Hemorheological consequences of hind limb ischemia-reperfusion differs in normal and gonadectomized male and female rats

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Abstract. It is known that hemorheological parameters show gender differences that might be altered by gonadectomy (GoE). Since micro-rheological parameters (erythrocyte deformability and aggregation) sensitively change during and after ischemia-reperfusion (I/R), the question arises whether the hemorheological effects of I/R may show gender differences and further changes might be expected when GoE and I/R are additive. Sprague-Dawley rats were divided into six groups: Control males and females, I/R males and females with 1-hour hind limb ischemia, GoE + I/R males and females when 3 months after bilateral gonadectomy the I/R was induced. Before and just after ischemia, and on the 1st-3rd-5th-7th postoperative days blood samples were taken (lateral tail vein, 0.3–0.5 mL) for analyzing hematological parameters, erythrocyte’s deformability (slit-flow ektacytometer) and aggregation (light-transmission aggregometer). Leukocyte and platelet counts raised markedly in gonadectomized animals during the investigated days. Hemorheological changes of I/R showed gender differences: significant impairment of erythrocyte deformability was found on the 1st-3rd postoperative days, expressed mostly in females. In gonadectomized females the postischemic deformability values were impaired. Erythrocyte aggregation index significantly raised by the 1st postoperative day, dominantly in males. It is suggested that gonadectomy may act as an additional rheological “risk factor” related to blood micro-rheological parameters in ischemia-reperfusion.

Keywords: Ischemia-reperfusion, gonadectomy, gender differences, red blood cell aggregation, red blood cell deformability

1. Introduction

Numerous literature data support the fact that ischemia-reperfusion (I/R) and related complex pathophysiological processes may cause significant changes in blood rheological parameters, especially in red blood cell deformability and aggregation, in which changes the harmful and cascade-like complex effects of oxygen-derived free radicals and leukocyte activation play pivotal roles [4, 8, 18–22, 34, 35, 38, 51]. These rheological changes may aggravate further the microcirculatory disorders [4, 19]. The magnitude of the changes are depending on the ischemic duration, the type of the affected tissue and the temperature [18, 35, 49]. Hemorheological changes during and after I/R could be observed in the early reperfusion

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period [18, 21, 38] and later on, depending on the magnitude of the inflammation during the 1st–3rd postoperative days, reflecting the local and systemic changes as well [22, 34, 35, 38, 49].

The peripheral vascular diseases as well as microcirculatory changes caused by I/R may show gender differences, underlying the importance of sex hormones as well as their concentration and the androgen-to-estrogen ratio [16, 17, 24, 28, 37, 48]. Several in vivo experimental studies proved the protecting effect of estrogen in myocardial, cerebral and hind-limb ischemia-reperfusion injuries [10, 15, 23, 41, 44, 47, 52]. The protective effect seems to be quite complex, including certain antioxidant properties, membrane stabilization effects, improvement of NO release, attenuating Ca\(^{2+}\) accumulation and preserving mitochondrial structure and function [27–30, 39, 47, 48, 52]. It was also demonstrated that estrogen may protect the red blood cells from damage and the presence of testosterone may render the erythrocytes more susceptible to oxidative stress after trauma-hemorrhagic shock [25]. However, the protective or non-protective effect of testosterone is still controversial [12]. Supposedly all of these in vivo effects are more complex than as it is in vitro.

Concerning the hemorheological gender differences in rats [32] and their changes after gonadectomy [33], it is supposed that the effect of I/R on erythrocyte deformability and aggregation may alter in female and males, that might differ further in gonadectomized animals. We aimed to investigate this question in a hind limb I/R follow-up study in rats.

2. Materials and methods

2.1. Experimental animals and study design

The experiments were approved and registered by the University of Debrecen Committee of Animal Research (permission Nr.: 37/2007, 17/2008), in accordance with the relevant Hungarian Animal Protection Act (Law XVIII/1998) and EU Directives (EEC 63/2010).

Twenty-five male and twenty-five female Sprague-Dawley rats (Janvier Co., France) were involved in the study. The animals were aged similarly to each other. The healthy female animals were in pro-estrus phase, according to the investigation of vaginal smears with Giemsa dying.

2.2. Operative techniques and sampling protocol

The general anesthesia was provided by intraperitoneal administration of Thiopental\(^{b}\) (60 mg/kg).

Six experimental groups were formed:

I. Control males (n = 8; 573.6 ± 82 g)

II. Control females (n = 8; 326 ± 11.8 g): besides of 2-hours anesthesia and blood samplings no further intervention was made.

III. Ischemia-reperfusion (I/R) males (n = 7; 586.2 ± 54.1 g) and IV. Ischemia-reperfusion (I/R) females (n = 7; 306.7 ± 14.9 g): a tourniquet was placed around the left thigh closed to the level of the inguinal ligament. The completion of ischemia was checked by laser Doppler tissue flowmetry (LD-1, Experimetria Ltd., Hungary) placed on the paws. The ischemia was maintained for 1 hour.

V. Gonadectomy + ischemia-reperfusion (GoE + I/R) males (n = 10; 508.8 ± 33.2 g) and VI. Gonadectomy + ischemia-reperfusion (GoE + I/R) females (n = 10; 384.8 ± 43.3 g): three months after bilateral orchidectomy or ovariectomy [33] the same protocol was completed as in I/R groups.
After the ischemia and on the 1st postoperative day the animals received analgesics (Flunixin, 2.5 mg/kg b.w., s.c.). In the control animals the same dosage was used, in parallel.

By puncturing the lateral tail vein 0.3–0.5 mL blood was taken as base sample (anti-coagulant: sodium-EDTA, 1.5 mg/ml) from each animal. By the same way, 5 minutes after removing the tourniquet (postischemic sample), on the 1st, 2nd, 3rd and 7th postoperative days further blood samples were taken (0.3–0.5 mL per each), and in parallel from the Control animals. On the 7th day the blood sampling was completed under general anesthesia and the animals were sacrificed, ending the experiment.

2.3. Laboratory investigations

A Sysmex F-800 microcell counter (TOA Medical Electronics Co. Ltd., Japan) was used for determining general hematological parameters. Red blood cell count (RBC [×10^6/l]), hematocrit (Hct [%]), hemoglobin (Hgb [g/dl]), mean corpuscular volume (MCV [fl]), mean corpuscular hemoglobin (MCH [pg]), mean corpuscular hemoglobin concentration (MCHC [g/dl]), white blood cell count (WBC [×10^3/l]), monocyte-granulocyte and lymphocyte ratio (Mo-Gr, %, Lymph %) and platelet count (Plt [×10^3/l]) were analyzed. A test requires approximately 70 μl of blood.

Red blood cell deformability was measured by a Rheoscan-D200 ektacytometer (Sewon Meditech Inc., Korea) [42, 43]. For the measurements isotonic solution of polyvinylpyrrolidone (360 kDa, viscosity = 28.8 mPa.s, osmolarity = 305 mOsm/kg, pH = 7.36) was prepared and 6 μl of native blood was taken into 0.6 mL of the PVP solution, and gently mixed. The sample suspension was taken into Rheoscan disposable slit-flow kit, in which the fluid is moving by the force of the vacuum generated by the device, creating the required shear stress profile (∼0.5–20 Pa) for elongating the red blood cells. The red blood cells—while elongating and changing shape—alter the laser diffraction pattern. The elongation index (EI) at a constant shear stress (SS [Pa]) is calculated from the length (L) and width (W) of the diffractogram: $EI = (L - W)/(L + W)$. EI increases with cell deformability [5]. For comparison EI values at 3 Pa were used and parameterization of individual curves Taniweaver-Burke analysis was completed, calculating the maximal elongation index (EI_{max}) and the shear stress at half EI_{max} (SS_{1/2}[Pa]) [7].

Red blood cell aggregation was tested using light transmission aggregometry (Myrenne MA-1 erythrocyte aggregometer, Myrenne GmbH, Germany). This aggregometer determines M (at 0 shear rate) and M1 (at shear rate of 3 s⁻¹) indices, reflecting the magnitude of red blood cell ‘clumping’ at the 5th or 10th seconds of the aggregation process [5, 40]. M and M1 indices increase with enhanced aggregation. The measurements require approximately 20 μl of blood. The measurements were carried out within 1 hour after sampling [5, 31].

2.4. Statistical analyses

Data are presented as mean ± standard deviation (S.D.). Based on the normality of data distribution, for inter-group comparison Student t-test or Mann-Whitney RS test, for intra-group comparison one-way ANOVA tests (Dunn’s or Bonferroni’s method) were used. A p value less than 0.05 was considered as statistically significant.

For analyzing the magnitude of inter- and intra-group changes, standardized differences were calculated: dividing the mean difference of values (EI at 3 Pa, EI_{max}, SS_{1/2}) between the base (before operation) data and the postischemic or postoperative values at a given SS by the pooled standard deviation of the base and postischemic or postoperative data. The pooled standard deviation is calculated as the square root of the mean of squared standard deviations of the two groups being compared [6, 46].
3. Results

3.1. Hematological parameters

In all I/R groups red blood cell count (RBC [×10⁶/µL]) and hematocrit (Hct [%]) slightly decreased by the 1st and 2nd postoperative day, and started to normalize from the 3rd day. There was no remarkable difference between groups. By the 7th day RBC and Hct values were similar to the normal, base data. Hemoglobin concentration, MCV and MCHC did not show important changes (data not shown).

The changes of white blood cell count (WBC [×10⁹/µL]) and platelet count (Plt [×10⁹/µL]) are shown in Table 1.

Compared to control animals, the ischemia-reperfusion resulted in a rise of WBC count by the 1st postoperative day with increasing monocyte-granulocyte ratio. In I/R males the increase was more expressed compared to females, showing significant gender difference on the 2nd and 3rd postoperative day (p<0.001 and p=0.001, respectively). Interestingly, in females the rise in WBC was diminished, presenting only a smaller, non-significant peak on the 3rd postoperative day.

In GoE + I/R males and females the changes were of larger magnitude. On the 1st day the increased WBC count of GoE + I/R males was significant compared to the base values (p<0.001) and versus control males (p=0.001), I/R males (p=0.002) and GoE + I/R females, too (p<0.001). These differences were existing over the investigated postoperative period, showing slight decrease. On the 7th day the elevated WBC count was still significant (p=0.007 vs. base, p=0.026 vs. I/R male, p=0.01 vs. GoE + I/R).

Additionally, in GoE + I/R females the increased WBC count was significant on the 1st (p<0.001 vs. base, Control females and I/R females), the 2nd (p=0.001), all, the 3rd (p=0.007 vs. base, p=0.006 vs. control females) and the 7th postoperative days (p=0.003 vs. base, p<0.001 vs. control and I/R females) (Table 1).

Compared to Control groups in Plt count of I/R and GoE + I/R groups there was a decrease on the 1st–3rd postoperative days, followed by definitive increase on the 7th day, mostly expressed in GoE + I/R females.

The decrease started in the early postischemic period, that was remarkable in I/R females (p=0.023) vs. base) and GoE + I/R females. On the 1st postoperative day the Plt count decrease was significant in all I/R groups (I/R males: p=0.006, I/R females: p=0.001, GoE + I/R males: p=0.005, GoE + I/R females: p=0.034), showing differences between the two GoE + I/R group, too (p=0.011). On the 2nd day further decrease was observed in all I/R and GoE + I/R groups, then from the 3rd day slight increase started, reaching markedly elevated Plt count on the 7th day, mostly in GoE + I/R groups. In GoE + I/R males the 7th-day Plt count was significant versus its base (p<0.001) and compared to Control males (p=0.003); in GoE + I/R females the rise was significant compared to base values (p<0.001) and versus 7th-day values of I/R females, too (p=0.004) (Table 1).

3.2. Red blood cell deformability

Figure 1 shows the cumulated elongation index (EI) – shear stress (SS) curves in control, I/R and GoE + I/R male and female groups.

In Control male and female groups there were no remarkable changes during the investigated period (Fig. 1A, B). In I/R groups significant impairment of red blood cell deformability was observed on the 1st–3rd postoperative days, expressed mostly by females (Fig. 1C, D). In GoE + I/R females the most decreased EI values were measured in postischemic samples (Fig. 1E, F).
Table 1
Changes of white blood cell (WBC) count and platelet (Plt) count in Control, Ischemia-Reperfusion (I/R) and Gonadectomy+Ischemia-Reperfusion (GoE + I/R) male and female group.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Group</th>
<th>Gender</th>
<th>Base</th>
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<th>Postoperative days</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1st</td>
<td>2nd</td>
</tr>
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<td></td>
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<td>5 ± 1.52</td>
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<td></td>
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<td>5.2 ± 1.08</td>
<td>5.9 ± 0.89</td>
<td>6.2 ± 0.32</td>
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<td>female</td>
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<td>7.7 ± 3.54</td>
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<td>8.2 ± 2.74</td>
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<td>707.9 ± 140.4</td>
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<td>759.6 ± 256.3</td>
<td>732.1 ± 108.6</td>
<td>785.1 ± 143.6</td>
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<td>505.3 ± 109.4</td>
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<td></td>
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<td>723.1 ± 124.9</td>
<td>655.8 ± 143.4</td>
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<tr>
<td></td>
<td>female</td>
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<td>606.4 ± 203.4</td>
<td>529.7 ± 229.4</td>
<td>662.8 ± 298.3</td>
</tr>
</tbody>
</table>

Means ± S.D.

* p < 0.05 vs. base; # vs. female; # vs. Control (same gender); # vs. I/R (same gender).
Fig. 1. Elongation index (EI) values of male and female Control (A, B), Ischemia-reperfusion (I/R) (C, D) and Gonadectomy + Ischemia-Reperfusion (GoEI/R) groups (E, F) in the function of shear stress (SS [Pa]) before (preop.) and after ischemia (postop.) on the 1st, 3rd, 5th and 7th postoperative days. Means ± S.D.

EI values at shear stress of 3 Pa remarkably decreased in GoE+I/R females just after ischemia that was significant compared to base values ($p = 0.003$), I/R females ($p < 0.001$) and versus GoE+I/R males ($p = 0.016$). On the 1st postoperative day I/R females as well as GoE+I/R males and females showed decrease EI values ($p < 0.05$). In GoE+I/R males the difference was significant compared to the I/R male group, too ($p = 0.025$). The 3rd day brought further impairment in I/R males and females. In I/R males the
decrease of EI was significant compared to GoE + I/R males \( (p = 0.013) \). In I/R females the impairment was more obvious, being significant versus base values \( (p < 0.001) \) and Control females \( (p < 0.001) \). By the 7th day EI values seemed to be normalized (Fig. 2A).

The postischemic standardized difference was the highest in GoE + I/R females \( (1.09) \), while in the 1st–3rd postoperative days the highest values were expressed by I/R females \( (1.06, 0.97 \text{ and } 2.28, \text{ respectively}) \) (Fig. 2B).

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**Fig. 2.** Elongation index values at shear stress of 3 Pa in Control (C), Ischemia-Reperfusion (I/R) and Gonadectomy + Ischemia-Reperfusion (GoE + I/R) male and female groups (A), and the standardized difference values (B) compared to base. means ± S.D. * \( p < 0.05 \) vs base; # vs female; + vs Control (same gender); % vs I/R (same gender).
The calculated $E_{\text{Imax}}$ reflected the above described changes. The lowest postichemic values were found in GoE + I/R females ($p = 0.011$ vs. base, $p < 0.001$ vs. I/R females). The difference between GoE + I/R males and females did not reach the significant level ($p = 0.068$). On the 1st postoperative day I/R females and both GoE + I/R male and female groups showed the lowest $E_{\text{Imax}}$ values, all being significant versus base ($p < 0.001$, $p = 0.002$ and $p = 0.032$, respectively). In I/R females the difference was significant compared to Control females ($p = 0.005$). In GoE + I/R significant difference was found versus Control.

Fig. 3. Calculated maximal elongation index values ($E_{\text{Imax}}$) in Control (C), Ischemia-Reperfusion (I/R) and Gonadectomy + Ischemia-Reperfusion (GoE + I/R) male and female groups (A), and the standardized difference values (B) compared to base. means ± S.D. * $p < 0.05$ vs base, # vs female; + vs Control (same gender); ł vs GoE (same gender); vs I/R (same gender).
females \((p<0.001)\) and GoE + I/R males \((p=0.044)\). GoE + I/R males expressed lower \(E_{\text{max}}\) values compared to Control males \((p=0.007)\) and I/R males, too \((p=0.002)\). On the 3rd postoperative day the lowest values were found in I/R females \((p<0.001\) vs. base, \(p=0.004\) vs. Control females, \(p=0.003\) vs. I/R males and \(p<0.001\) compared to GoE + I/R females) (Fig 3A).

The standardized difference was the highest in GoE + I/R females in posts ischemic samples (0.77), and in I/R females on the 1st (1.39), the 3rd (2.43) and by the 7th (1.76) postoperative days (Fig. 3B).

Shear stress values at half \(E_{\text{max}}\) \((SS_{1/2} \text{[Pa]})\) were increased in GoE + I/R posts ischemic \((p=0.006\) vs. base) and 1st-day samples \((p=0.021\) vs. base, \(p=0.002\) vs. Control females), and in I/R females on

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**Fig. 4.** Calculated shear stress values at half maximal elongation \((SS_{1/2} \text{[Pa]})\) in Control (C), Ischemia-Reperfusion (I/R) and Gonadectomy + Ischemia-Reperfusion (GoE + I/R) male and female groups (A), and the standardized difference values (B) compared to base. means ± S.D. \(* p<0.05\) vs base, \# vs female; \# vs Control (same gender); \$ vs GoE (same gender).
the 2nd day ($p=0.005$ vs. base, $p<0.001$ vs. GoE + I/R females), furthermore, in I/R males on the 3rd postoperative day ($p=0.012$ vs. GoE + I/R males) (Fig. 4A).

The standardized difference was the largest in magnitude in postischemic samples of GoE + I/R females (−0.8) and in I/R females on the 2nd day (−1.27) (Fig. 4B).

3.3. Red blood cell aggregation

Figure 5 shows changes of aggregation index values M and M1 at 5- or 10-second mode in I/R and GoE + I/R male and female groups.

In postischemic samples of GoE + I/R females M (5 s and 10 s) index values increased, being significant versus base (10 s: $p=0.019$) and I/R females (5 s: $p<0.001$).

On the 1st postoperative day all the index values remarkably increased, showing significance compared to base values (M 5 s: I/R males $p=0.003$, I/R females $p=0.008$, GoE + I/R males $p<0.001$, GoE + I/R females $p=0.031$; M 10 s: I/R males $p<0.001$, I/R females $p=0.002$, GoE + I/R males $p<0.001$, GoE + I/R females $p=0.005$; M 15 s: I/R males $p<0.001$, I/R females $p=0.012$, GoE + I/R males $p<0.001$, GoE + I/R females $p=0.03$; M 110 s: I/R males $p<0.001$, I/R females $p<0.001$, GoE + I/R males $p=0.03$, GoE + I/R females $p=0.016$).

On the 3rd postoperative day significant increase of aggregation was found interestingly in I/R male group when measuring M 10 s ($p=0.015$ vs. base and $p=0.015$ vs. I/R females), M1 5 s ($p=0.003$ vs.

![Figure 5](image_url)
base, \( p < 0.001 \) vs. I/R females and \( p < 0.001 \) vs. GoE + I/R males) and M1 10 s values \( (p < 0.001 \) vs. base, I/R females and GoE + I/R males).

4. Discussion

It is known that red blood cell deformability is determined by cell volume, surface-to-volume ratio, morphology, cytoskeletal properties, inner viscosity as well as membrane viscosity \[26\]. The other important micro-rheological parameter, the red blood cell aggregation is influenced by both cellular (deformability, cell morphology, membrane glyocalyx structure) and plasmatic factors (fibrinogen concentration, micro-environmental conditions) \[36\]. These factors can be influenced by pathophysiological processes during I/R at several points. Oxygen-derived free radicals are among the most important agents causing impaired cell deformability. The reactive free radicals cause direct and cascade-like harmful effects, including lipid peroxidation of the membrane, methemoglobin and Heinz-body formation – so increasing the inner viscosity of the cells as well as protein modification (e.g. by sulfhydryl cross-linking) – and consequent functional changes in transmembrane proteins, ion channels, structural proteins \[4, 8, 13, 34, 45\]. Changes in micro-environmental conditions (e.g. pH, osmolarity, lactate concentration) as well as inflammatory processes, and as a part of acute phase reaction, the increased fibrinogen may cause enhanced red blood cell aggregation \[4, 8, 18, 21, 22\].

Growing number of data underline the importance of hemorheological gender differences not only in the clinical investigations but also in the experimental medicine \[32\]. Furthermore, gonadectomy may additionally influence these micro-rheological gender differences, reflecting the influence of sex hormones, too \[33\]. The protective effect of estrogen has been demonstrated in numerous I/R studies including acute \[47\] and chronic hind limb \[23\], myocardial \[10, 27, 30, 37, 52\], brain \[15, 41\] and intestinal ischemia \[48\]. The effect of testosterone (protective or non-protective influence) is still controversial \[12, 39, 48\]. However, little is known related to micro-rheological changes.

In current study we aimed to investigate the effect of hind limb I/R on erythrocyte deformability and aggregation focusing on potential differences in female and males, that might differ further in gonadectomized rats.

The main findings were the following: (1) I/R resulted in a rise of WBC count by the 1st postoperative day with increasing monocyte-granulocyte ratio. In I/R males the increase was more expressed compared to females, showing significant gender difference on the 2nd and 3rd postoperative day. In GoE + I/R males and females the changes were of larger magnitude. (2) Compared to Control groups in Plt count of I/R and GoE + I/R groups there was a decrease on the 1st-3rd postoperative days, followed by definitive increase on the 7th day, mostly expressed in GoE + I/R females. (3) In I/R groups significant impairment of red blood cell deformability was observed on the 1st–3rd postoperative days, expressed mostly by females. (4) In GoE + I/R females the most decreased EI values were measured in postischemic samples, together with significantly increased aggregation index values. (5) On the 1st postoperative day aggregation index values remarkably increased in all I/R groups, while on the 3rd day significant enhancement in aggregation was found only in I/R male group.

In a rat model of hind limb I/R marked impairment of red blood cell deformability has been observed on the 1st and 2nd postoperative day, together with rise in leukocyte count and also an elevation of platelet count over the early postoperative days \[34\]. In that study by using microclips on femoral vessels 1-hour ischemia was induced. On the 1st–7th postoperative days small quantity of blood samples were taken for determining red blood cell deformability (by bulk filtrometry) and hematological parameters. The highly
significant increase of relative cell transit time, reflecting impairment of red blood cell deformability, was seen on the 1st and 2nd postoperative days. These changes could be prevented by giving allopurinol, inhibitor of xanthine oxidase enzyme known to be the major source of superoxide during reperfusion [34].

According to these findings we have chosen the follow-up period of one week and the critical days to be investigated: 1st – 3rd and 7th postoperative days. The current findings nicely correlate with the previous results: the 1st and 3rd postoperative days were critical regarding the impairment of red blood cell micro-rheological properties.

The magnitude of changes in red blood cell deformability and red blood cell aggregation and their correlation to each other raise further questions. It is difficult to estimate the real relation between red blood cell deformability and red blood cell aggregation, and their changes. Interestingly, the females with lower bodyweight showed larger changes during the early postoperative days compared to males, while gonadectomized females expressed more obvious impairment in erythrocyte deformability just after ischemia.

The gonadectomized females had significantly augmented bodyweight compared to the same-age females of control and I/R groups. The obesity is known to affect numerous physiological parameters and pathways, causing abnormal insulin sensitivity, increased vasomotor tone, dyslipidemia, abnormalities in organs (e.g., liver, kidney), endothelial dysfunction and inflammation [1, 3, 50]. It has been reported that in obesity the bioavailability of nitric oxide (NO) is reduced, due to peroxynitrite production by deliberating superoxide anion [3, 9]. Furthermore obesity may affect guanylate cyclase pathways, formation of angiotensin II, expression of endothelin-1, and vasoconstrictor prostanoids [3, 50]. Obesity is also known to reduce the effect of ischemic preconditioning by enhanced mitochondrial oxidative stress, abnormal function of mito-KATP channels as well as due to the adverse effects of leptin, activating NADPH oxidase [2, 14].

Since NO is known to improve [11], reactive oxygen-derived free radicals are known to impair red blood cell deformability [4, 8, 34], it is supposed that the existing obesity in ovariectomized females could affect the magnitude of red blood cell deformability changes in the early reperfusion period.

5. Conclusions

Hemorheological effects of 1-hour hind limb ischemia and the following reperfusion showed gender differences in rats, reflecting significant impairment of red blood cell deformability on the 1st–3rd postoperative days, mostly expressed in females. In gonadectomized females the posts ischemic elongation index values were the lowest with enhanced red blood cell aggregation. Aggregation index values significantly raised by the 1st postoperative day after I/R, dominantly in males. White blood cell count and platelet count raised more markedly in gonadectomized animals during the early postoperative days.

According to the findings, further question arises whether the gonadectomy may act as an additional rheological risk factor related to blood micro-rheological parameters in ischemia-reperfusion. This issue needs further study to clarifying the findings.

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