ORIGINAL RESEARCH

Oxidative alterations during human platelet storage

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SUMMARY: During storage of platelet obtained by apheresis several changes occur. The aim of this study was to investigate the effect of storage on activation, apoptosis, protein pattern, lipid peroxidation, and the levels of nitric oxide (NO) and glutathione (GSH) of platelets. In this study, platelets obtained from healthy donors (n=7) by apheresis were kept in an agitator for nine days at 20-24°C. The samples were taken on the 1st, 3rd, 5th and 9th days and platelets were precipitated. Platelet activation with PAC-1 and CD62-P antibodies and platelet apoptosis were measured with Annexin-V using flow cytometer. Platelets were frozen and thawed four times and then NO, GSH and malondialdehyde (MDA) levels were assayed by spectrophotometry. Platelets protein pattern was investigated via sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) procedure. When compared to the 1st day, platelet CD62-P, PAC-1 expressions and Annexin-V levels significantly increased on the 3rd, 5th and 9th days, while platelet NO and GSH levels significantly decreased on the 3rd, 5th and 9th days. Furthermore MDA levels significantly increased on only the 5th and 9th days. Mild changes occurred in the density of platelet protein bands. In conclusion, our results show that alterations of platelet activity during storage period may enhance platelet procoagulant activity which increases trombogenic risk. Therefore, for transfusion using fresh platelets or adjusting platelet preservation are strongly important for platelet behavior in vivo conditions.

KEY WORDS: Transfusion, platelet, activation, apoptosis, protein

INTRODUCTION
Platelets play an important role in maintaining hemostasis. Platelet transfusions are essential for the treatment of patients with thrombocytopenia and thus routinely used during surgery, chemotherapy and for bleeding disorders. Different separation methods are described to prepare therapeutic platelet concentrates from human donor blood (1,2). In platelets obtained with these methods including apheresis, several changes may occur in morphology, adhesion and aggregation, membrane features (protein pattern), activation and apoptosis markers during prolonged storage (1). There are several receptors on platelet membrane which mediate adhesion, aggregation (Glycoprotein (Gp) IIb/IIIa/ fibrinogen receptor), activation (P-selectin) and other cellular events. Activation signals lead to variations in platelet membrane Gp receptor expressions and this situation may affect platelet functions (3). In recent years, by using fluorescently labelled antibodies receptor changes are easily detected on platelets. It has been shown that platelet membrane Gp expressions and platelet GSH levels change during the preparation of platelet concentrates and storage (4,5).

On the other hand, nitric oxide synthase (NOS) has been identified in platelets and it is important in the regulation of platelet recruitment (6). Moreover, the decrease in platelet NO increases platelet activation and cellular production of O2 radicals thus lipid peroxidation. Lipid peroxidation of membranes induced by reactive oxygen species alters the structure and function of membrane components followed by the platelet signaling pathway activation. Additionally, although platelets are anucleated cells, apoptosis like events take place both in vivo and in vitro and they can also synthesize proteins by using several transport mechanisms and the amino acids contents in their cytoplasms (7,8,9). PS (phosphatidylserine) exposure is recognized as a marker of cell death as well as being an activation marker.
Chronically elevated or prolonged exposure of PS on the cell surface increases vascular damage and results in the formation of a hypercoagulable environment in platelets (10,11,12).

The aim of this study was to investigate the effect of storage on activation, apoptosis, protein pattern, lipid peroxidation, NO and GSH levels of platelets obtained from aphaeresis; and in case there were significant alterations, the aim was to interpret how they would influence platelet functions in vivo.

**MATERIAL AND METHODS**

**Materials**

SDS-6H Protein Standard, bisacrylamide, bovine serum albumin (BSA), bromophenol blue coomassie brilliant blue R 250, ethylene diamine tetra-acetic acid (EDTA), HEPES, glycerin, Phosphate-buffered saline (PBS), sodium citrate, reduced glutathione (GSH), Triton X-100, 5,5-dithiobis-2-nitrobenzoic acid (DTNB), adenosine diphosphate (ADP), paraformaldehyde (PFA) were from Sigma (St.Lous,MO,USA); methanol, ammonium persulfate, sodium carbonate, TRIS, sodium potassium tartrate, copper sulphate, thiobarbituric acid (TBA), were from Merck (Darmstadt,Germany); Annexin-V-FITC (Annexin-V-fluorescein isothiocyanate) Apoptosis Detection Kit, binding buffer and FITC anti-human CD62-P were from Becton Dickinson Pharmingen (San Diego, CA,USA); PAC-1 FITC was from Becton Dickinson Biosciences (San Jose, USA).

**Subject Criteria**

The participants of this study are healthy voluntary blood donors who regularly donate blood at the Blood Center of Istanbul University Cerrahpasa Medical School and have given consent. The subjects’ ages were between 20 and 40. The routine anamnesis and physical examination of the subjects were performed. Serological scanning tests were applied. Subjects didn’t have any medication 10 days prior to the aphaeresis.

**Obtaining Platelet Suspension with Apheresis and Preparation of Platelet Samples**

In this study, the platelets obtained from healthy donors (n=7) by the aphaeresis method were kept in an agitator under in-vitr0 conditions for nine days at 20-24°C. The samples were taken on the 1st, 3th, 5th and 9th days. After the centrifugation (9000 rpm, 4°C, 15 min, Hettich, Universal 32R, DLB Labcare, Newport Pagnell, England) process, platelets were isolated and the platelet pellet was washed with Tris-NaCl buffer (0.03 M Tris, 0.12 M NaCl, pH 7.4) containing 5 mM EDTA. The platelets were frozen and thawed 4 times. After centrifugation, the supernatant was obtained. Then the protein concentration was determined with the Lowry method (13) and SDS-PAGE was applied to examine the protein alterations.

**Measurement of GSH**

GSH levels were assayed according to the method of Mergel et al. (14) using DTNB. GSH contents of platelets were determined with GSH (2–30 μg/ml) as the standard. The results were expressed as μg per 10⁹ platelets.

**Measurement of Lipid Peroxidation**

For the measurement of lipid peroxidation, the precipitate was solubilized for 5 h with Tris-NaCl buffer containing 1% Triton X-100 at 4°C and then centrifuged. Lipid peroxidation was evaluated with TBA according to the method of Buege and Aust by spectrophotometry (15). The results were expressed as nmol/10 mg protein.

**Nitrite Assay**

The washed platelets were incubated with 1.44 mmol/L NADPH for 1 h at 37°C. Then, each sample was incubated for 1 h at 37°C after the addition of 20 mU nitrate reductase, which reduced nitrate to nitrite. The platelets were frozen and thawed 4 times. After centrifugation, the supernatant was allowed to react with Griess reagent to form a chromophore; its absorption was measured subsequently at 546 nm. Sodium nitrite (0.2 to 4 μM) was used as the standard (16).
Flow cytometric Analysis of Platelet Activation and Apoptosis

Briefly, washed platelets were diluted in phosphate-buffered saline (PBS) (8 mM NaH2PO4, 5 mM KCl, 125 mM NaCl, 5 mM glucose and 0.5 g/L albumin) and mixed gently. Platelets were identified by staining with FITC conjugated CD41a and by gating from logarithmic scaled forward scatter/side scatter scattergram. FITC-labeled CD62-P (P-Selectin) and PAC-1 (activated GpIIb/IIIa) antibodies were added to each platelet suspension. The tubes were incubated at room temperature, in the dark for 15 minutes for platelet activation analysis. For fixation, PFA (%1 vol/vol) was added and then the samples were diluted with PBS. All samples were stored at 4°C until analysis and analyzed in an hour.

For the measurement of PS exposure, annexin-V-FITC was used. Annexin-V-FITC (25 μg/ml) was added to the diluted samples. Sample tubes were kept on ice until analysis for 10 min in the dark after the addition of an equal volume of cold binding buffer (10mM HEPES/NaOH (pH7.4) 140 mM NaCl, 2.5 mM CaCl2) (17).

The analysis of all the samples was carried out in the FACS Calibur flow cytometry system (Becton-Dickinson, Franklin Lakes, NJ). The system was equipped with 488 nm argon ion laser. Calibrite beads (BD Biosciences, San Jose, CA) were used for daily quality control. 50,000 cells were counted in each tube. The results were expressed as the percentage of positive cells. The negative control cursor was set to 2 % of cells on histograms.

Statistical Analysis

The results were presented as mean ± SD. For the daily platelet alterations, analyses were compared using the Wilcoxon signed-rank test. Statistical analyses were performed with Graph Pad Prism 4 software. P values <0.05 were considered significant.

RESULTS AND DISCUSSION

Platelet transfusions are routinely used during surgery and during several conditions resulting in thrombocytopenia such as chemotheraphy (1, 5). It has been pointed out in several studies that platelets that are kept under the blood bank conditions begin to lose their functions (18,19). Platelets undergo several modifications during storage that reduce their post transfusion functionality. This loss can be observed in the plasma concentrations even in the first 24 hours. There has been debates whether this loss of functions is caused by the platelet activation during the preparation and storage process or by the changes in the pH and enzyme activation of the plasma envi-
environment. Perhaps both mechanisms are responsible for this phenomenon.

In this study, we first detected percentage of protein levels of SDS-PAGE bands determined from densitometry in platelets (figure 1). In several studies carried out so far, approximately 2,300 proteins have been detected in platelets. Some abnormalities can be seen in platelet functions resulting from several protein deficiencies and defects. Since signalling proteins trigger major processes in platelets, especially loss of signalling proteins is very important for their survival. George NJ et al. (20) demonstrated major glycoprotein loss (later this protein is termed GPIb) after electrophoresis in storage platelets. In the present study, the alterations on the membrane protein bands between the first and ninth days were investigated and mild changes were observed in the density of platelet protein bands. However, we should point out that we examined only the platelet membrane proteins in this study; we didn’t determine the changes in the cytoplasmic proteins. Further studies are needed in order to show all the changes in all the protein content.

It is known that activation is the most abundant process in platelets and essential for platelet aggregation and coagulation. On the other hand, platelets undergo apoptosis in various conditions and contain at least some of the machinery necessary for apoptosis, such as caspases and death receptors. PS becomes exposed on the outer cell membranes during the early stages of apoptosis. At the same time, PS exposure is reported to be a predictor of platelet activation as well as being an apoptotic marker (21). For measurement of platelet activation and apoptosis, several monoclonal antibodies have been described that recognize antigens on platelets. PAC-1 has been used identifying fibrinogen receptor GpIIb/IIIa in activated platelets (22). Additionally, P-selectin is only expressed on the platelet surface after α-granule secretion thus after activation (23). Recent studies proved that the increased expression of the P-selectin correlates with the in vivo recovery of transfused platelets (24). PS exposure on the surface of cells is commonly measured by flow cytometry of fluorescently labeled Annexin-V binding to this procoagulant phospholipid (25). In several studies, it was shown that platelet membrane markers including CD62-P, CD63 and Annexin-V increased during prolonged storage (26). Platelets exhibit changed membrane features, increased expression of pro-apoptotic markers and they lose viability during in vitro storage. In fact, the life span of platelets in the human circulation is estimated to be 10 to 12 days. However, after 5 to 6 days of in vitro storage, platelets lose their viability and activity. Therefore, we detected by flow cytometry binding of Annexin-V for platelet apoptosis (at the same time a platelet activation marker) (figure 3) and the expression of PAC-1 and CD62-P for platelet activation (figure 4) in platelets prepared by aphaeresis. In this study, when compared to the 1st day, platelet CD62-P, PAC-1 expressions (the percentage of CD-62 and PAC-1 positive cells) and Annexin-V levels significantly increased (p<0.01) on the 3rd, 5th and 9th days. Our results are consistent with other studies. The activation criteria that we found show that as the storage time of platelets is prolonged, activation potentiality will increase, there will be more sensitivity to agonists following transfusion, and thus they will be inclined to aggregation. Additionally, in prolonged storage increased PS exposure increases procoagulant activity and may cause a prothrombotic condition thus thromboembolic risk.

In this study, we also determined oxidative stress parameters and platelet NO levels during platelet storage. There is strong evidence that oxidative stress is a mediator of apoptosis. Additionally, in this study, when compared to the 1st day, platelet NO and GSH levels significantly decreased on the 3rd, 5th and 9th days; whereas, MDA levels significantly increased on only the 5th and 9th days (p<0.05). It was shown that the decrease in platelet NO increases platelet activation and lipid peroxidation (27). Low nitrite levels observed in the study may indicate that some factors are responsible for the increase in platelet activation, platelet lipid peroxidation and apoptosis. It is still a complicated phenomenon whether resulting oxidative changes trigger activation and apoptosis or activation and apoptosis trigger oxidative changes depending on storage.
In conclusion, our results show that the increase of apoptosis, activation and lipid peroxidation and decrease of NO and GSH in platelets prepared for transfusion will enhance platelet procoagulant activity thus thrombogenic risk in in vivo circulation. On the other hand, storage leads to defects in platelet activation properties, thus their clearance by macrophages. The use of updated storage methods will help to generate platelets for transfusion with optimal haemostatic function and a long circulation time after transfusion. At the same time using fresh platelets and adjusting platelet preservation are strongly important for platelet functions in in vivo conditions.

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REFERENCES