

Effect of tramadol on lung injury induced by skeletal muscle ischemia-reperfusion: an experimental study*

Efeito do tramadol na lesão pulmonar induzida por isquemia-reperfusão de músculo esquelético: um estudo experimental

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Abstract

Objective: To determine whether tramadol has a protective effect against lung injury induced by skeletal muscle ischemia-reperfusion. **Methods:** Twenty Wistar male rats were allocated to one of two groups: ischemia-reperfusion (IR) and ischemia-reperfusion + tramadol (IR+T). The animals were anesthetized with intramuscular injections of ketamine and xylazine (50 mg/kg and 10 mg/kg, respectively). All of the animals underwent 2-h ischemia by occlusion of the femoral artery and 24-h reperfusion. Prior to the occlusion of the femoral artery, 250 IU heparin were administered via the jugular vein in order to prevent clotting. The rats in the IR+T group were treated with tramadol (20 mg/kg i.v.) immediately before reperfusion. After the reperfusion period, the animals were euthanized with pentobarbital (300 mg/kg i.p.), the lungs were carefully removed, and specimens were properly prepared for histopathological and biochemical studies. **Results:** Myeloperoxidase activity and nitric oxide levels were significantly higher in the IR group than in the IR+T group ($p = 0.001$ for both). Histological abnormalities, such as intra-alveolar edema, intra-alveolar hemorrhage, and neutrophil infiltration, were significantly more common in the IR group than in the IR+T group. **Conclusions:** On the basis of our histological and biochemical findings, we conclude that tramadol prevents lung tissue injury after skeletal muscle ischemia-reperfusion.

Keywords: Tramadol; Muscle, skeletal; Ischemic attack, transient; Lung Injury.

Resumo

Objetivo: Investigar se o tramadol tem um efeito protetor contra a lesão pulmonar induzida por isquemia-reperfusão de músculo esquelético. **Métodos:** Vinte ratos Wistar machos foram divididos em dois grupos: grupo isquemia-reperfusão (IR) e grupo isquemia-reperfusão + tramadol (IR+T). Os animais foram anestesiados com cetamina e xilazina (i.m., 50 mg/kg e 10 mg/kg, respectivamente). Todos os animais foram submetidos a 2 h de isquemia por oclusão da artéria femoral e 24 h de reperfusão. Antes da oclusão da artéria femoral, foram administrados 250 UI de heparina pela veia jugular para impedir a coagulação. Os ratos do grupo IR+T foram tratados com tramadol (20 mg/kg i.v.) imediatamente antes da reperfusão. Após o período de reperfusão, os animais foram sacrificados com pentobarbital (300 mg/kg i.p.), os pulmões foram removidos cuidadosamente, e os espécimes foram preparados adequadamente para estudos histopatológicos e bioquímicos. **Resultados:** A atividade de mieloperoxidase e os níveis de óxido nítrico foram significativamente maiores no grupo IR que no grupo IR+T ($p = 0,001$ para ambos). Anormalidades histológicas, como edema intra-alveolar, hemorragia intra-alveolar e infiltração neutrofílica, foram significativamente mais frequentes no grupo IR que no grupo IR+T. **Conclusões:** Com base nos resultados histológicos e bioquímicos deste estudo, concluímos que o tramadol tem um efeito protetor contra o dano ao tecido pulmonar após isquemia-reperfusão de músculo esquelético.

Descritores: Tramadol; Músculo esquelético; Ataque isquêmico transitório; Lesão pulmonar.

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Introduction

Ischemia-reperfusion injury is one of the most common types of cell injury that occurs in a variety of surgical practices. Reperfusion of ischemic organs can result in tissue injury, which manifests as microvascular and parenchymal cell dysfunction. The mechanisms underlying ischemia-reperfusion injury have been previously described; polymorphonuclear leukocytes and reactive oxygen metabolites have been indicated to have pivotal roles in the etiology.⁽¹⁻³⁾

Skeletal muscle ischemia-reperfusion resulting from trauma, limb revascularization, orthopedic surgery, free flap reconstruction, or any other etiology not only leads to muscle damage itself but also causes injury involving a severe destruction of remote organs. Considerable advances have been made in the understanding of the mechanisms of this systemic response regarding the skeletal muscle ischemia-reperfusion sequence. Remote organs with intense microcapillary systems, such as the lungs, are prone to developing this type of systemic injury.^(3,4)

Various investigators have demonstrated that the opioid pathway is involved in tissue preservation during hypoxia or ischemia, and this protection is mediated via the delta opioid receptor.^(5,6) Tramadol hydrochloride is an effective analgesic drug used for severe acute and chronic pain conditions. It has a weak affinity to the μ -opioid receptor and inhibits the reuptake of monoamines in the central nervous system, thereby activating the descending inhibitory systems.^(7,8) Recent research discloses that tramadol decreases lipid peroxidation and regulates noradrenalin uptake, and, therefore, these therapeutic properties are used for the management of myocardial ischemia.⁽⁹⁾

The aim of the present study was to investigate the potential protective effect of tramadol hydrochloride on lung ischemia-reperfusion injury induced by the hind limb model by means of histopathological evaluation and determination of inflammatory responses via myeloperoxidase (MPO) activity and nitric oxide (NO) levels in the lung tissue of rats.

Methods

All of the animals used in the present research were properly cared in accordance with the norms of the Laboratory of Animal Experimentation at the Islamic Azad University Faculty of Specialized Veterinary

Sciences, in Tehran, Iran. The study was approved by the Animal Research Ethics Committee of the Department of Veterinary Surgery of the university.

Twenty Wistar male rats, weighing 250-300 g (12-15 weeks old), were used in the present study. All of the rats were maintained under constant room temperature and standard conditions, with *ad libitum* access to water and commercial food, and placed in individual plastic cages with soft bedding. The animals were divided randomly into two experimental groups of ten rats each: ischemia-reperfusion (IR) group and ischemia-reperfusion + tramadol (IR+T) group. Anesthesia was induced using ketamine and xylazine (i.m., 50 mg/kg and 10 mg/kg, respectively). After induction of anesthesia, the left hind limb was completely clipped. After clipping, disinfecting, and dropping, a skin incision was made on the medial surface of the left hind limb. The femoral artery and vein were isolated from the surrounding tissues, and the femoral artery was exposed and clamped with a mini bulldog forceps. Prior to the occlusion of the femoral artery, 250 IU heparin was administered via the jugular vein in order to prevent clotting. All of the animals underwent 2-h ischemia by the occlusion femoral artery with a vascular clamp and 24-h reperfusion. The animals were maintained in dorsal recumbency and kept anesthetized throughout the duration of the ischemic period. Additional doses of the anesthetics were given as necessary in order to maintain anesthesia during the experiment. Body temperature was maintained with a heating pad under anesthesia. The animals in the IR+T group were administered tramadol i.v. (20 mg/kg)⁽¹⁰⁾ immediately before reperfusion. Following the ischemic period, the vascular clamp was removed, and the surgical site was routinely closed. After the surgery, fluid losses were replaced by intraperitoneal administration of 5-mL warm isotonic saline, and the rats were returned to their cages with *ad libitum* access to commercial food and water during the reperfusion period. After 24 h of reperfusion, the rats were euthanized with an overdose of intraperitoneal pentobarbital injection (300 mg/kg), and the left lungs were harvested and fixed in 10% formaldehyde for histopathological examination under light microscopy. The right lungs were removed and stored at -20°C for analysis. The lung tissue homogenate and supernatant samples were prepared as described by Yildirim et al.⁽¹¹⁾

The biochemical assay consisted of determining MPO activity and NO levels in lung tissue. The activity

of MPO⁽¹²⁾ was analyzed spectrophotometrically as described elsewhere, whereas NO levels in lung tissue were measured by the Griess reaction.⁽¹³⁾

All of the left lung tissue samples were fixed in 10% formalin solution and processed routinely (embedded in paraffin blocks, the anterior lung region being sectioned into 6- μ m sections, and stained with H&E). The severity of lung injury was determined by a pathologist who was blinded to the experiment. Lung injury was classified into four levels, as follows: level 0, no diagnostic change; level 1, mild neutrophil leukocyte infiltration and mild to moderate interstitial congestion; level 2, moderate neutrophil leukocyte infiltration, perivascular edema formation, and partial destruction of pulmonary architecture; and level 3, dense neutrophil leukocyte infiltration and complete destruction of pulmonary structure.⁽¹⁴⁾ A total of four slides from each lung sample were randomly screened, and the mean level was considered representative of the sample.

Statistical analyses were carried out with the Statistical Package for the Social Sciences, version 11.2 (SPSS Inc., Chicago, IL, USA). The distribution of the groups was analyzed with one-sample Kolmogorov-Smirnov test. Biochemical results showed normal distribution, and one-way ANOVA was used. Histopathological results were analyzed using Kruskal-Wallis and Mann-Whitney U tests. Values of $p < 0.05$ were considered as statistically significant.

Results

All of the rats survived until the end of the study period.

Regarding biochemical results, NO levels were significantly higher in the lungs of the rats in the IR group than in those of the rats in the IR+T group ($p = 0.001$; Figure 1). Likewise, MPO activity, a novel indicator for neutrophil function, was significantly higher in the IR group than in the IR+T group ($p = 0.001$; Figure 2).

Figure 3 illustrates a representative photomicrograph of lung tissue of the rats in the IR group 24 h after reperfusion. Histological changes in the IR group included intra-alveolar edema, intra-alveolar hemorrhage, and neutrophil infiltration. The mean level of lung injury in the IR group was 2.10 ± 0.89 . These pathological changes, particularly neutrophil infiltration, were much less common in the IR+T group (Figure 4). One animal in the IR+T group presented no injury, whereas the level of lung injury in the other animals ranged from 1 to 2 (mean, 1.70

± 0.23). Histopathologically, there was a significant difference between two groups ($p = 0.035$).

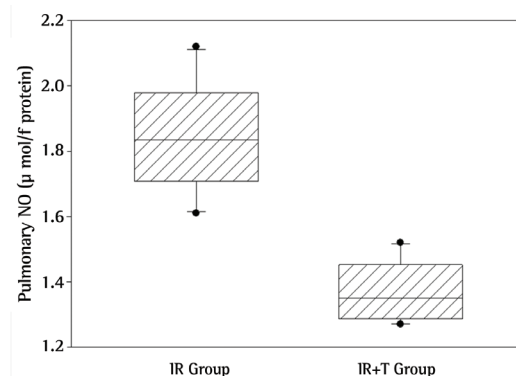


Figure 1 – Levels of nitric oxide (NO) in lung tissue between the groups studied ($p = 0.001$).

IR: ischemia-reperfusion; and IR+T: ischemia-reperfusion + tramadol.

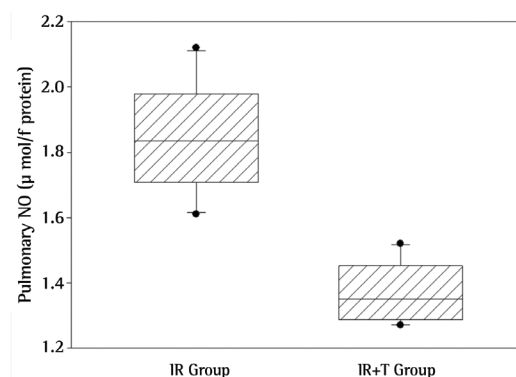


Figure 2 – Myeloperoxidase (MPO) activity in lung tissue between the groups studied ($p = 0.001$).

IR: ischemia-reperfusion; and IR+T: ischemia-reperfusion + tramadol.

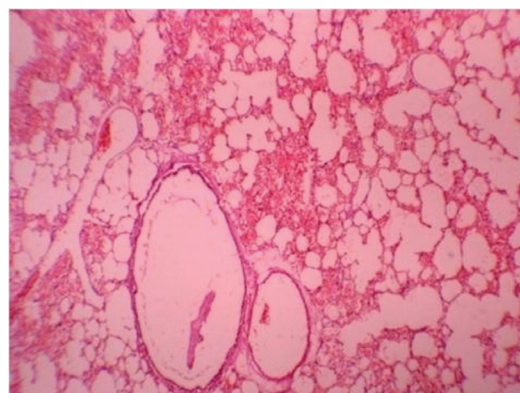


Figure 3 – Photomicrograph under light microscopy. Lung tissue of a rat in the ischemia-reperfusion group showing extensive intra-alveolar hemorrhage (H&E; magnification, $\times 40$).

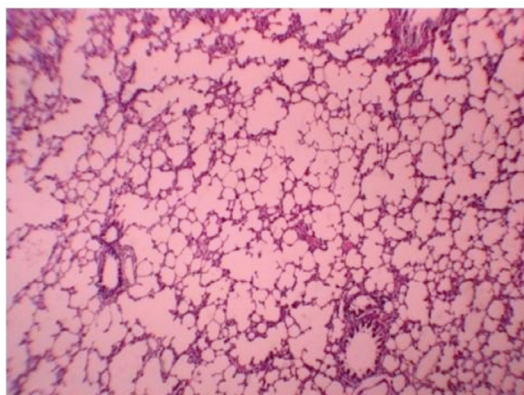


Figure 4 – Photomicrograph under light microscopy. Lung tissue of a rat in the ischemia-reperfusion + tramadol group showing fewer histological changes and better preserved, practically normal structures (H&E; magnification, $\times 40$).

Discussion

The lung is one of the most important target organs in multiple organ dysfunction syndrome or multiple system organ failure caused by severe injury. Lungs can be damaged by indirect injuries caused by the intestine, liver, and skeletal muscle reperfusion, as well as by circulatory shock.^(15,16) The mechanism of respiratory failure after ischemia-reperfusion injury is a complex process which is associated with the activation of systemic inflammatory mediators, including bacteriotoxins, immunocytokines, and inflammatory mediators, such as TNF and interleukins.^(17,18) TNF and NO are significant determinants of the lung injury process, which is caused by lower extremity ischemia-reperfusion,^(19,20) whereas MPO is an index for the accumulation of activated leukocytes in tissues and is associated with an overproduction of reactive oxygen species (ROS); therefore, leukocyte accumulation, high MPO activity, and excessive ROS production exist together in the inflammatory process. Overproduction of ROS results in a quick depletion of antioxidant capacity of the body, which consequently leads to the damage of target organs.^(21,22)

Animal studies showed that opioids can act as a trigger for both phases of ischemic preconditioning,⁽²³⁾ and serotonin augments⁽²⁴⁾ or attenuates⁽²⁵⁾ this phenomenon depending on the concentration. Mayfield et al.⁽⁶⁾ and Chien et al.⁽⁵⁾ have demonstrated that the opioid pathway is involved in tissue preservation during

hypoxia or ischemia. It has been proven that morphine has cardioprotective effects during ischemia-reperfusion.^(26,27) Factors such as respiratory depression and histamine release are the disadvantages of morphine usage during the postoperative period.⁽²⁸⁾ Tramadol is a centrally acting analgesic drug with negligible respiratory depressant action, very low tolerance, and physical dependence liability. The use of tramadol (10 and 20 mg/kg) showed a protective effect against transient forebrain ischemia in rats.⁽¹⁰⁾ In the present study, we tested the hypothesis that 20 mg/kg of tramadol could protect the lungs from remote organ injury after skeletal muscle ischemia-reperfusion. Higher doses of tramadol should be investigated in order to determine whether higher doses would have a higher protective effect.

The present study, in concert with previous ones,⁽²⁹⁻³¹⁾ confirmed that lower limb ischemia-reperfusion could induce acute lung injury in rats. We demonstrated that the acute lung injury induced by lower limb ischemia-reperfusion could be mitigated by tramadol. Our data demonstrate that tramadol significantly decreases the severity of acute lung injury, the infiltration of macrophages and polymorphonuclear leukocytes in the lungs, pulmonary vascular permeability, and intra-alveolar hemorrhage, as well as inhibiting cellular apoptosis in the lungs after skeletal muscle ischemia-reperfusion injury. These results suggest the possibility of clinical application of tramadol in ischemia-reperfusion injury of the lung. Different dosages, alternative time protocols, and forms of tramadol administration for lung injury induced by skeletal muscle ischemia-reperfusion should be investigated in future studies.

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