Red blood cell microrheological changes and drug transport efficiency

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Abstract. The purpose of this study was to estimate drug effect on red blood cell (RBC) microrheological properties making for blood transport efficiency. It is well known that blood flow in microcirculation, tissue perfusion and oxygenation depend on red blood cell (RBC) microrheological properties, namely deformability (RBCD) and aggregability (RBCA). In turn the changed red cell mechanical properties can influence the tissue oxygenation as well as the drug delivery to tissues. The exposure of RBCs to agonist of β-adrenergic receptors (isoproterenol) caused RBCA decrease and an increase of their deformability. The prostaglandin E1, prostacyclin and insulin showed the similar positive microrheological effect. Neutral microrheological effect was fixed after cell incubation with 5-fluorouracil, rolipram and phorbol 12-myristate 13-acetate; no significant alterations of microrheological properties were found. The agonists of α-adrenergic receptors and prostaglandin F2α showed the negative microrheological effect: RBCD was markedly worsened and RBCA was enhanced by 49–60%. It was found that three PDE inhibitors: vinpocetine, rolipram and cilostasol – significantly decreased RBCA and increased RBCD. Therefore PDEs might be considered as red blood cell molecular targets for some drugs. Stimulated by A23187 Ca2+ influx was accompanied by an increase of RBCA and a decrease of RBCD. The blocking of Ca2+ entry into the RBCs by verapamil led to a significant positive RBC changes. Therefore the red cell membrane Ca2+ channels can be considered as molecular targets responsible for red cell microrheology modifications too. Taken together the obtained data showed that the drugs infused in blood flow can interact with blood cells, including the most numerous cell population – red blood cells. This interaction can lead to an alteration of RBC microrheological properties and blood transport possibilities at least according to three abovementioned scenarios.

Keywords: Red blood cell aggregation and deformability, drugs, microrheological effect, cellular signaling pathways

1. Introduction

Blood flow in microcirculation, tissue perfusion and oxygenation depend on red blood cell (RBC) microrheological properties; namely deformability and aggregation [8, 19]. Drugs and signaling molecules can affect microrheological properties of blood cells including red cell aggregability and deformability [4, 6]. It is known that at the capillary level the blood flow efficiency depends on red cell microrheological properties [21], and their alterations can influence not only on the tissue oxygenation [11] but also affect the drug delivery to tissue. Three variations of drug effect on red cell microrheology can be distinguished by the probable red cell microrheological changes after drug treatment: 1) drug has no effect on the red cell aggregability and deformability and blood transport capacity doesn’t change – it is neutral effect; 2) drug decreases red cell deformability and increases red cell aggregation – it is negative effect; and 3) drug increases red cell deformability and decreases red cell aggregation (RBCA) – it is positive effect. One of the well-known rheologically “positive” drugs is pentoxifylline [8]. The

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main mechanism of pentoxifylline blood rheological efficiency is connected with its phosphodiesterase (PDE) inhibitory activity [7]. The other drugs, having phosphodiesterase as intracellular targets, can be microrheologically active both in vitro and in vivo [16]. It is important to note that the red blood cells were historically considered inert to the regulatory signals from other cells. However they are surprisingly well equipped with the machinery required for intercellular communication [13, 15, 20]. It has been proven experimentally that the RBC membrane contains α- and β-adrenoceptors [20, 23], insulin receptors [3], and cholinergic muscarinic receptors [26]. Note that the marked changes of red blood cell microrheological properties such as red cell deformability and aggregation are mediated by an activation of the molecular control mechanisms [12, 13, 15, 20].

The present study was designed to explore the effects of some drugs and chemicals as the signaling molecules on red blood cell microrheological properties.

2. Materials and methods

2.1. Preparation of blood samples

Venous blood samples (15 ml) were drawn via sterile venipuncture using heparin (5 IU/ml) as anticoagulant. The study was approved by the local ethic committee at the Yaroslavl State Pedagogical University, and an informed consent of all the subjects were obtained according to the recommendations of the Declaration of Helsinki (The International Response to Helsinki VI, The WMA's Declaration of Helsinki on Ethical Principles for Medical Research Involving Human Subjects, as adopted by the 52nd WMA General Assembly, Edinburgh, October, 2000).

Red blood cells (RBCs) were separated by centrifugation at 1,400 g for 20 min and washed 3 times with 10 mM phosphate buffered saline (PBS) (pH = 7.4). The washed RBCs were then resuspended in PBS at a hematocrit (Hct) of approximately 40% for incubation with drugs or chemicals (experimental panel) and without any drugs (control panel). After an incubation period the supernatant was removed by centrifugation and cells were resuspended in autologous plasma at Hct = 40% for measurement of RBC aggregation and deformability.

2.2. Protocols for in vitro aggregation studies

It was organized fourth research sessions. In the first research session RBC suspension was divided into four aliquots and exposed to: 1) Isoproterenol (1.0 μM); 2) Insulin 0.1 μM; 3) Prostaglandins (PGE1, 0.01 μM); 4) Prostacyclin (PGI2, 0.01 μM).

In the second research session RBC suspension aliquots exposed to: 1) 5-fluorouracil (5-FU, 5 μM); 2) Inhibitor of the protein kinase C (PKC) activity, phorbol 12-myristate 13-acetate (PMA, 3 μM; n = 24); 3) PDE4 inhibitor rolipram (10 μM; n = 24).

In the third research session RBC suspension aliquots exposed to: 1) prostaglandin F2α (PGF2α, 0.01 μM); 2) phenylephrine – alpha-1-adrenoceptor agonist (1.0 μM); 3) clonidine alpha-2-adrenoceptor agonist (1.0 μM).

In the fourth research session RBC suspension aliquots exposed to: 1) calcium ionophore A23187 (3 μM); 2) verapamil – Ca2+ channel blocker (10 μM). In this research session the Ringer’s solution was used as suspension medium, containing Ca2++.
To study the role of phosphodiesterases (PDEs) in red cell microrheology alterations the cells were incubated with: PDE1 inhibitor vinpocetine (10 μM; PDE3 inhibitor cilostazol (10 μM) and PDE4 inhibitor rolipram (10 μM). The cell shape was controlled with the light microscopy. The cell incubation was performed at 37°C for 15 min. The red blood cells, incubated in the phosphate buffered saline (or in Ringer’s solution) without any drugs, were used as a control. Stock solutions of drugs were prepared in DMSO or water. All analyses were completed within 4 h after the blood collection. Drugs and chemical compounds were mainly purchased from Sigma.

2.3. Red blood cell aggregation measurement

Red blood cell aggregation (RBCA) in native plasma was assessed by the Myrenne Aggregometer which provides an index of RBC aggregation facilitated by low shear. In brief, the suspension was subjected to a short period of high shear to disrupt pre-existing aggregates, following which the shear was abruptly reduced to 3 s⁻¹ and light transmission through the suspension was integrated for 5 seconds; the resulting index, termed “M5” by the manufacturer and “RBCA” herein, increased with enhanced RBC aggregation.

To estimate the deformability of RBCs the latter were placed into a flow microchamber. The cells were attached to the bottom part of the chamber with “one point” and then were deformed by shear flow under constant shear stress (τ). At a given volume flow rate Q (determined by weighing the amount of saline which flows through the flow channel in a given time period) the shear stress at the surfaces is given by [2]:

\[ \tau = \frac{6 \eta Q}{wh^2}, \]

where \( \eta \) is the viscosity of the perfusate, \( w \) is the width and \( h \) is the height of the flow passageway. In our experiments, \( \eta = 1.07 \text{ mPa}\cdot\text{s}, w = 0.90 \text{ cm}, h = 0.01 \text{ cm}. \)

2.4. Miscellaneous techniques

The whole blood and red cell suspension hematocrite was determined via the microhematocrit method (i.e., 12,000 × g for 7 minutes).

2.5. Statistics and data presentation

The results are presented as mean SEM. The differences between the mean values were evaluated using an ANOVA test. The values of \( p < 0.05 \) indicate statistical significance.

3. Results

3.1. The relatively positive microrheological effect of the drugs

Catecholamines as stress hormones ensure an effective adaptation to various factors of the environmental. It is important to note that human erythrocyte membranes contain both \( \alpha_1 \) - and \( \beta_2 \) - adrenergic receptors. In the presence of 1.0 μM Isoproterenol the red blood cell aggregation was decreased to some extent versus control but not statistically (Table 1), while RBCD was significantly increased by 26% (\( p < 0.05 \)) under these conditions. It is well known that insulin is a functional antagonist of the catecholamines, and erythrocytes possess insulin receptors. Therefore it may be assume that the red cell microrheology will be altered after the cell incubation with insulin.
The RBC micro rheology alterations after cell incubation with some drugs (Neutral hemorheological effect) (M ± m; n = 24)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control</th>
<th>5-FU (5 μM)</th>
<th>Rolipram (10 μM)</th>
<th>PMA (3 μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBCA</td>
<td>7.14 ± 0.15</td>
<td>7.24 ± 0.18</td>
<td>6.68 ± 0.36</td>
<td>7.29 ± 0.26</td>
</tr>
<tr>
<td>RBCD</td>
<td>0.208 ± 0.006</td>
<td>0.202 ± 0.007</td>
<td>0.220 ± 0.007</td>
<td>0.215 ± 0.005</td>
</tr>
</tbody>
</table>

Abbreviations: 5-FU – 5-fluorouracil; PMA - phorbol 12-myristate 13-acetate.

We have tested this hypothesis and found the RBCA was decreased by 44% ($p < 0.05$) and RBCD was increased by 16% ($p < 0.05$) versus control. The similar positive effect was observed after RBC incubation with Prostaglandin E1 and Prostacyclin (Table 1).

3.2. The relatively neutral micro rheological effect of the drugs

5-fluorouracil is a drug for cancer chemotherapy and it employs for this purpose. It was found that this chemical has a neutral micro rheological effect. Neither RBCA nor RBCD were changed significantly after the cell incubation with this drug (Table 2). PDE4 inhibitor rolipram also possess the relatively neutral effect on RBC micro rheological behavior. Changes of red cell micro rheology (up to 6%) were not significant (Table 2). PMA is an activator of cellular protein kinase, mainly PKC. It was shown minor alteration (by 2-3%) of red cell micro rheological properties, both RBCA and RBCD after PMA treatment (Table 2).

3.3. The relatively negative micro rheological effect of the drugs

The prostaglandin of F family – PGF2α intensified the red cell aggregation. In presence of PGF2α erythrocyte aggregation increased by 66%, $p < 0.05$ (Table 3). PGF2α is known as a calcium cell entry stimulator [20]. Therefore the aggregation rise may be associated with an augmentation of Ca²⁺ influx.

In the presence of 1.0 μM α₁- and α2-receptor agonists (phenylephrine and clonidine) the red blood cell aggregation was markedly increased – from 49 to 60% (vs. control; $P < 0.05$). The most significant aggregation effect had α2-agonist clonidine. After the cells incubation with this drug the RBCA was increased by 60% (Table 3). Therefore, it concerns mostly α2-adrenergic receptor stimulation by epinephrine or specific α2-agonists. On the contrary, these alpha-receptor agonists had the lowering RBCD effect (Table 3). On the whole, the obtained data make us believe that an activation of the red blood cell alpha-receptor is associated with a relative negative micro rheological effect.
Table 3
The RBCA and RBCD alterations after cell incubation with some drugs (Negative hemorheological effect) (M ± m; n = 24)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control</th>
<th>PGF2α (0.01 μM)</th>
<th>Phenylephrine (1.0 μM)</th>
<th>Clonidine (1.0 μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBCA</td>
<td>7.34 ± 0.26</td>
<td>12.18 ± 0.433 *</td>
<td>10.96 ± 0.324 *</td>
<td>11.74 ± 0.344 *</td>
</tr>
<tr>
<td>RBCD</td>
<td>0.206 ± 0.005</td>
<td>0.189 ± 0.0055 *</td>
<td>0.192 ± 0.0044 *</td>
<td>0.196 ± 0.0064</td>
</tr>
</tbody>
</table>

*p < 0.05 versus control. Abbreviations: RBCA – red blood cell aggregation; RBCD – red blood cell deformability; PGF2α – Prostaglandin F2α.

Table 4
The red blood cell microrheology changes after cell incubation with Ca2+ cell entry blocker Verapamil and calcium ionophore A23187 (M ± m, n = 24)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control</th>
<th>Verapamil (10 μM)</th>
<th>A23187 (3 μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBCA</td>
<td>7.18 ± 0.28</td>
<td>5.88 ± 0.255 *</td>
<td>12.78 ± 0.544 *</td>
</tr>
<tr>
<td>RBCD</td>
<td>0.212 ± 0.004</td>
<td>0.250 ± 0.0055 *</td>
<td>0.193 ± 0.0044</td>
</tr>
</tbody>
</table>

*p < 0.05 versus control. For abbreviations, see Table 1.

4. Conclusion: some probable molecular targets in RBCs for drug microrheological effect

4.1. Role of Ca2+ in red blood cell microrheology alteration

The measurement of the total red cell calcium concentration has yielded values between 5 and 50 nM [22], while only a few percent of total red cell Ca2+ is in an ionized form [27]. Any rise in intracellular Ca2+ in erythrocytes activates a specific K+ channel (‘Gardos’ channel) which normally makes little contribution to K+ fluxes [10]. RBC treatment with stimulators of Ca2+ influx as well as the inhibition of Ca2+ efflux resulted in a significant increase of their aggregation [17]. Contrasting with it, Ca2+ entry blocking into the red cells by verapamil led to a significant RBCA decrease and RBCD rise (Table 4). Our experimental data indicated that the stimulation of Ca2+ influx by A23187 caused a significant increase of red blood cell aggregation and some RBCD decrease (Table 4).

The changes in RBC Ca2+ and cAMP content are not simply two isolated linear signaling pathways, but they actually interact at multiple levels to form an effective signaling network.

4.2. Effect of phosphodiesterase (PDE) activity inhibition on red blood cell microrheology

It is known that the intracellular cAMP level is regulated by phosphodiesterases (PDEs) [1, 5]. To study their role in red cell aggregation alterations the cells were incubated with: PDE1 inhibitor vinpocetine; PDE3 inhibitor cilostazol; PDE4 inhibitor rolipram. The red cell aggregation was reduced after the cell incubation with the above-mentioned drugs, having the PDE inhibitory activity. The most significant effect was found under cell incubation with PDE1 inhibitor – vinpocetine. Taken as a whole RBCA reduction was on the averaged 25% under these conditions (Fig. 1). On the whole, the obtained data make us believe that PDEs are a part of the cell regulatory system “the cell membrane receptor – G-protein – AC-cAMP complex” and they involved in the RBC microrheological control mechanisms.
Although the concept that the cAMP and Ca$^{2+}$ signaling pathways interact at multiple levels was appreciated for some time, an emerging body of data has begun to define specific molecular loci where the two pathways interface. Spotlighting these important molecular loci at which cAMP and Ca$^{2+}$ signaling pathways converge in non-excitable cells can help us further understand and appreciate the complexity and specificity of stimulus–response coupling in these cells. In the capacity of the potential sites of the cross-talk between Ca$^{2+}$ and cAMP signaling systems in red blood cells may be PDEs [14]. For example, the constitutive type 4 phosphodiesterase activity rapidly hydrolyzes cAMP so the Ca$^{2+}$ inhibition of AC is difficult to resolve, indicating that high phosphodiesterase activity works coordinately with AC to regulate membrane-delimited cAMP concentrations, which is important for control of cell-cell apposition [5]. Moreover there are data that clearly indicate that the activation of AC and cAMP increase lead to the Ca$^{2+}$ influx decrease and red cell aggregation and leukocyte adhesion lowering [17, 28]. The similar results were found under the blocking Ca$^{2+}$ entry into cell or PDE activity inhibition [16].

Taken together the obtained data showed that the drugs infused in blood flow can interact with blood cells, including the most numerous cell population – red blood cells. This interaction can lead to an alteration of RBC micro rheological properties and blood transport possibilities at least according to three abovementioned scenarios.

Acknowledgments

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References


