Erythrocyte membrane microviscosity and phospholipid composition in lead workers

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ABSTRACT The microviscosity and fluidity of erythrocyteghost membranes from lead workers and control subjects was measured by fluorescence polarisation using the fluorophore, 1,6-diphenyl-1,3,5-hexatriene (DPH). Increased blood lead was associated with a significant decrease in the average microviscosity of resealed and unsealed erythrocyte membranes. Since DPH fluorescence reflects the organisation of lipids in the central core of the membrane, two aspects of phospholipid metabolism were investigated. Phospholipids were extracted from red blood cell ghost membranes and identified by high performance liquid chromatography. The ratio of phosphatidylcholine to phosphatidyl ethanolamine, an established correlate of membrane fluidity, was significantly increased in lead workers. This is attributed to the known increases in red blood cell cholesterol in lead workers and the structural incompatibility of phosphatidyl ethanolamine and cholesterol, which result in a compensatory increase of phosphatidylcholine. Erythrocyte ghost membranes from control subjects were resealed with the intermediates in phospholipid synthesis that increase with a lead inhibited decrease in red blood cell pyrimidine 5'-nucleotidase. Membrane fluidity was not modified by incubation with cytidine triphosphate, uridine triphosphate, cytidine diphosphate choline, or cytidine diphosphate ethanolamine. Alterations in the microviscosity of the lipid regions of the hydrophobic core of the erythrocyte membrane bilayer and in the phospholipid composition of the membrane may be defects which contribute to the clinical and biochemical instability of the red blood cell on exposure to lead.

Mechanisms that produce shortened red blood cell (RBC) survival, increased shear viscosity, and a decrease of osmotic fragility in erythrocytes from lead workers are incompletely defined. Fukumoto et al found that lead modifies the percentage of globular proteins that affect membrane ion transport or fluidity.¹ Conformational changes in the proteins of the RBC membrane believed responsible for sodium/potassium ATPase function were reported by Selhi et al.² Using erythrocyte membranes and electron spin resonance with a 16-doxyl-stearate label to probe the inner core of the RBC membrane, Valentino et al found a decrease in membrane fluidity associated with exposure to lead and attributed this to changes in membrane protein or to the increased cholesterol.³

The present study of the effects of lead exposure on RBC membrane used 1,6-diphenyl-1,3,5-hexatriene (DPH) fluorescence as a primary index of alterations in membrane lipids of the central core and examined its relation to blood lead and to phospholipid composition of the membranes. We also tested the in vitro effect of loading and sealing erythrocyte ghost membranes with the pyrimidine nucleotides and esters that accumulate in RBC with lead induced inhibition of pyrimidine 5'-nucleotidase and are known intermediates in phospholipid metabolism.

Subjects

All subjects were adult white men aged 24–45 who were employed as lead workers (PbW) or in health care (C). Informed consent was provided under a protocol approved by the University of Nebraska Medical Center review board.

Methods

BLOOD LEADS

Heparinised venous blood was collected in lead free vacutainer tubes (Becton, Dickinson, Rutherford,
Ten microlitre aliquots of whole blood diluted in NH₄H₂PO₄ and Triton X-100 were assayed in duplicate using a Perkin Elmer 2380 atomic absorption spectrophotometer with a graphite furnace.⁴

**MEMBRANE PREPARATION**

*Unsealed ghosts*—All procedures were performed at 0–4°C. Saline (0·9% NaCl solution) washed, packed red cells were lysed with 20 volumes 5 mM sodium phosphate, pH 8. The resulting erythrocyte ghosts were centrifuged at 15 000 rpm for 15–20 minutes and the supernatant removed by aspiration. The membrane pellet was resuspended in phosphate buffer and recentrifuged at 15 000 rpm for 15–20 minutes. This washing procedure was repeated at least three times to produce colourless ghosts.

*Sealed right side out ghosts*—Buffer loaded unsealed ghosts (see A) were resuspended in 40 volumes of phosphate buffered saline (150 mM NaCl, 5 mM sodium phosphate, pH 8) and incubated at 37°C for 40 minutes.⁵ The ghosts were pelleted and washed twice with phosphate buffered saline before fluorescence polarisation measurements.

**NUCLEOTIDE LOADING**

Three hundred microlitres of unsealed erythrocyte membrane ghosts (see A) were suspended in 700 μl 50 mM Tris/glycylglycine buffer, pH 8, containing 5 mM MgCl₂, 1270 μM ATP, and the following nucleotides: CTP 770 μM; UTP 590 μM; CDPA-C 930 μM, and CDP-E 410 μM. All concentrations were comparable to those in congenital deficiency of pyrimidine 5'-nucleotidase.⁶ Reaction mixtures were held for 14–16 hours at 4°C and then incubated one hour in a shaking water bath at 37°C. The resealed ghosts or vesicles were washed with 10 volumes of a solution containing 150 mM NaCl, 10 mM Tris-HCl, pH 8, and 5 mM MgCl₂. Vesicles were pelleted at 15 000 rpm and the supernatant aspirated before microviscosity measurements.⁷

**PROTEIN DETERMINATION**

Suspensions of membrane ghosts were diluted to obtain protein concentrations suitable for assay. One hundred microlitres of diluted membrane suspensions and bovine serum albumin as standards were incubated with 100 μl 0·1 N NaOH for one hour at 37°C. Bio-Rad protein reagent (Bio-Rad, Richmond, CA), 5 ml, was added and absorbance at 595 nm was used to determine protein concentrations.⁸

**FLUORESCENCE POLARISATION**

RBC membrane microviscosity was measured by the method of Shnitzky and Barenholz.⁹ Three microlitres of DPH, in 0·001 M tetrahydrofuran, were mixed with 3 ml degased membrane buffer. Fifteen microlitres of membrane suspension were added, followed by gentle mixing. Fluorescence polarisation measurements were carried out at 25°C using a Perkin Elmer LS-5 fluorescence spectrophotometer. Excitation and emission wavelengths were 330 and 449 nm, respectively. Recordings were made of fluorescence intensities parallel (I₂) and perpendicular (I₁) to the direction of the polarisation of the exciting light. Calculations were made of fluorescence anisotropy, r, where \( r = \frac{I_2}{I_1} \), and the anisotropy parameter, \( \alpha_p = \left( \frac{I_0 - 1}{r} \right)^{-1} \), which provides a similar index of the resistance to rotational motion. The greater the \( \alpha_p \) the higher the microviscosity of the environment.¹⁰ The term \( r_0 \) has an experimental value of 0·362 for DPH.

**PHOSPHOLIPID ANALYSIS**

One hundred microlitres of resealed membranes (buffer loaded) were extracted with 2 ml chloroform/methanol/2M HCl (6:3:1). The chloroform layer was separated and the aqueous layer re-extracted twice with 2 ml chloroform/methanol (2:1). The combined chloroform extracts were dried under nitrogen. Lipid was resuspended in 25–34 μl hexane and injected into a Beckman (Berkeley, CA) chromatographic system consisting of two model 110 A pumps, a model 420 microprocessor (Beckman Instruments, Palo Alto, CA) and Partasil 10 SAX anion exchange columns with a UV detector (Laboratory Data Control, River Beach, FL) operating at 206 nm. Phospholipids were eluted by high performance liquid chromatography using a continuous gradient from 100% solvent A (hexane, propanol, water—6:8:0·75) to 100% solvent B (hexane, propanol, H₂O—6:8:1·5).¹¹ The separation into phosphatidyl ethanolamine (PE), phosphatidyl inositol (PI), phosphatidyl serine (PS), and phosphatidyl choline (PC) took 18 minutes.

**STATISTICAL ANALYSIS**

The SAS program (SAS Institute Inc, Cary, NC) was used to calculate t tests for significant difference, F tests for analysis of variance (ANOVA), and univariate and bivariate regressions for multiple measurements on the same subjects. Data are presented throughout as the mean ± the standard deviation. Blood lead values were logarithmically transformed because they were not normally distributed.

**Results**

RBC membrane viscosity, expressed as the anisotropy parameter (ap) of membrane fragments, was...
Table 1  Anisotropy parameter (ap) of red blood cell membrane fragments

<table>
<thead>
<tr>
<th></th>
<th>Lead workers (n = 6)</th>
<th>Controls (n = 5)</th>
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<tbody>
<tr>
<td>Blood lead (µM)</td>
<td>1.54 ± 0.47</td>
<td>0.39 ± 0.13*</td>
</tr>
<tr>
<td>Anisotropy parameter</td>
<td>1.96 ± 0.54</td>
<td>2.70 ± 0.27*</td>
</tr>
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*p < 0.05 by t test.

Effect of storage of mean ap of RBC fragments of 2 lead workers:
ap (fresh) 1.78
ap (24 h) 2.78
ap for DPH = \[0.362 - \frac{r}{r} \]^{-1}

Table 2  Anisotropy parameter (ap) of resealed, haemoglobin free red blood cell ghosts

<table>
<thead>
<tr>
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<th>Lead workers (n = 25)</th>
<th>Controls (n = 23)</th>
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<tbody>
<tr>
<td>Blood lead (µM)</td>
<td>1.38 ± 0.33</td>
<td>0.29 ± 0.06*</td>
</tr>
<tr>
<td>Anisotropy parameter</td>
<td>1.54 ± 0.15</td>
<td>2.03 ± 0.56**</td>
</tr>
<tr>
<td>Protein content (mg/ml)</td>
<td>0.97 ± 0.42</td>
<td>0.94 ± 0.35</td>
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*p < 0.01 by t test; **p < 0.0001 by F Ratio, ANOVA.

The multiple regression for the bivariate correlation of ap with PbB and the protein concentration of each sample was:
ap = 2.6237 - 0.6234 log10 PbB µg/dl - 0.3600 mg/ml protein
r2 log PbB: 0.29, SEE 0.4753, r2 protein concentration: 0.09, SEE 0.3021; r2 total: 0.37; adjusted r2 total: 0.34.

Discussion

Changes in the rheological properties of the RBC after lead exposure such as increased blood viscosity and decreased deformability are well documented but not well understood.1–3 Studies have shown that alterations in the protein cytoskeleton or lipid composition of erythrocyte membranes can affect membrane fluidity or microviscosity and ultimately the macroviscosity or flow properties of the RBC.12

In this study membrane fluidity was examined through the technique of fluorescence polarisation. Fluorophores such as DPH, which are immobilised in the hydrophobic core of a membrane and excited by polarised light, will emit fluorescence that may be interpreted through the ap \[\frac{r_0 - r}{r} \]^{-1}, (see fluorescence polarisation under methods). The greater the value of this parameter, the higher the apparent “microviscosity” of the environment. The present studies were undertaken to characterise the

Table 3  Phospholipid ratios of red blood cell ghosts

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<tr>
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<th>Lead workers (n = 12)</th>
<th>Controls (n = 10)</th>
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<tbody>
<tr>
<td>Blood lead (µM)</td>
<td>1.17 ± 0.27</td>
<td>0.32 ± 0.12*</td>
</tr>
<tr>
<td>Phosphatidyl choline:</td>
<td>2.00 ± 0.69</td>
<td>1.10 ± 0.21**</td>
</tr>
<tr>
<td>Phosphatidyl ethanolamine:</td>
<td>0.74 ± 0.29</td>
<td>0.78 ± 0.58</td>
</tr>
<tr>
<td>Phosphatidyl ethanolamine:</td>
<td>0.96 ± 0.37</td>
<td>1.10 ± 0.22</td>
</tr>
<tr>
<td>Phosphatidyl inositol:</td>
<td>1.7 ± 0.87</td>
<td>1.7 ± 0.57</td>
</tr>
<tr>
<td>Fluidity index</td>
<td>2.6 ± 0.71</td>
<td>3.6 ± 1.1</td>
</tr>
</tbody>
</table>

*p < 0.01 by t test; **p < 0.0001 by F ratio, ANOVA.

Fluidity index = ap/mg protein.
microviscosity of the non-polar regions of the erythrocyte ghost membranes. The demonstrated decrease in the microviscosity is interpreted as a perturbation in the lipid organisation of the central core of the bilayer.

Membrane fluidity as determined by fluorescence polarisation is distinct from assessment by electron spin resonance (ESR), nuclear magnetic resonance, and other techniques, which better define the interactions of lipids with aqueous components, other lipids and proteins, the rotational and lateral mobility of proteins, and the effects of divalent cations. Valentino et al found a modest decrease in the fluidity of RBC from PbW when measured by ESR with 5 doxyl-stearate (5NS) and 16 doxyl-stearate (16NS). There was a greater increase in viscosity of the inner core (16NS) than the outer layer but no significant direct correlation with PbB which ranged from 2.5 to 3.9 μM, considerably higher than that of our subjects. Fiorini et al found an effect on the ESR of 16NS in liposomes incubated with 2.85–11.5 μM Pb⁺⁺. The effect of high concentrations of in vitro Pb⁺⁺ on ESR viscosity is attributed to an increase in charge density of the central core and intramembranous aggregation of membrane bound proteins.

Decreases in microviscosity as measured by the fluorescence polarisation of DPH are primarily related to cholesterol depletion, to the length and unsaturation of the fatty acid ester chains in phospholipids, and to increases in the relative amount of phosphatidylcholine expressed either as the PC:PS ratio or the PC:PE ratio.

Given the characteristics of DPH fluorescence as an index of fluidity, we propose that the increase in PC and the direct correlation of a decrease in DPH viscosity with PbB relate to the well established increase of RBC cholesterol in PbW. Both PE and cholesterol have truncated cone shapes that are structurally incompatible. This results in a preferential association with PC in PC-PE cholesterol mixtures and is why high amounts of PE and cholesterol do not occur in the same membrane. A similar association of high PC and cholesterol is found in the RBC membranes of lecithin:cholesterol acyl transferase deficiency and in hereditary haemolytic anaemia with high red cell membrane phosphatidyl choline.

Decreased microviscosity as measured by DPH does not contradict the increased macroviscosity of the RBC in PbW. It is an index of alterations in the rotational mobility of lipids in the central hydrophobic core of the bilayer that may well contribute to the haemolytic effects of lead exposure.

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References

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