Effect of Perturbation of Specific Folate Receptors during In Vitro Erythropoiesis

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Abstract

Although antisera to specific placent al folate receptors inhibits the uptake of 5-methyltetrahydrofolate into cultured malignant human cells, little is known of the functional significance of folate receptors in normal human cells. Human bone marrow cells were therefore assayed for erythropoietic burst-forming units in the presence of an antihuman placent al folate receptor serum and preimmune serum to determine the role of such a receptor in erythroid differentiation. When marrow cells were assayed in the presence of anti-receptor antiserum, there was (i) a threefold increase in erythropoietic burst formation and a twofold increase in the number of cells per erythroid burst; (ii) morphological evidence for nuclear/cyttoplasmic dissociation of orthochromatic normoblasts composing erythroid bursts (megablastic erythropoiesis); (iii) intracellular folate deficiency with a 70% reduction of intracellular folate in antiserum treated cells as compared with control cells; and (iv) complete reversal of antiserum-induced changes on preincubation of antiserum with purified human placent al folate receptor. These data support the conclusion that folate receptors on marrow cells provide an important function in the cellular uptake of folates during in vitro erythropoiesis. This process of folate uptake also appears to play a pivotal role in the differentiation and proliferation of erythroid progenitor cells.

Introduction

All human cells require folates intracellularly for enzyme reactions that are crucial to the synthesis of deoxyribonucleic acid (1). Recently, we established that specific, high affinity, externally-oriented, membrane-associated folate-binding proteins (FBPs) on the surface of malignant human nasopharyngeal carcinoma (KB) cells act as specific transport proteins in the cellular uptake of folates (2). However, little is known about the existence or function of similar proteins in normal human cells.

Earlier, we identified by indirect immunofluorescence studies that immunologically similar proteins to human placent al folate receptors (PFR) were present on circulating human erythrocytes (3). Since erythrocytes have a 30-fold higher intracellular folate concentration than serum (1), we suggested that this gradient could be mediated by specific folate transport proteins (3). Although subsequent investigations identified that human erythrocyte membranes (4), ghosts and intact cells (5), contained FBP s that had similar characteri stics of folate binding as KB cell FBPs (2), the magnitude of folate uptake in circulating erythrocytes (in contrast to KB cells) was much lower and not consistent with a physiologically significant specific FBP-mediated uptake system (5a). Furthermore, the majority of erythrocyte FBPs identified by a specific radioimmunoassay to PFR (6) were nonfunctional in that they did not bind radiolabeled folate (5a). These studies suggested that the higher intracellular folate content in the erythrocyte was probably achieved at an earlier stage in development. Using in vitro hematopoietic progenitor cell assays, we asked the following questions: (i) Is the development of erythroid progenitor and precursor cells dependent on the functional integrity of specific folate receptors? and (ii) Can perturbation of specific folate receptors on these cells lead to qualitative and/or quantitative changes consistent with folate deficiency? Our studies appear to support the conclusion that specific folate receptors mediate folate uptake in normal human marrow cells in vitro.

Methods

Iscove’s modified Dulbecco’s medium (IMDM) containing 4 mg folic acid per liter, and preservative-free sodium heparin were obtained from Gibco, Grand Island, NY. Partially purified human urinary erythropoietin (72 U/mg of protein) was obtained from Toyobo Co., New York, NY. Fetal calf serum and rabbit preimmune serum were purchased from HyClone Laboratories, Inc., Logan, UT, whereas 125I-labeled folic acid (an iodinated histamine derivative of folic acid, purity 98%, with a specific activity of 2,200 Ci/mmole) was obtained from DuPont Co., Diagnostic & BioResearch Systems, Wilmington, DE, and New England Nuclear, Boston, MA. Folic acid (99% pure), β-lactoglobulin, and monobasic and dibasic potassium phosphate were obtained from Sigma Chemical Co., St. Louis, MO. Norit charcoal, methylcellulose, and Wright Giemsa “Accustain” were obtained from Fisher Scientific Co., Fairlawn, NJ. Ficoll–Hypaque was purchased from Pharmacia Fine Chemicals, Piscataway, NJ, whereas 2-mercaptoethanol was obtained from Eastman Kodak Co., Rochester, NY.

Preparation of antifolate receptor antibody. Human PFR was purified to apparent homogeneity based on its migration as a single stained band during sodium dodecyl sulfate–polyacrylamide gel electrophoresis (3). Rabbit antihuman PFR serum was raised by injecting the purified PFR (emulsified with Freund’s adjuvant) into a New Zealand White rabbit, as previously described (3, 7).

Cultures of erythroid progenitor cells in vitro. Human bone marrow cells were aspirated from the posterior iliac crest of hematologically normal volunteers according to guidelines established by the Human Investigations Committee of Indiana University School of Medicine. The marrow aspirates were diluted 1:1 with IMDM containing sodium.
heparin at 20 U/ml. This mixture was layered over an equal volume of Ficoll–Hypaque (specific gravity of 1.077 g/cm³) and density centrifugation performed at 500 g for 25 min at 4°C in a centrifuge (model TJ-6R; Beckman Instruments, Fullerton, CA). The interface mononuclear cell layer was collected, washed with 20 vol of IMDM, and cells were enumerated before use in culture. Assays for the human burst-forming unit–erythroid (BFU-E) were carried out according to the method of Fauser and Messner (8). In brief, 1 × 10⁴ low density bone marrow mononuclear cells were suspended in 35-ml standard tissue culture dishes, containing a 1:ml mixture of IMDM, 1.1% methylocellulose, 30% fetal bovine serum, 5 × 10⁻³ M 2-mercaptoethanol, 1 U of erythropoietin, and 10% of either preimmune serum or varying dilutions of antihuman PFR antisera. All sera were routinely filtered with 0.22 μm Millex-GV filters (Millipore/Continental Water Systems, Bedford, MA) before addition to culture. To test for the specificity of antisera, 800 μl antihuman PFR antisera or preimmune serum was incubated with 2 μg purified human PFR (stock solution of 1 mg/ml protein in 0.1 M potassium phosphate, pH 7.5, containing 1% Triton X-100 and 0.15 M NaCl) for 2 h at 37°C before addition to the erythroid progenitor cell assay system.

The culture plates were incubated at 37°C in a light-protected, humidified atmosphere of 5% CO₂ in air. BFU-E were scored after 14 d using standard criteria (8, 9). To determine the number of cells per erythroid colony, plates containing enumerated numbers of erythroid bursts were harvested into sterile PBS (10 mM potassium phosphate, pH 7.5, containing 150 mM NaCl) and the cells sedimented at 1,000 g for 10 min at 22°C. The resulting supernatant was aspirated, discarded, and the sedimented cells were washed with 50 vol of PBS for three additional centrifuge-wash cycles, as described above. They were then resuspended in 0.5 ml PBS and total erythroid cell counts were determined manually with a hemocytometer. Based on the number of cells derived from a known number of erythroid bursts per plate, the approximate number of erythroid cells per erythroid burst was determined. Erythroid colonies were also directly plated from methylocellulose cultures with a sterile tapered pasteur pipette under direct microscopic visualization, and cells (~10,000) were resuspended in 100 μl PBS and transferred to slides using a centrifuge (Shandon Cytocentrifuge II; Shandon Southern Instruments Inc., Sewickley, PA) at 750 rpm for 5 min. After drying and fixation with methanol, the stained cells were observed for morphological changes. Orthochromat normoblast cell sizing was performed using a digital filar eyepiece (Los Angeles Scientific Instrument Co., Los Angeles, CA) adaptable to a standard light microscope. Also, the number of nuclei per cell was determined in each group of cells described above.

Assay for endogenous intracellular folate. Erythroid bursts were plucked from culture plates containing cells incubated with either rabbit preimmune or antihuman PFR antiserum. The percentage of erythroid cells in these bursts were >98%. Cells from 14 antisera and 18 preimmune sera-treated plates were separately pooled, washed with 100 vol of PBS for two centrifuge-wash cycles as described above, resuspended in 5 ml PBS, and the number of cells/ml determined. The cells were subsequently frozen at −70°C in dry ice–acetone and thawed at 22°C in a water bath for two cycles to lyse the cells. The mixture was centrifuged at 30,000 g for 30 min at 4°C in a centrifuge (model J-21; Beckman Instruments), and the supernatant containing released intracellular material was filtered through 0.22 μm Millex-GV filters to retain particulate material. The filtrate was subsequently assayed for folate content, as previously described (7, 10, 11). The standard curve for the radioisotope dilution assay was based on the ability of known concentrations of unlabeled folate acid (pteroylglutamate [PteGlU]) to competitively inhibit the binding of 125I-labeled folic acid (histamine derivative) to bovine FBP (present as a contaminant in β-lactoglobulin). Briefly, a final reaction volume of 1 ml contained 160 μmol borate-KOH, pH 9.5, 2 μmol diithirotol, 10 μmol 2-mercaptoethanol, 5 fmol 125I-labeled folic acid (histamine derivative), and increasing concentrations of unlabeled PteGlU (0.05–5 pmol) or the sample to be measured for endogenous folate. The mixtures were boiled at 100°C and cooled at 22°C and β-lactoglobulin (4 μg) was added to each tube (Eppendorf Microfuge; Brinkmann Instruments Inc., Westbury, NY). After incubation for 30 min at 22°C, 0.8 mg dextran-coated charcoal was added and the mixtures were incubated another 10 min at 22°C. The samples were subsequently centrifuged in a centrifuge (model 59A Micro-Centrifuge, Fisher Scientific Co.) for 2 min at top speed, and 0.5 ml of each supernatant (containing radiolabeled folate-bound FBP) was counted for radioactivity. The radioactivity of the supernatant in the sample was directly compared with that of the standard curve obtained with PteGlU. Under the conditions of the assay, both PteGlU as well as 5-methyltetrahydrofolate bind bovine FBPs with comparably high affinity (10). This assay however fails to detect other folate forms (such as 5-formyltetrahydrofolate) that have lower affinity and consequently lower ability to displace 125I-labeled folic acid (histamine derivative) from bovine FBP relative to PteGlU. The assay had an upper limit of sensitivity of 100 fmol and a lower limit of sensitivity of 5 pmol PteGlU. Each sample for folate analysis was assayed in triplicate at various dilutions, and the concentration of folate in the unknown sample that was well within the standard curve was determined. Based on the number of cells in each sample (treated with preimmune and immune serum) and the results from the assay for endogenous folate (released from these cells), the concentration of intracellular folate per cell was determined.

Statistical analysis. Each of the experiments described above were carried out on three or more occasions using bone marrow cells from different human subjects. The results of cloning efficiency, and morphology of erythroid progenitor cells in the presence of rabbit preimmune and immune serum was quantitatively and qualitatively comparable with <10% variation. The data for BFU-E–derived colonies is expressed as the mean±SD from quadruplicate values of colony assays, and levels of statistical significance were determined using the student's t test. The values for endogenous folate performed in triplicate in preimmune and immune serum-treated cells are expressed as the mean±SD from cells of one volunteer. In two subsequent experiments with bone marrow cells from different donors, the values for endogenous folate did not significantly differ (<10% variation) from one another in immune and preimmune serum-treated cells.

Results

Rabbit preimmune serum resulted in slight but insignificant augmentation of formation of erythroid bursts in vitro compared with control cells grown in the absence of rabbit serum. Incubation of cells with increasing concentrations of antihuman PFR antisera, however, resulted in a progressive increase in the number of erythroid bursts over concomitant controls assayed in the presence of preimmune serum alone (Fig. 1). With complete replacement of preimmune serum by antihuman PFR antisemur, there was a 3.9-fold greater number of BFU-E–derived colonies per plate as compared with controls containing preimmune serum alone. Also, the erythropoietic bursts developed in the presence of antihuman PFR antisemur were larger and contained ~twofold greater number of cells (6,100±300 cells/BFU-E–derived colony) as compared with controls developed in the presence of preimmune serum (3,450±150 cells/BFU-E–derived colony).

Analysis of Wright–Giemsa stained cells from BFU-E–derived colonies assayed in the presence of antihuman PFR antisemur, but not in preimmune serum, revealed findings that were consistent with morphological changes associated with megaloblastic erythropoiesis (Fig. 2 top, middle, bottom). This was best appreciated in orthochromat normoblasts that contained fine nuclear chromatin that was more characteristic of nuclei from proerythroblasts and early normoblasts (Fig. 2 bottom); also, many such cells contained evidence of dyseryth-
erythropoiesis with two or more nuclei per cell that were joined by fine internuclear bridging material (Fig. 2 middle). Also, there was marked retardation of nuclear maturity and pyknosis in antiserum-treated cells; these morphological findings are consistent with functional nuclear-cytoplasmic asynchrony, which is the hallmark of megaloblastic erythropoiesis (1).

To be certain if the effects of the antihuman PFR antiserum on in vitro erythropoiesis were specifically due to antibody-mediated perturbation of a protein immunologically similar to the PFR, immune serum was preincubated with excess purified PFR before incubation with marrow cells. Under these circumstances, cultures containing PFR mixed with immune serum had no evidence of increased cloning efficiency and were similar to control cells grown in preimmune serum both in the absence or presence of purified PFR (Table I). These results indicated that the unique effect of antihuman PFR antiserum alone was mediated by specific antibodies that bound to folate receptors on marrow cells that were immunologically cross-reactive with human PFR.

To avoid any observer bias in the morphological evaluation of normoblasts in cytopsin preparations of antiserum and preimmune serum-treated cells, we introduced quantitative measurements of dyserythropoiesis. As shown in Table I, antihuman PFR antiserum-treated cells had a significantly ($P < 0.001$) greater (a) cell diameter, (b) percent of cells having more than one nucleus, and (c) number of nuclei per cell when compared with control cultures or rabbit preimmune serum-treated cultures, or antihuman PFR antiserum plus PFR-treated assays. Also, smears from all experimental groups (Table I) were observed and scored by three other certified hematopathologists who were “blinded” to the study. Both they and two of the authors (Antony and Hoffman) could reproducibly distinguish morphological characteristics of megaloblastic erythropoiesis in antiserum-treated vs. preimmune serum-treated cells.

To determine if perturbation of folate receptors on BFU-E by specific antihuman PFR antiserum resulted in intracellular folate deficiency (thereby accounting for megaloblastic changes), the intracellular folate content of erythroid cells composing the resultant bursts was determined. Bursts from plates developed in the presence of preimmune serum and antiserum were harvested, washed, and lysed, and the released intracellular folates were quantitated by a radioisotope dilution assay for endogenous folate. Preimmune serum-treated cells contained 0.104 fmol folate/cell (3 pmol±0.20 SD per 2.8 × 10⁶ cells), whereas anti-human PFR antiserum-treated cells, 0.035 fmol folate/cell (3.8 pmol±0.15 SD per 1.1 × 10⁶ cells); thus, there was 67% less intracellular folate in antiserum-treated erythroid cells as compared with the intracellular folate content in cells incubated with preimmune serum.

Taken together, these studies strongly suggest that specific folate receptors play a major functional role in the accumulation of folate in erythroid progenitor cells in vitro.

**Discussion**

Circulating human erythrocytes are unique among human cells in that they have a single external plasma membrane without any intracellular organelles. The fact that these cells contain a 30-fold higher intracellular folate content as compared with that in serum has remained unexplained. The major form of serum folate is 5-methyltetrahydrofolate (monoglutamate), whereas intracellular folates are predominantly polyglutamated (12). The enzyme responsible for polyglutamation of folate, PGlu synthetase, is however absent in mature cells (13). This suggests that the existing folate in mature cells was taken up as 5-methyltetrahydrofolate and subsequently polyglutamated in (immature) erythroid precursors within the bone marrow. However, the component(s) of a specific folate transport system in erythroid progenitors has not been identified.

During studies with high affinity, specific FBPs isolated from human placental membranes, we raised rabbit antiserum to the purified FBPs and identified the presence of immunologically cross-reactive moieties on mature human erythrocytes (3). Subsequent ligand binding studies with ³H- and ¹⁴C-labeled folates did not detect significant specific radioligand binding to isolated erythrocyte membranes. With the recent synthesis of a stable radioiodinated histamine derivative of folic acid with a specific activity that was 50-fold higher than previously available radiolabeled folates, as well as the development of a sensitive and specific radioimmunoassay for placental folate receptors and related high affinity specific FBPs (6), it was possible to extend earlier studies of FBPs on intact erythrocytes. We found (4, 5) that FBPs from purified membranes of circulating human erythrocytes possessed similar biochemical characteristics with respect to ligand binding, antigenic identities, molecular weights, external orientation, and hydrophobicity as physiological KB cell folate receptors (2, 11). However, the number of functional FBPs that participated in the binding and internalization of folates in erythrocytes was < 1% of the total number of immunoreactive FBPs per cell. Furthermore, folate uptake in circulating erythrocytes was four to six orders of magnitude less than that identified in KB cells. Taken together, these findings suggested that the
Figure 2. Alteration of erythroid burst morphology in the presence of (top) preimmune serum (× 400) and (middle) antihuman PFR antiserum (× 400). (bottom) Higher magnification (× 1,000) of a megaloblastic orthochromatic normoblast from an erythroid burst developed in the presence of antihuman PFR antisera.

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Table I. Effect of Antifolate Receptor Antisera on In Vitro Erythropoiesis

<table>
<thead>
<tr>
<th>Addition to culture</th>
<th>BFU-E-derived colony formation/1 × 10^3 cells plated*</th>
<th>Normoblast cell diameter†</th>
<th>Normoblasts having &gt; 1 nucleus‡</th>
<th>Mean number of nuclei/cell§</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>39.6±19.4</td>
<td>8.4±0.4</td>
<td>5</td>
<td>1.3±0.6</td>
</tr>
<tr>
<td>Normal rabbit serum</td>
<td>43.8±20.5</td>
<td>9.4±0.3</td>
<td>12</td>
<td>1.4±0.8</td>
</tr>
<tr>
<td>Antifolate receptor antisera</td>
<td>101.2±47.5**</td>
<td>13.2±0.4**</td>
<td>47</td>
<td>1.8±0.8**</td>
</tr>
<tr>
<td>Normal rabbit serum plus purified folate receptor</td>
<td>34.6±10.8</td>
<td>8.0±0.2</td>
<td>2</td>
<td>1.3±0.8</td>
</tr>
<tr>
<td>Antifolate receptor antisera plus purified folate receptor</td>
<td>44.0±13.3</td>
<td>9.0±0.3</td>
<td>13</td>
<td>1.2±0.5</td>
</tr>
</tbody>
</table>

* Each value represents the mean±SD of assays pooled from eight individual studies. In individual studies, assays were performed in quadruplicate. † Each point represents the mean±SD of measured cell diameters of 200 orthochromat normoblasts composing erythropoietic bursts in individual treatment groups. ‡ Each point represents the percentage of 200 orthochromat normoblasts composing erythropoietic bursts having > 1 nucleus in each treatment group. § Each point represents the mean±SD number of nuclei per 200 orthochromat normoblasts composing the erythropoietic bursts in each treatment group. ** P < 0.001 compared with all other experimental groups.

existing FBPs in circulating erythrocytes were probably vestigial remnants of a prior functional folate transport system in actively dividing erythroid precursors within the bone marrow. With this background, we attempted this study to determine if specific FBPs were functionally important in the proliferation and differentiation of erythroid progenitor cells.

With the advent of methods to specifically assay for human bone marrow erythroid progenitor cells in vitro, we reasoned that the FBPs on the surface of primitive erythroid lineage cells could be perturbed by antihuman PFR antiserum. This hypothesis was based on our earlier studies (4, 5, 5a), which showed (i) cross-reactive material to PFR on circulating erythrocytes; (ii) immunoprecipitation of radioligand-bound solubilized erythocyte FBP with antihuman PFR antiserum (4); and (iii) the demonstration that antihuman PFR antiserum could inhibit the binding and internalization of radiolabeled folate in circulating erythrocytes and sealed-right-side-out ghosts (5). The results from the present studies (Table I and Fig. 1) suggested that antihuman PFR antiserum enhanced BFU-E cloning efficiency as well as increasing the number of cells comprising individual erythroid bursts. The cells comprising the bursts were also megaloblastic by morphologic criteria (Fig. 2 and Table I). Significantly, these changes were associated with a marked reduction of intracellular folate content in antihuman PFR antiserum-treated cells, where ~ one-third the amount of folate was accumulated intracellularly as compared with control cells. Therefore, it is likely that folate-deficient megaloblastic erythropoiesis was induced in vitro by antihuman PFR antiserum.

It is important to emphasize that the method to quantitate endogenous folate has definite limitations in that it relies solely on the ability of (released) intracellular folates to displace 125I-labeled folic acid (histamine derivative) from bovine FBP. Since various folates differ significantly in their affinity for FBPs (3, 7, 10), it follows that the assay will identify only those folates (like 5-methyltetrahydrofolate) having high affinity for FBPs. Conversely, other folates (e.g., 5-formyltetrahydrofolate) having lower affinity for FBPs will not be measured even if they were present in significantly increased concentrations intracellularly. Moreover, since the forms of folates in erythroid precursor cells during in vitro erythropoiesis is not known, a decrease in measured endogenous intracellular folate (in antiserum-treated cells as compared with preimmune serum-treated cells) may be interpreted as being due to either (i) a true decrease in total cellular folates with comparable high affinity for FBPs like 5-methyltetrahydrofolate, or (ii) a relative decrease in folate forms like 5-methyltetrahydrofolate due to their conversion to folate forms like 5-formyltetrahydrofolate. However, the fact that antiserum-treated cells exhibited megaloblastic erythropoiesis is consistent with the conclusion that there was a true intracellular folate deficiency of 5-methyltetrahydrofolate–like forms after interaction of antihuman PFR antibodies with specific folate receptors on BFU-E and BFU-E-derived erythroid bursts. Since excess of antiserum was present during the development of BFU-E-derived colonies in vitro, we suggest that the antibody–antigen interaction led to a perturbation of the function of folate receptors resulting in inhibition of folate uptake in erythroid progenitor cells.

In contrast to our data, it has recently been shown (14) that growth of erythroid colony formation can be inhibited by assaying human bone marrow cells in the presence of antitransferrin receptor monoclonal antibodies. At present, we have no data to explain why BFU-E cloning efficiency and colony size were increased by the antihuman PFR antiserum. Nevertheless, these results with antihuman PFR antiserum-induced folate deficiency with resultant increased cloning efficiency of erythroid progenitors have correlated with the clinical findings characteristic of human folate deficiency, where hypercellularity with megaloblastic erythropoiesis is commonly observed (1). Little information is available concerning in vitro erythropoiesis in megaloblastic anemia due to cobalamin (vitamin B12) or folate deficiency. However, it has been reported (15) that when bone marrow cells from patients with megaloblastic anemia were assayed for erythroid progenitor cells in vitro in the presence of erythropoietin, there was a fourfold increase in the number of erythroid colonies as compared with normal cells. Also, such colonies appeared earlier and formed even in the absence of exogenously added erythropoietin. Furthermore, although these erythroid colonies had megaloblastic morphologic features after 3 d of incubation, these characteristics were not evident by day 7 suggesting a correction of nutritional deficiency under conditions of in vitro growth. It is therefore of significant interest that we obtained similar results with cells assayed in the presence of antihuman PFR anti-
serum (two- to threefold increase in erythroid bursts, twofold increase in cells per erythroid burst, megaloblastic features) as compared with cells grown in preimmune serum. Therefore, we hypothesize that in addition to mediating folate uptake in erythroid progenitors, interaction of antifolate receptor antibodies with folate receptors may result in augmentation of cell proliferation. The in vitro experimental model of intracellular folate deficiency as induced by perturbation of folate receptors with antihuman PFR antiserum may be used to address fundamental questions pertaining to the metabolism of folate receptors in erythroid progenitor cells at the biochemical and molecular level. Our recent studies with KB cells appear to indicate profound changes in intracellular folate and folate receptor metabolism as a consequence of altering the folate content in the growth media of these cells (16); therefore, it is possible that an analogous process may occur with erythroid progenitor cells in vitro.

One cautionary note should be considered when interpreting this data. We have presumed that the primary effects of the antihuman PFR antiseras were directed against erythroid progenitor cells. Since it is likely that all marrow cells express similar proteins and the marrow cells assayed were relatively heterogeneous, an alternative explanation for our findings might involve the additional perturbation of folate receptors on marrow accessory cell populations that influence erythropoiesis. The effect of the antireceptor antiseras on enriched populations of marrow cells will be pursued to explore these possibilities. Preliminary studies appear to suggest that cells comprising CFU-granulocyte-macrophage-derived colonies assayed in the presence of antihuman PFR antiseras also exhibit features characteristic of megaloblastic hematopoiesis such as neutrophil hypersegmentation.

In summary, the findings that antihuman PFR antiserum mediates the induction of megaloblastic erythropoiesis in vitro provides strong evidences for a functional role of folate receptors in normal erythropoiesis.

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References


