivative) after they were boiled; secondly, boiling membranes released 100% of bound $^{125}$I-labeled PteGlu (histamine derivative). Thus it was unlikely that endogenous folate remained bound to membrane FBPs to account for our inability to detect it in the assay for endogenous folate. We therefore hypothesized the presence of nonfunctional but immunoreactive folate binding proteins in these cells.
Cytes of cyte. It was determined that all of the observed folate binding was accounted for by contaminating reticulocytes in the relatively heterogeneously aged population of erythrocytes in a given sample of blood. We determined (a) the folate binding capacity and (b) amount of immunoreactive FBP in membranes from cells separated on the basis of age. The age-related fractionation of leukocyte-free red cells was based on earlier findings.

To test this possibility, a RIA was constructed that was based on the competitive binding of purified iodinated (Fig. 8 A) and unlabeled placental folate receptor by crude rabbit anti-human PFR serum; 125I-antigen-antibody complexes were immunoprecipitated with protein A-bearing Staphylococcus aureus cells. The standard RIA assay curve (Fig. 8 B) had an upper limit of sensitivity of 5 ng (0.1 pmol) and a lower limit of sensitivity of 500 ng (10 pmol) PFR. When solubilized erythrocyte membranes were tested in the RIA, they were found to contain 160±20 ng cross-reacting material/mg protein; this constituted 0.016% of the total solubilized erythrocyte membrane proteins. When the value of cross-reacting material per milligram of protein was converted to molar equivalents of FBPs and thence to molecules of protein per erythrocyte, assuming that the true M₀ of the FBP was ~44,000 as has been shown with solubilized particulate FBPs from milk [18] and KB cells [30], there were ~1,500±150 molecules/erythrocyte. It was therefore evident that there was a significant component of nonfunctional but immunoreactive FBP in erythrocytes.

Relationship between folate binding protein and red cell age. To determine if all of the observed folate binding was accounted for by contaminating reticulocytes in the relatively heterogeneously aged population of erythrocytes in a given sample of blood, we determined (a) the folate binding capacity and (b) amount of immunoreactive FBP in membranes from cells separated on the basis of age. As described in Methods were assessed for folate binding (o) using 0.1 nM 125I-labeled PteGlu (histamine derivative). Control (unfractionated) erythrocyte membranes (from 2×10⁶ cells) bound 1.1 fmol 125I-labeled PteGlu (histamine derivative). Membranes of known protein content from each separate were solubilized and assessed for cross-reacting material to the PFR by RIA (e). Inset shows the Pyruvate kinase activity and percent reticulocytes in each erythrocyte fraction after density gradient separation. -/+σ on the abscissa shows the position in the gradient from top (-σ youngest cells) to bottom (+σ aged cells). Mean cell age was calculated as previously described (31). Each data point was the mean±SE of experiments performed in triplicate. Similar results were obtained on three different occasions.

**Figure 8.** (A) Sodium dodecyl sulfate polyacrylamide gel electrophoresis and autoradiography of iodinated placental folate receptor 100,000 cpm of affinity purified 125I-PFR (see Methods) were detritonized and run on 15% slab gels. After drying, the gels were subjected to autoradiography using Kodak X-Omat AR film and Cronex Lightning-Plus intensifying screens for 8 h at ~70°C. The gel was run from left to right and the major band corresponds to a M₀ = 40,000, while the minor band (containing aggregated material that failed to enter the gel) corresponds to the origin. (B) Standard RIA curve for the PFR. Tubes containing 20 μl anti-human PFR serum (diluted 1:100 in preimmune serum) and increasing concentrations of unlabeled PFR diluted in a final volume of 500 μl 0.01 M KPO₄, pH 7.5, containing 1% Triton X-100 were incubated for 18 h at 4°C. 125I-PFR, 10,000 cpm, was then added and incubation continued for an additional 18 h. IgG Sorb, 20 μl, was subsequently added and after 2 h at 4°C, the 125I-PFR-antibody complexes bound to IgG Sorb were sedimented and counted for radioactivity. Preimmune serum, 20 μl, nonspecifically precipitated 28% of the total 125I-PFR, and this value, when subtracted from that precipitated with 0.2 μl immune serum, gave the 100% value (4,000 cpm) on the ordinate.

**Figure 9.** 125I-Labeled PteGlu (histamine derivative) binding to erythrocyte membranes and RIA on solubilized membranes from erythrocytes separated by Stractan density gradients. 2×10⁶ cells from each of seven layers of erythrocytes separated by Stractan density gradients were lysed and after exposure of cryptic folate binding sites as described in Methods were assessed for folate binding (o) using 0.1 nM 125I-labeled PteGlu (histamine derivative). Control (unfractionated) erythrocyte membranes (from 2×10⁶ cells) bound 1.1 fmol 125I-labeled PteGlu (histamine derivative). Membranes of known protein content from each separate were solubilized and assessed for cross-reacting material to the PFR by RIA (e). Inset shows the Pyruvate kinase activity and percent reticulocytes in each erythrocyte fraction after density gradient separation. -/+σ on the abscissa shows the position in the gradient from top (-σ youngest cells) to bottom (+σ aged cells). Mean cell age was calculated as previously described (31). Each data point was the mean±SE of experiments performed in triplicate. Similar results were obtained on three different occasions.
that red cells became more dense as they aged and could be separated on this basis by Stractan density gradients. Parameters of aging red cells included the measurement of reticulocytes (youngest circulating erythrocytes) and the activity of the enzyme pyruvate kinase (PK), which decreases with increasing cell age. As shown in Fig. 9, inset, the concentration of reticulocytes in the fractions declined rapidly from top to bottom and virtually no measurable reticulocytes were demonstrated in the bottom fraction (which contained aged red cells), whereas the maximum reticulocytes (12%) were concentrated in the top fraction. The PK activity was also maximum in the top fraction of the gradient and progressively declined; cells in the bottom fraction contained a 10-fold lower PK activity thus confirming the validity of the gradient in separating cells on the basis of their age. Also shown in Fig. 9, a 6.75-fold higher binding capacity of $^{125}$I-labeled PteGlu (histamine derivative) was observed in membranes from young cells when compared with an equivalent number of unfractonated (control) red cells. No correlation was identified between the binding of $^{125}$I-labeled PteGlu (histamine derivative) to the various cell cohorts after fractionation and the number of reticulocytes per fraction. These findings excluded the possibility that the observed $^{125}$I-labeled PteGlu (histamine derivative) to unfractonated cells was due to contamination by reticuloocytes. However, these results also showed that the degree of $^{125}$I-labeled PteGlu (histamine derivative) binding to red cells markedly decreased at the onset of red cell aging. Scatchard analysis (32) of $^{125}$I-labeled PteGlu (histamine derivative) binding isotherms on erythrocyte membranes from the youngest cohort of erythrocytes revealed a single high affinity binding site and an essentially similar dissociation constant ($K_d = 4 \times 10^{-11}$ M) compared with unfractonated and aged erythrocytes ($K_d = 3.3 \times 10^{-11}$ M). Therefore, the observed differences in folate binding by younger erythrocytes were due to true differences in the number of functional (in folate binding) FBPs and not due to significant differences in affinity. To determine if the observed decrease in folate binding was accompanied by a proportionate decrease in the amount of immunoreactive FBP in red cells separated on the basis of age, solubilized membranes obtained from age-separated cells were analyzed by the RIA to detect cross-reacting material to PFR. The results showed that each cohort of aging erythrocytes contained essentially similar amounts of cross-reacting material (Fig. 9). Pretreatment of erythrocyte membranes with low pH (5.5) and pH 8 lysis buffer did not result in any change (< 5% variation) in the content of immunoreactive folate binding protein. When corrected for the number of the cells, the data indicated 1,500±150 molecules of immunoreactive FBP per cell. Interestingly, Scatchard analysis of ligand binding data revealed ~2 molecules of functional FBP per cell in the cohort of reticulocyte-rich young cells. (Assuming a 1:1 molar stoichiometric ratio of ligand binding), these data indicated that (a) the observed folate binding was only to a fraction of the total immunoreactive FBP per cell; (b) >99% of the immunoreactive protein was nonfunctional; (c) ligand binding studies were 100-200-fold more sensitive than the RIA to detect small differences in functional FBP in young versus aged erythrocytes. The circulating erythrocyte FBP could be degraded shortly before (or after) its release from the bone marrow if it was functional in erythroid precursors; this was not investigated in bone marrow cells. Taken together, the ligand binding studies and the RJA support the hypothesis that with the onset of red cell aging there is a rapid decrease in the functional capacity of the erythrocyte FBP to bind folate that is not due to a concomitant loss of immunoreactive FBP. The use of high specific activity $^{125}$I-labeled PteGlu (histamine derivative) has nevertheless identified that aged red cells have much lower capacity to bind folate than younger cohorts.

**Discussion**

Using a highly purified, stable, novel folate analogue, the data on high affinity, specificity, saturability, pH, time- and temperature-dependence of erythrocyte membranes as presented in this paper extends earlier observations (3) of shared antigenic determinants between the placental folate receptor and a moiety on intact erythrocytes. These specific folate binding moieties were solubilized out of erythrocyte membranes by non-ionic detergents where they eluted predominantly as 160,000 $M_r$ species. Anti-FFR antiserum also immunoprecipitated solubilized 160,000 $M_r$ iodinated membrane proteins. Evidence that this specific antibody interaction was with FBPs on erythrocyte membranes came from experiments where antiserum to PFR specifically immunoprecipitated $^{125}$I-labeled PteGlu (histamine derivative)-bound proteins in a dose-dependent manner. The data that support the possibility that these erythrocyte folate binding sites may be bound by folates present in serum was shown by experiments with "acid stripping" that uncovered cryptic folate binding sites.

The binding constants of erythrocyte membrane FBPs with various folate analogues were comparable to that reported earlier in particulate membranes from human placenta, milk, KB cells, leukemic cells, goat milk, rabbit and porcine choroid plexus, and rat kidney (3, 5-7, 12, 15, 24-29). However, studies on folate uptake systems in several other intact mammalian cells including L1210 cells (33-35), PHA stimulated lymphocytes (36), and human bone marrow cells (37) and human erythrocytes (10) suggest that transport was equally efficient and maximal for 5-methyltetrahydrofolate, 5-formyltetrahydrofolate and amethopterin, while that for PteGlu was poor. With the exception of L1210 cells, where the affinity of the transport system for folates correlates with affinity for folates by surface membrane FBPs (34), competitive binding inhibition studies (similar to that reported with P-FBPs) using crude membranes or purified folate binding proteins from these cells (lymphocytes, bone marrow cells, and erythrocytes) have not been reported. Therefore, it is not known if other unique folate transport proteins in these cells can bind and internalize various reduced folates more efficiently than PteGlu. Along these lines, we are currently investigating the basis for the apparent dichotomy between the high binding constants of erythrocyte FBPs for PteGlu and 5-methyltetrahydrofolate and the marked differences reported in the affinity of the transport system in intact human erythrocytes (10). It should nevertheless be pointed out that in our recent studies with human KB cells, membrane-associated FBPs bound PteGlu and 5-methyltetrahydrofolate with high affinity, and intracellular transport of both folates were comparable and mediated by these FBPs (8, 24, 38). Kamen and Capdevila (39) have also recently demonstrated that similar FBPs on monkey kidney cells in culture specifically bind PteGlu and 5-methyltetrahydrofolate and transport the latter by a process that is consistent with
receptor-mediated endocytosis. Thus, there appears to be no controversy regarding the relatively high binding constants of FBPs and FBP-mediated transport efficiencies for PteGlu and 5-methyltetrahydrofolate in human KB cells and monkey kidney cells when both parameters (binding and transport) were directly analyzed.

Recently, hemoglobin has been shown to possess a folate binding site (40). Evidence to rule out the possibility that the observed binding of folate to erythrocyte membranes was through hemoglobin, which contaminated the membrane preparations are as follows: (a) the dissociation constant of erythrocyte folate binding moieties for folate were more consistent with that of a high affinity, low capacity system in the nanomolar range, while folate binding to hemoglobin was more consistent with a lower affinity, high capacity system (micromolar range); (b) there was apparently no difference in \( ^{125}\text{I}\)-labeled PteGlu (histamine derivative) binding to “pink” membranes (which contained 100-fold more hemoglobin) compared to “white” membranes; (c) the specific immunoprecipitation of \( ^{125}\text{I}\)-labeled PteGlu (histamine derivative) bound to solubilized erythrocyte folate binding proteins by antiplacental folate receptor serum; and (d) the apparent molecular weight of the erythrocyte folate binding protein on gel filtration in detergent \( (M_r, 160,000) \) compared to hemoglobin. The observations that intracellular folate dissociate from binding sites on hemoglobin at lower oxygen tension while binding was increased at higher oxygen tension \( (41) \) raised the possibility that the membrane-associated folate binding proteins may somehow participate in this process. However, preliminary studies on folate binding to lysed (white) membranes under low and high oxygen tension appear not to support an alteration in affinity for folate under these circumstances (unpublished observations).

Due to the heterogenous nature of the red cells with respect to age in a given sample of whole blood (ranging from 1 to 120 d), it was necessary to determine if the observed folate binding was predominantly to younger erythrocytes or reticulocytes. When separated on the basis of age, the reticulocyte-rich fractions in a Strattract density gradient had a 6.75-fold greater folate binding capacity compared to unfraccionated cells that was not due to differences in affinity compared to aged cells. However, even older cells (containing no contaminating reticulocytes) possessed folate binding capacity. To determine if this apparent decrease in radioligand binding with erythrocyte aging was accompanied by parallel changes in immunoreactive FBP, cohorts of cells separated on the basis of age were analyzed by a RIA which measured cross-reacting material to placental folate receptors. The results indicated that the amount of immunoreactive FBP did not decrease with age. These findings suggested a progressive loss of functional FBP as erythrocytes aged. The amount of erythrocyte FBP per erythrocyte as estimated by RIA indicated \( \sim 1,500 \) immunoreactive FBP molecules per cell and constituted 0.016% of the total solubilized membrane protein. These studies also documented that this novel radioligand, the iodinated histamine derivative of folate, was at least two orders of magnitude more sensitive than the RIA in detecting small differences in the number of functional FBPs. Use of both assay methods (ligand binding to membranes and RIA on solubilized membrane proteins) has supported the hypothesis that with the onset of aging in the circulation, the erythrocyte rapidly loses its functional FBPs without a concomitant loss of the membrane FBP.

These findings are similar to those observed with the insulin receptor on erythrocytes that decrease in direct proportion to the red cell aging process (31). However, no direct studies of immunoreactive insulin receptor were reported to identify if there were similar findings as reported in our studies. The data presented for erythrocyte transferrin receptors is also different from our results. In that case (42) there is a proportionate and abrupt decrease in both transferrin binding and immunoreactive transferrin receptor when reticulocytes matured to erythrocytes. Since evaluation of transferrin receptor content was made on reticulocytes from patients with hemolytic anemias, (where the overall age of erythrocytes contaminating the reticulocytes would be younger), it is not known if the immunoreactive transferrin receptor is present in older erythrocytes as shown with FBPs. Nevertheless, immunofluorescent analysis of erythrocyte FBPs and transferrin receptors also showed major differences where FBPs were present in all red cells in varying degrees while transferrin receptors were only identified in reticulocytes. A discrepancy between the ligand binding studies and immunoreactive FBP has previously been observed with serum FBPs in normal individuals as well under various disease states (43). The significance of excess immunoreactive nonfunctional FBPs in serum is unclear; in erythrocyte membranes it may be a reflection of FBP degradation during the aging process.

The first indirect evidence for a receptor role for plasma membrane-associated folate binding moieties in vivo were performed by Zamierowski and Wagner (44) who showed that shortly after administration to rats, radiolabeled folate was first localized in liver cells on the external plasma membrane before its translocation intracellularly. McHugh and Cheng (12) subsequently confirmed folate binding to particulate membranes in cultured human KB cells. We have recently demonstrated a transport role for these particulate folate binding moieties in KB cells in the cellular uptake of 5-methyltetrahydrofolate \( (8) \). However, as shown by McHugh and Cheng (12) and confirmed by us \( (8) \), these FBPs may also be intracellular proteins that could function as folate storage proteins. The functional erythrocyte FBP is antigenically related to and appears to share most of the functional ligand binding characteristics as the 5-methyltetrahydrofolate transport proteins in KB cells \( (24) \). The striking difference however, is the quantitatively higher content of these P-FBPs in KB cells \( (12) \) and relatively minute quantities in erythrocyte membranes; the major difference being the folate requirement during the rapid growth rate of KB cells \( (\text{where a doubling time of } \sim 24 \text{ h}) \), while erythrocytes are nondividing cells that completed DNA synthesis and their requirement for folates at younger stages within the bone marrow. An issue raised by the identification of P-FBPs in mature human erythrocyte membranes is therefore related to the physiological role of these proteins. These cells contain predominantly polyglutamated forms of folate and yet do not contain the enzyme pteroyl polyglutamate synthetase, which is only present in immature red cells. This enzyme is responsible for the addition of the polyglutamate chain to 5-methyltetrahydrofolate taken up from serum. This implies that the existing folate in the mature cell was taken up at a more immature stage of erythrocyte development. This has been verified experimentally by earlier observations (and that
presented in this paper) that more folate is taken up by younger erythrocyte forms (reticulocytes) than mature cells (9, 11). However, the fact that mature erythrocytes can take up 5-methyltetrahydrofolate (10) by a specific, saturable carrier-mediated process, and the clinical observations that the intracellular folate content is 30-fold higher than of serum (1), support the possibility that the erythrocyte has the capacity to concentrate folate against a gradient by a folate transport system. Whether specific folate transport in intact erythrocytes is mediated by FBPs is currently under study.

Without the availability of purified serum folate binding proteins, we cannot rule out the possibility that the observed functional and nonfunctional FBPs in erythrocyte membranes are due to adsorption of FBPs from the serum. Such a process could arise either through specific (binding) interactions or by hydrophobic or ionic interactions with erythrocyte membrane proteins. Investigation of membrane-associated FBPs and the folate transport system in this simplest of human cells may highlight their physiological expression in more complex human cells. Finally, tissues that have previously not been found to contain functional FBPs by ligand binding studies should be reevaluated using the higher specific activity folate analogue and the RIA described here to identify their distribution in normal and diseased states.

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