Effect of picroside II on hind limb ischemia reperfusion injury in rats

Introduction: Many structural and functional damages are observed in cells and tissues after reperfusion of previously viable ischemic tissues. Acute ischemia reperfusion (I/R) injury of lower extremities occurs especially when a temporary cross-clamp is applied to the abdominal aorta during aortic surgery. Research regarding the treatment of I/R injury has been increasing day-by-day. In this study, we aimed to investigate the effect of picroside II on skeletal muscle of rats experiencing simulated I/R.

Materials and methods: Twenty-four male Wistar albino rats weighing between 210 and 300 g were used in this study. Rats were randomly divided into 4 groups of 6 rats each (control, I/R, control + picroside II, and I/R + picroside II). The infrarenal section of the abdominal aorta was occluded with an atraumatic microvascular clamp in I/R group. The clamp was removed after 120 minutes and reperfusion was provided for a further 120 minutes. Picroside II (10 mg kg⁻¹) was administered intraperitoneally to the animals in control + picroside II and I/R + picroside II groups. At the end of the study, skeletal muscle tissue was obtained for the determination of total oxidant status (TOS) and total antioxidant status (TAS) levels. Apoptosis was evaluated by TUNEL experiment.

Results: TOS levels were significantly higher in I/R group than that of control and I/R + picroside II groups (P=0.014, P=0.005, respectively). TAS levels were significantly higher in I/R group than that of control and I/R + picroside II groups (P=0.007 P=0.005, respectively). TUNEL assay revealed that picroside II reduced cell necrosis.

Conclusion: The results of this study demonstrated that picroside II plays a critical role to prevent I/R injury. Even though our results were found to be satisfactory, it should be encouraging to those who want to conduct future research on this topic.

Keywords: ischemia reperfusion, picroside II, hind limb skeletal muscle, TOS, TAS
Peroxynitrite, known to oxidize sulphhydryl groups and to yield products indicative of hydroxyl radical (OH) reaction with deoxyribose and dimethyl sulfoxide, causes direct cellular injury by inducing lipid and protein peroxidation and damage nucleic acids.\textsuperscript{1,2}

Oxygen free radicals are the most important and toxic substances produced in various clinical conditions.\textsuperscript{3} The common characteristics in all of the clinical situations is either hypoxic microenvironments followed by reoxygenation or ischemic microenvironments followed by reperfusion. In general, as reperfusion progresses, systemic inflammatory response syndrome and multiple organ failure (excretory, respiratory, circulatory system, etc.) follow local edema and muscle necrosis.\textsuperscript{4,8}

Picroside II is a glycoside derivative, and its chemical structural formula is $\text{C}_{23}\text{H}_{28}\text{O}_{13}$. It is one of the primary active constituents isolated from \textit{Picrohiza scrophulariiflora}. It belongs to the family, Scrophulariaceae. Several pharmacological effects of picroside II, such as antioxidant,\textsuperscript{9-11} anti-apoptotic,\textsuperscript{12,13} anti-inflammatory,\textsuperscript{12,13} anticarcinogenic,\textsuperscript{14} neuroprotective,\textsuperscript{15,16} hepatoprotective,\textsuperscript{17} anticholestatic effects, and immune modulating activities\textsuperscript{18} have been demonstrated in various studies.

While most of the studies focused on myocardial ischemia/reperfusion (I/R) injury, the primary aim of this study was to investigate the role of picroside II in an experimental model of rat lower limb muscle I/R injury.

Materials and methods

Animals and experimental protocol

This study was performed at Gazi University GUDAM Laboratory with the approval of the Ethics Committee of Experimental Animals of our university. All of the procedures were performed according to accepted standards of Guide for the Care and Use of Laboratory Animals.

In this study, 24 Wistar Albino rats weighing between 210 and 300 g, raised under the same environmental conditions, were used. The rats were kept under 24°C in cycles of 12-hour daylight and 12-hour darkness and had free access to food until 2 hours before the anesthesia procedure. The animals were randomly separated into four groups of 6 rats each. Midline laparotomy was performed under ketamine (100 mg kg\textsuperscript{-1}) anesthesia.

Control (C) group

A midline laparotomy was performed without any extra surgical intervention. After 4 hours of follow-up, blood sample from abdominal aorta and tissue samples from hind limb skeletal muscles were collected. Tissue samples from hind limb skeletal muscles of rats were obtained for immunohistochcmical and biochemical evaluations, and then animals were sacrificed.

Control + picroside II (P) group

All steps were performed similar to C group but in addition picroside II was intraperitoneal administered (10 mg kg\textsuperscript{-1}) 30 minutes before ischemia. After collecting blood and tissue samples, rats were sacrificed at the end of 4 hours.

I/R group

Midline laparotomy was performed as previously mentioned. Infrarenal aorta was clamped for 2 hours. The clamp was removed and then reperfusion was started. Reperfusion lasted for 2 more hours. In the end, after blood and tissue sampling, rats were sacrificed.

I/R + picroside II (I/R + P) group

All steps were performed similar to C group but in addition to the procedure, picroside II was intra-peritoneal administered (10 mg kg\textsuperscript{-1}) 30 minutes before ischemia. After collecting blood and tissue samples, rats were sacrificed at the end of 2 hours of reperfusion period.

Preparation and application of picroside II

Picroside II was purchased from Sigma-Aldrich Co. Ltd. (St Louis, MO, USA; CAS No: 39012-20-9, purity >98%, molecular formula: $\text{C}_{23}\text{H}_{28}\text{O}_{13}$). It was diluted to a 1% solution with normal saline. The animals in the treatment group were administered an intraperitoneal injection of picroside II 30 minutes before ischemia (10 mg kg\textsuperscript{-1}); the animals in the C and I/R groups were injected with the same volume of 0.09% NaCl. After collecting samples, rats were sacrificed at the end of reperfusion period that lasted for 2 hours.

Homogenization of tissues

The hind limb tissue was collected in a sterile Eppendorf tube and was stored at –80°C until total antioxidant/oxidant status and oxidative stress index analysis. Without being allowed to thaw, the tissues were quickly weighed on a precision scale and separated as 80–100 mg using a no 22 lancet (PLUSMED\textsuperscript{4}). The frozen tissue samples were crushed in the presence of liquid nitrogen in a porcelain bowl. The powdered tissue was transferred to the homogenization tube, and for each gram of tissue, 140 mM KCl solution was added such that the dilution was 1/10. To prevent increase in temperature during homogenization, the tube was held in a snow-filled glass beaker before and after homogenization.
with a homogenizer (Glas_Col K5424®) at a speed of 50 revolutions per minute (rpm) for 2 minutes. Homogenates were transferred to Eppendorf tubes, and the tubes were coated with parafilm and then centrifuged (Hettich Micro 200R®) for 10 minutes at 3,000 rpm. After centrifugation, the supernatant was collected in another Eppendorf tube and the total oxidative status (TOS) and total antioxidant status (TAS) were measured.

**TAS measurement**

TAS was measured by using TAS test kit (RelAssay Diagnostic®, Gaziantep, Turkey), and the test was performed according to the manufacturer’s instructions. Briefly, 500 μL of reagent 1 (measurement buffer) and 30 μL of sample were mixed and absorbance was measured at 660 nm by a spectrophotometer (NanoDrop® ND-1000) (A₂). Next, 75 μL of reagent 2 (colored 2,2-azino-bis-3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) was added to the mixture. The tube was covered with parafilm and incubated for 5 minutes in a hot water bath at 37°C (Lightning Laborteknik®). After incubation, absorbance was measured at 660 nm (A₁). For standard measurement, the Trolox Eq solution at a concentration of 1 mmol/L was used instead of the sample. The first and second measurements were performed thrice, and their averages were calculated. The absorbance change (ΔAbs) was calculated by subtracting the first absorbance value (A₂) from the second absorbance value (A₁). TAS levels were calculated using the formula given in the kit and expressed as mmol H₂O₂ Eq/L.

\[
\text{TAS} = \frac{\Delta \text{Abs standard} - \Delta \text{Abs sample}}{\text{ΔAbs standard}}
\]

**TOS measurement**

TOS was measured by the TOS test kit (RelAssay Diagnostic®, Gaziantep, Turkey), and the test was performed according to the manufacturer’s instructions. Briefly, 500 μL of reagent 1 (measurement buffer) and 30 μL of sample were mixed and absorbance was measured at 530 nm by a spectrophotometer (NanoDrop® ND-1000) (A₂). Next, 25 mL of reagent 2 (Pro-chromogenic solution) was added to the mixture. The tube was covered with parafilm and incubated for 5 minutes in a hot water bath at 37°C (Lightning Laborteknik®). After incubation, absorbance was measured at 530 nm (A₁). A standard solution containing 10 μmol/L hydrogen peroxide (H₂O₂) equivalent/liter given in the kit was used for standard measurement. The first and second measurements were performed thrice, and their averages were calculated. The absorbance change (ΔAbs) was calculated by subtracting the first absorbance value (A₂) from the second absorbance value (A₁). TOS levels were calculated using the formula given in the kit and expressed as mmol H₂O₂ Eq/L.

\[
\text{TOS} = \frac{\Delta \text{Abs sample} \times \text{Standard Concentration (10 μmol/L)}}{\Delta \text{Abs standard}}
\]

**Preparation of histological samples and light microscopic investigations**

For light microscopy, muscle tissue samples were placed in 10% neutral buffered formalin for 72 hours to fix the samples. After fixation routine histological evaluation of tissues was performed. After rinsing tissues in running tap water for an hour, all samples were dehydrated in a graded series of ethanol (75%, 80%, 90%, and 100%, respectively). Xylene was used to clear the tissue, and then tissue was embedded in paraffin blocks. Sections (5 μm) were cut using the Leica RM 2125RT model microtome and then placed on the slide. The slides were held overnight for deparaffinization at 60°C. Xylene was used to clear the slides, and then they were stained by H&E. The sections were examined and photographed by two different histology and embryology specialists who were blinded about the groups on the AxioScope. A. (Carl Zeiss, Oberkochen, Germany) brand light microscope in terms of their morphological and histomorphometric properties. To determine the histological damage score, 3 different sections of the same animal were evaluated according to degeneration and disorganization findings of muscle fibers observed at ×100 magnification, and they were evaluated as follows: 0= normal, (0%); 1= mild, (1%–33%); 2= moderate (34%–66%); and 3= severe (67%–100%). In order to determine the severity of mononuclear cell infiltration in the same section, the aforementioned score table was used.19

**Histomorphometric study for gastrocnemius muscle**

Samples from transverse muscle sections stained with H&E for each muscle tissue were examined by light microscopy in five different microscopic fields using ×200 magnification and the cross-sectional area of each muscle fiber in this area was measured using the AxioVision computer program. The results were evaluated statistically.

**Terminal deoxynucleotidyl transferase-mediated dUTP nick end-labeling (TUNEL)**

TUNEL (in situ end labeling) method was used to determine the apoptotic cells in the tissues. In situ Cell Death Detection Kit (POD Roche Applied Science, Cat # 11684817910) was used according to the manufacturer’s directions.
Sections (4 μm thick) were transferred to polylysine-coated slides and incubated overnight at 60°C. Then, they were deparaffinized in xylene. Sections were rehydrated in graded series of 75%, 80%, 90%, and 100% ethanol and washed with PBS. Slides immersed in 0.1 M citrate buffer were microwaved at 350 W for 5 minutes, and antigen retrieval procedure was performed. PBS was used for washing. Sections were incubated with the TUNEL reaction mixture (labeling solution + terminal deoxynucleotidyl transferase (TdT) enzyme solution) at 37°C for 1 hour in a humidified chamber and washed twice in PBS. For negative control, some sections were incubated with label solution only (without terminal transferase) instead of TUNEL reaction mixture. After this step, Converter-POD was added, and the slides were incubated for 30 minutes at 37°C and then rinsed twice with PBS. The sections were then treated with 3,3-diaminobenzidine at room temperature and rinsed twice with PBS. Sections were counterstained with Harris’ Hematoxylin for 10 minutes. The sections were dehydrated by passing them through series of graded alcohol concentrations. Xylene was used as a clearing agent, and then, the sections were coverslipped with Entellan for light microscopical examination.

**TUNEL outcome evaluations**

Immunohistochemically, immunoreactive cells for TUNEL staining in tissues were counted in 5 different fields in each section with a AxioScope.A1 (Carl Zeiss, Oberkochen, Germany) light microscope at a magnification of x 40. The apoptosis index of each group was calculated as percentage (%) according to the following formula:

\[
\text{Apoptosis index (%)} = \frac{\text{Number of TUNEL positive cells}}{\text{Total number of cells}} \times 100
\]

**Statistical analyses**

Statistical analyses were performed with SPSS (Chicago, IL, USA) 20.0 program. *P*-value <0.05 was considered statistically significant. Kolmogorov–Smirnov test was used to compare the distribution of all variable groups. The variations in TOS and TAS levels were assessed by using the Kruskal–Wallis test. To determine which group differs from the others, we used the Bonferroni-adjusted Mann–Whitney *U* test after the Kruskal–Wallis test. The data were expressed as mean ± standard deviation. 

**Results**

When the groups (skeletal muscle tissue) were compared in terms of TOS enzyme activity, there was a significant difference between the groups (*P*=0.023). Significantly higher TOS enzyme activity was found in I/R group than that of C and I/R + P groups (*P*=0.014, *P*=0.005, respectively) (Table 1).

There was a statistically significant difference between the groups when they were compared among themselves by means of TAS enzyme activity in skeletal muscle tissue (*P*=0.017). TAS enzyme activity in I/R group was found significantly higher than that of C and I/R + P groups (*P*=0.007 *P*=0.005, respectively) (Table 1).

There was a significant difference between the groups when number of TUNEL-positive cells in skeletal muscle tissue were compared (*P*<0.0001). The number of TUNEL-positive cells was significantly higher in I/R group than that of C, P, and I/R + P groups (*P*<0.0001, *P*<0.0001, *P*<0.0001, respectively). In addition, the number of TUNEL-positive cells in I/R + P group was significantly higher than that of C and P groups (*P*<0.0001, *P*<0.0001, respectively) (Table 2, Figures 1–4).

There was a significant difference in mononuclear cell infiltration in skeletal muscle tissue when the groups were compared (*P*<0.0001). The mononuclear cell infiltration was significantly higher in I/R group than that of C, P, and I/R + P groups (*P*<0.0001, *P*<0.0001, *P*=0.027, respectively). In addition, the mononuclear cell infiltration in I/R + P group was significantly higher than that of C and P groups (*P*<0.0001, *P*=0.002, respectively) (Table 2, Figures 5–11).

When the groups were compared in terms of skeletal muscle injury, there was a significant difference between the (*P*<0.0001). Skeletal muscle injury was significantly higher in I/R group than that of C, P, and I/R + P groups

| Table 1 Oxidant status parameters in rat skeletal muscle tissue (mean ± standard deviation) |
|-----------------------------------------------|----------------|----------------|----------------|----------------|----------------|
| (n=6)                                        | (n=6)          | (n=6)          | (n=6)          | (n=6)          | P-value**      |
| TOS (mmol H₂O₂ Eq/L)                         | 0.49±0.08*     | 0.57±0.12      | 0.66±0.07      | 0.48±0.10*     | 0.023          |
| TAS (mmol Trolox Eq/L)                       | 9.99±2.09*     | 11.63±2.94     | 14.35±3.19     | 9.23±2.83*     | 0.017          |

*Notes:* *P*<0.05, compared with group I/R; *P*-value, Kruskal–Wallis test significance level *P*<0.05. 

*Abbreviations:* C, control; P, control + picroside II; I/R, ischemia reperfusion; I/R + P, ischemia reperfusion + picroside II; TAS, total antioxidant status; TOS, total oxidative status.
Table 2 TUNEL positive cell data in skeletal muscle of rats (mean ± standard deviation)

<table>
<thead>
<tr>
<th></th>
<th>Group C (n=6)</th>
<th>Group P (n=6)</th>
<th>Group I/R (n=6)</th>
<th>Group I/R + P (n=6)</th>
<th>P-value**</th>
</tr>
</thead>
<tbody>
<tr>
<td>TUNEL positive cell</td>
<td>9.00±5.76**</td>
<td>12.00±5.59**</td>
<td>90.17±6.97</td>
<td>48.17±10.28*</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Mononuclear cell infiltration</td>
<td>0.00±0.00</td>
<td>0.33±0.18</td>
<td>2.00±0.68</td>
<td>1.33±0.52*</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Injury</td>
<td>0.00±0.00</td>
<td>0.16±0.16</td>
<td>2.67±0.52</td>
<td>1.67±0.52*</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Cross-section</td>
<td>2.821.49±79.92</td>
<td>2.829.06±155.64</td>
<td>1.167.88±373.73</td>
<td>2.870.25±294.32</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Notes: *P<0.05, compared with group I/R; **p-value, Kruskal–Wallis test significance level P<0.05; ***P<0.05, compared with group I/R + P.

Abbreviations: C, control; P, control + picroside II; I/R, ischemia reperfusion; I/R + P, ischemia reperfusion + picroside II.

(P<0.0001, P<0.0001, P=0.003, respectively). In addition, skeletal muscle injury in the I/R + P group was significantly higher than that of C and P groups (P<0.0001, P<0.0001, respectively) (Table 2, Figures 5–11).

There was a significant difference between the groups in terms of cross-section of skeletal muscle (P<0.0001). The cross-section was significantly higher in I/R group than that of C, P, and I/R + P groups (P<0.0001, P<0.0001, P<0.0001, respectively) (Table 2, Figures 6–11).

In cross-sections examined in the C and P groups, it was observed that the muscle fibers were rounded, or they were found to have a slightly angular polygonal shape if they were compressing to each other. It was observed that the muscle fibers in the fascicle surrounded by the perimysium were tightly packed together, and the interstitial area was extremely narrow. In addition to the sections containing one or more nuclei located just below the sarcolemma, there were also nucleus-free cross-sections. Blood vessels and nerve sections were observed in connective tissue sheaths. Cross-sections of myofibrils were seen on the cross-sectional surfaces of muscle fibers (Figures 5 and 6).

In the I/R group, it was seen that the cross-section of the muscle fibers was round or irregular; the distance between the muscle fibers surrounded by the endomysium, where the core was placed mid-point, was increased. The presence of atrophied muscle fibers attracted our attention. Mononuclear cell infiltration was observed with thickening in the perimysium surrounding the muscle fascicles, more prominently around the vessel (Figures 7–9).

In the I/R + P group cross-sectional surfaces of the muscle fibers were observed to be rounded occasionally, and in some of the fibers, mid-point settlement of the core was remarkable. Increase in the area between muscle fibers and mild mononuclear cell infiltration were observed. Although the atrophic muscle fibers were seen in the specimens occasionally, there were muscle fibers showing a normal multifaceted cross-sectional surface in some areas (Figures 10 and 11).

Discussion

Picroside II (β-d-glucopyranoside, 1a,1b,2,5a,6,6a-hexahydro-6-{(4-hydroxy-3-methoxybenzoyl)oxy}-1-a(hydroxymethyl)oxireno[4,5]cyclopenta[1,2-c]pyran-2-yl) is a major iridoid glycoside isolated from P. scrophulariiflora Pennell (Scrophulariaceae). Several pharmacological effects of picroside II such as antioxidant, anti-apoptotic, anti-inflammatory, anticarcinogenic, neuroprotective, anti-inflammatory, anticarcinogenic, neuroprotective, anti-inflammatory, anticarcinogenic, neuroprotective.
Figure 3 In the micrograph of the ischemia reperfusion group, brown staining is observed in apoptotic cell nuclei showing immunoreaction by TUNEL staining. Magnification: ×400.

Figure 4 In the micrograph of the ischemia reperfusion + picroside II group, brown staining is observed in apoptotic cell nuclei showing immunoreaction by TUNEL staining. Magnification: ×400.

Figure 5 The cross-sectional faces of striated skeletal muscle fibers are observed in a multi-sided appearance in the micrograph of the control group. Peripheral nuclei (arrow) under the sarcolemma of muscle fibers, cross sections of myofibrils (asterisk) in the cytoplasm are observed and in the perimysium (P) surrounding the fascicles formed by muscle fibers, blood vessels (arrowheads) draw attention. H&E ×200.

Figure 6 The cross-sectional faces of striated skeletal muscle fibers are observed in a multi-sided appearance in the micrograph of the picroside group. Peripheral nuclei (arrow) under the sarcolemma of muscle fibers and blood vessels (arrowheads) in the perimysium (P) surrounding the fascicles formed by muscle fibers are observed. H&E ×200.

Figure 7 In the micrograph of the ischemia reperfusion group, the cross-sectional faces of the muscle fibers are observed to have a rounded (arrow) shape. Increase in endomysium distance between the fibers (star) and atrophy in the muscle fibers (tailed arrow) are also observed. H&E ×200.

Figure 8 Cross-sections of the fibers in the micrograph of the ischemia reperfusion group show a rounded shape of the cross-sections and middle position of the core (arrow), advanced atrophy in the muscle fibers are observed. H&E ×400.
Picroside II in the prevention of ischemia reperfusion injury

Unpaired electrons and free radicals are highly reactive and readily take part in chemical reactions with virtually all cell components (lipids, proteins, complex carbohydrates, and nucleic acids) in the body. These reactions occur through a chain of oxidative reactions to cause tissue injury. For most biological structures (such as lipids, proteins, and nucleic acids), free radical damage is closely associated with oxidative damage, causing direct cellular injury by inducing lipid and protein peroxidation and damaging nucleic acids. I/R results in the generation of toxic ROS in the organs. It reduces the activity of cellular defense enzymes against ROS, and reperfusion or the introduction of oxygen further disturbs the delicate balance of oxidants/antioxidants and intracellular calcium overload, adenosine triphosphate depletion, myocardial apoptosis, and endothelial dysfunction are all observed consequently. Furthermore, oxidative products such as ROS, reactive nitrogen species, hydrochloric acid, malondialdehyde (MDA), and lipid peroxides constitute TOS.

Zhang et al performed animal experiments and showed that the amount of MDA, nitric oxide (NO), and H₂O₂ can directly reflect the changes in free radical content in vivo in cerebral ischemic injury. The amount of MDA, NO, and H₂O₂ increased significantly in serum and brain tissue after establishing the animal models, and the amount decreased after injecting picroside II intraperitoneally. It was suggested that picroside II can enhance the activity of endogenous antioxidant enzymes and clear the excess oxygen radicals to protect brain tissue from oxidative injury.

In our study, TAS, as the marker of total antioxidant protection against the attack of free radicals in the organism and TOS, as the marker of total value of oxidative stress were used. Studies have shown that TAS measurement provides more valuable information than individual measurements of antioxidants. When we compared skeletal muscle tissue in terms of TOS enzyme activity in our study, there was a significant difference between the groups. TOS enzyme activity was significantly higher in I/R group than that of C and I/R + P groups. Oxidative stress, which is the primary actor of I/R injury, was found to be lower in the picroside II-treated groups in TOS measurement.

In addition, when we compared skeletal muscle tissue in terms of TAS enzyme activity in our study, there was a significant difference between the groups. TAS enzyme activity was found to be significantly higher in I/R group than that of C and I/R + P groups, which is parallel to the fact that antioxidant systems are impaired by I/R damage, and that the low activity of TAS enzyme in the groups given picroside II
indicates that the antioxidant systems are protected, in other words active. Kim et al. found that administration of 2.5 mg vitamin C for 5 days increased the total antioxidant status during that period, and after the administration of 2.5 mg vitamin C for 5 days, TAS was found to be decreased. They thought that this decrease in TAS was related to decreased antioxidant stress. Ayada et al. also observed that nesfatin-1 reduced TOS and also TAS levels under laparotomy and I/R conditions. They thought that chronic peripheral nesfatin-1 administration can reduce oxidant level in both experimental conditions in parallel with inhibition of NO production.

TUNEL method is a staining method used for in situ detection of DNA fragments (breaks) that is an indicator of apoptosis. Li et al. reported that picroside II could have a neuroprotective effect by inhibiting apoptosis and the expressions of NFκB and IκB following middle cerebral artery occlusion/reperfusion in rats. Wang et al. reported that cerebral ischemic injury activated ERK1/2 to induce apoptosis, whereas picroside II down-regulated ERK1/2 expression following cerebral ischemia to inhibit apoptosis in rats. The number of positive inflammatory cells (histiocytes) in epithelioid granulomas in 1 mm² area was the number of TUNEL positive cells, and in our study, there was a significant difference between the groups when the skeletal muscle tissues were compared in terms of number of TUNEL cells. The number of TUNEL cells was significantly higher in I/R group than that of C, P, and I/R + P groups. In addition, the number of TUNEL cells in I/R + P group was significantly higher than that of C and P groups.

When we compared skeletal muscle tissues in terms of mononuclear cell infiltration, in our study, there was a significant difference between the groups. It was significantly higher in I/R group than that of C, P, and I/R + P groups. In addition, mononuclear cell infiltration in I/R + P group was significantly higher than that of C and P groups.

When we compared injury to skeletal muscle tissue, in our study, there was a significant difference between the groups. The injury was significantly higher in I/R group than that of C, P, and I/R + P groups.

There was a significant difference between the groups in terms of skeletal muscle cross-section in our study.

In conclusion, our findings suggest that picroside II has a protective effect against I/R injury of skeletal muscle, but we still think these promising results should further be supported by more detailed studies with larger volumes.

Disclosure
The authors alone are responsible for the content and writing of the paper. The authors report no conflicts of interest in this work.

References