Original Article

A Comparative Pathomorphological Findings Between Leiurus abdullahbayrami and Androctonus crassicauda (Scorpion: Buthidae) Envenomation in Rabbit Animal Model

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Abstract

Background: The aim of study was to compare macroscopical and histopathological findings between venoms belonging to two scorpion species, Androctonus crassicauda, and the newly discovered Leiurus abdullahbayrami.

Methods: The animals used in this experimental study were fifteen New Zealand bred rabbits. Three groups were constituted as group I (L. abdullahbayrami group, n= 6), group II (A. crassicauda group, n= 6) and group III (control group, n= 3). The animals in the L. abdullahbayrami group and the A. crassicauda group were envenomed through an intravenous route. The rabbits were monitored for the first 24h following the envenomation. The animals dead within that time period were examined and all animals were sacrificed and standard necropsy process was performed at 24h.

Results: The pathomorphological findings from group I were found to be more severe than those observed in group II. The venom from the newly identified L. abdullahbayrami has a greater effect than the venom from the A. crassicauda. Moreover, as this was a rabbit modeling study, the L. abdullahbayrami might pose the most serious health threat to infants in particular due to their smaller body weight.

Conclusion: These findings will provide a better understanding of envenomation of human beings in terms of the possible consequences of scorpion toxication on the organs.

Keywords: Leiurus abdullahbayrami; Androctonus crassicauda; Venom; Pathomorphology; Rabbit

Introduction

In many parts of the world, millions of people are stung by various scorpion species yearly. These stings can result in death, particularly in children (1, 2). This is because of the serious health problems that result, such as cardiovascular, respiratory, and/or neurologic complications. Nowadays, despite advances in medical science, scorpion envenomation cases still continue to be a current public health problem all over the world, including in Turkey. Scorpions are considered to be life-threatening venomous animals. Arguably, the medical knowledge of the scorpion's species is most critical for the scorpion species that are typically found in Mexico, the middle and northern regions of South America, North and South Africa, India, and the Middle Eastern countries because those areas have scorpion species with the most potent venoms as they are classified as neurotoxic, hemotoxic, cardiotoxic, nephrotoxic, and myotoxic (3-5).

In Turkey, the most hazardous scorpions are the yellow scorpion, Leiurus abdullahbayrami, which is endemic in Southeastern Anatolia, and a black scorpion, Androctonus crassicauda, commonly found in Southeastern Anatolia and in a part of Eastern Anatolia (Fig. 1) and neighbouring countries Iran, Iraq and Syria (2, 5-7).

Furthermore, in these regions, the species are responsible for the most deadly cases, particularly those involving children. According-
ly, these scorpion species are described as medically significant to both the world and to Turkey (1, 3, 5, 7).

In the toxicokinetic studies, radiolabeled venom from the *L. quinquestriatus* and venom from the *A. crassicauda* scorpion have reached the maximum value of total plasma radioactivity in less than two minutes after a subcutaneous injection and begin distribution within four to seven minutes after the injection with a half-life of between 4–21h. At that time, the degree of the clinical symptoms depends on the amount of venom circulating in the body (7-9).

A low venom dose triggers an adrenergic effect while high scorpion venom concentrations result in venom cholinergic symptoms. The venom results in an excessive acetylcholine (Ach) release or a decrease in the destruction of Ach, which is acting on the postganglionic nerve endings. Thus, cholinergic activity begins. On the other side, catecholamine is released into the peripheral sympathetic nerve endings from the adrenal medulla in response to the venom. Then, an autonomic storm begins because of the renin secretion by the alpha adrenergic receptor. The receptor stimulation plays a significant role because it increases blood pressure and ultimately results in the pathogenesis of pulmonary edema. Additionally, it causes an inflammatory reaction in the vital organs (8, 10-13).

Scorpion envenomation has been reported to mainly culminate in a syndrome of fuel-energy deficiency. This situation develops as a deficiency during the usage of existing metabolic substrates. As an aftermath, a failure occurs within vital organs including a multi-organ system deficiency that can lead to death (14, 15). The main reason for this failure is the triggering of an inflammatory response cascade that results in a release of several mediators, such as prostaglandins, cytokines, and nitric oxide, and an infiltration of inflammatory cells (16-19). The highest venom concentrations can be found in the kidneys, liver, heart, and lungs following a venom injection (20-22). In envenomations from different scorpion species, the lungs, heart, liver, intestines, and pancreas were documented as the most affected organs in addition to the recently described effects on the brain following a *Leiurus* envenomation (23-27).

As a result of the effect of scorpion venom, different clinical tables have reported varying effects from localized signs to more serious autonomic and neurologic findings as well as fatal cases due to multisystem organ failure. According to medical records from Turkey, most deaths were caused by the results of cardiopulmonary complications, such as myocarditis and acute pulmonary edema following a scorpion’s sting (1, 2, 7).

In animal models, scorpion envenomation symptoms were similar to those described by the clinical findings in humans. Therefore, to mimic human victims, especially children, rabbits were chosen as the animal model for experimental scorpion envenomation to mimic a natural sting accident. Thus, in this study, different pathomorphological findings developed by two scorpion venoms were revealed out *L. abdullahbayrami* and *A. crassicauda*, which species were most encountered in southeast region of Turkey and also were showed *L. abdullahbayrami* having unusual harmful effects onto several organs during long term.

### Materials and Methods

#### Scorpions Origin

Scorpions were collected using an ultraviolet lamp at night in the Sanliurfa and Gaziantep Provinces, southeastern part of Turkey. The animals were kept in plastic boxes at the Department of Entomology, Faculty of Veterinary Medicine, Ankara. The scorpions were received fresh water daily and fed crickets or cockroaches weekly.
Scorpion venom and the median lethal dose ($LD_{50}$)

The venom of scorpions was milked by electrical stimulation of 24 Volt. The venoms were added sterile double-distilled water, dissolved and centrifuged at 14000rpm for 15 min at 4 °C. Supernatant was dissolved in PSS. The median lethal doses ($LD_{50}$) of the scorpions were determined as in previous studies (28, 29).

Animals

This study was conducted on 15 of healthy New Zealand rabbits 12 months old of both sexes and between 2.4±0.1kg body weight for this experiment. The rabbits were fed with a special rabbit pellet diet ad libitum until envenomation. The animals were kept in room temperature set to 19±1 °C and with 12h light/12h dark schedule. The experimental protocol was approved by the local Laboratory Animal Ethics Committee. After 24h all animals were subjected to euthanasia for macroscopical and histopathological findings.

Experimental envenomation of the animals

The animals were envenomed by the venom of $L.\ abdullahbayrami$ in group I (GI, n: 6), and the venom of $A.\ crassicauda$ in group II (GII, n: 6). Finally, three rabbits were selected as a control group and will be referred to as GIII. The venoms ($2\times LD_{50}$ for both) were injected into the marginal ear vein by intravenous (IV) route after dissolving the venoms within 0.5mL PSS. For control group, any envenomation was not performed. 0.9% Physiological saline solution only was administrated to rabbit of this group. The animals were monitored and kept in individual cages up to 24h following the injections because the deaths were not shown during first 24h of critical period.

Histopathological Examination

After providing sedation by xylazine (10 mg/kg Intramuscular route) and ketamine hydrochloride (5mg/kg intramuscular route), Sodium Pentobarbital injection (100mg/kg Intravenous route) were done and then cervical dislocation were performed for each one. Necropsy process was done systematically and all organs and tissues were examined routinely. Lesions were photographed. For histopathological examination, tissue samples were taken systematically from each organ in 10% buffered formalin. After fixation for 48h, tissue samples were processed routinely through alcohol and xylol series and mounted in paraffin. Processed tissues were embedded in paraffin wax. Five-micron thickness-sections were cut from paraffin blocks and Haematoxylin-Eosin (H and E) staining method.

Results

Macroscopical Findings

In all groups, cadavers were well-fed, and rigor mortis did not happen. However, the blood in group I was lately clotted in contrast to the other groups. Conjunctiva was hyperemic in two of the animals from group I. The gut walls were thickened and filled with a yellowish content in all of the members of group I and II. The liver, kidneys, and lungs were congested in all members of both groups I and II. However, there was haemorrhagia that happened in the kidneys in the members of group II, and also atrophia and capsular contraction in the spleen was noticed in all animals within group II. In group I, a meningeal vessel was congested in one animal. In contrast, none of these macroscopical findings were evident in the animals of group III.

Histopathological Findings

Heart

Myocardioocytes in some areas were degenerated or partly necrotic in the members of group I. There was a hyalinosis in appearance in the degenerated myocardioocytes in the group I. Haemorrhagia between cardiomyocytes was formed in two animals from group I and four
animals from group II. However, there was focal mononuclear cell infiltration in one animal from group II. There were no changes in control group.

**Lungs**

All veins and vessels were hyperemic in members from group I. Furthermore, there was edema in all of the animals in group I. Microhaemorrhages were noticed in three of the members of group I and all of the members of group II. BALT hyperplasia was noted in one member of group I and all of the members of group II. It was not encountered with any findings in control group.

**Kidneys**

In the kidneys, there was hyperemia in all vessels and glomerules in all animals from group I, there were no such findings in the group II animals. On the other hand, vacuolar and hydropic degeneration in tubular epithelium were common findings for members of the first group except for one of group I animals. There is no any changes in control group.

**Liver**

Hepatocytes contained vacuoles in different sizes in their cytoplasm and degenerated nuclei in especially the periphery of lobules for all animals from the first two groups. In some animals of group II, reticular degeneration was also noted in addition to those findings. Additionally, mononuclear cell infiltrations in the portal field were evident in the group I findings and in all animals of group II. Finding in control group in terms of this organs were unremarkable.

**Spleen**

Although no prominent findings were observed in animals of group II, focal follicular necrosis and haemorrhagia were noted in each animal of group I. Finding in control group in terms of this organ were unremarkable.

**Central Nervous System**

Lesions on the central nervous system were predominantly seen in group I. Meningeal and parenchymal vessels were severely hyperemic in one animal of group I. Stratum pyramidal cells of the brain often seemed to have lost their nuclear chromatin in almost all foci. Perivascular mononuclear cell infiltration was observed in substantia alba of the cerebrum within one of the animals in group I. In the cerebellum, cytoplasms and nuclei of Purkinje cells generally degenerated in both groups of animals. Moreover, some of the cells were necrotic in group I animals, seen in Fig. 2 and Fig. 3A–F. However, there were no conspicuous findings in the control group in terms of any of these organ.

**Gastrointestinal system**

Regarding the intestinal lesions, in group I, lymphocyte-plasma cell infiltration was noted in three animals, and neutrophil leucocyte infiltration was evident in one animal in the duodenum. In the group II, lymphocyte and macrophage infiltrations were seen in the jejunum as well as the duodenum in two of the animals. In addition, aggregate lymphoid follicles were hyperplasic in three animals from group I. There were no findings in the stomach of any of the members of group I in contrast to the focal periarteriolar haemorrhagia noted in one animal from group II. In control group, there were no any pathologies.
Fig. 1. The distribution of the black colored scorpion is the *Androctonus crassicauda* (black dots), and a yellow colored scorpion the *Leiurus abdullahbayrami* species (red dots) are found in Hatay, Kilis, Sanliurfa, Mardin, Gaziantep provinces of the Southeastern Anatolia as shown map of Turkey. The scorpions were captured in Gaziantep and Sanliurfa.

Fig. 2. A. Vacuolar degeneration (arrows), periphery of lobules, liver, x100, H and E. B. Hyperemia (asterisks), glomerules and capillary vessels, kidney, x100, H and E. C. Vacuolar degeneration (arrows), kidney, x400, H and E. D. Haemorrhagia (arrows), lung, x100, H and E. E. Necrosis in myocardiocytes (arrows), x100, H and E. F. Degeneration (arrows) and necrosis (arrowhead) in Purkinje cells x100, H and E
Discussion

Scorpion envenomation continues to be a global problem despite some national measures taken by authorities in various countries. Regarding the clinical symptoms, the degree of the envenomation depends on some factors related to the scorpion such as the species and the size of the scorpion as well as the number of stings, the content of the venom, and the amount of venom injected. Another factor associated with a patient’s prognosis include the patient’s health status, the part of the body that has been stung, the patient’s body mass, and the patient’s age. After the sting, the symptoms of envenomation can be evident within a few minutes, and maximum severity is usually achieved within five hours. Consequently, the clinical table has shown localized symptoms at the sting site up to a significantly severe generalized envenomation (7, 11, 28).

In Turkey, the scorpion sting is very common in the provinces of Southeastern and Eastern Anatolia due to social and geographical factors. In these areas, the scorpion sting cases have been reported as coming from both the yellow colored scorpion, which is the L. abdullahbayrami, and a black colored scorpion, which is the A. crassicauda species. In addition, scorpions have been the main animal responsible for medically important envenomation cases (5, 11-13).

In the Kilis Province, southeastern Turkey, recently, acute pulmonary edema and cardio
myopathy were due to the excessive catecholamine release that resulted from a sympathetic overstimulation in an 11-yr-old male child following an *L. abdullahbayrami* stung (7). Another study conducted at the Sanliurfa Children’s Hospital reported on a patient with symptoms that included fever, hypersalivation, mydriasis, tachycardia, lethargic, and respiratory distress during admission time. Later, this patient had to be intubated because of pulmonary edema and myocarditis. Another girl had neurological signs similar to the first victim as well as anisocoria and seizure. Two girls, ages 2-yr-old and 4-yr-old, had pulseless ventricular tachycardia and died (13). In the province of Hatay in the Southeast Anatolia region, the patients were categorized according to the color of the scorpion that had stung them as yellow (54.2%) and black (28.7%). Two children passed away due to cardiac and respiratory complications resulting from a scorpion sting (30). In this same province, aPE older boy died because of pulmonary edema (31). Furthermore, in Adiyaman Province, the patients were categorized by the color of the scorpion as well with black consisting of 40% of the cases and yellow resulting in 60% of the cases (12). One of these patients died due to cardiac and respiratory arrest. 50.8% of their scorpion cases were *A. crassicauda* stings (5). Additionally, four of the deaths in children were caused by the *A. crassicauda* species, and only one was caused by an *L. abdullahbayrami* (32). Especially in children, similar findings have been shown in previous studies concerning these two important scorpion species. These scorpions were responsible for serious medical cases and deaths in Turkey as well. Therefore, our study aimed to determine and compare the pathological effects of the venom based on an animal model.

The *L. abdullahbayrami* scorpion venom-induced potassium and sodium voltage-gated ion channels and then led to excessive catecholamine release due to the over stimulation of the sodium and potassium ion channels. As a result of the catecholamines storm, cardiomyopathy and acute pulmonary edema were caused by severe envenomation, especially among small children or infants because of their smaller body weight (7). Namely, following envenomation, an acute failure leads to multi-organ system deficiency among the vital organs, which leads to death (14, 15), especially within the heart and lungs. After acute envenomation, a cardiac dysfunction occurred, and as a result of this cardiac failure, the formation of edema and hemorrhages happened in the lungs (8, 9, 33). Thrombosis of cardiac vessels, endothelial cells of those vessels, edema, and necrosis in cardiac muscle cells were all observed during the 24h following envenomation (33). In another acute envenomation study with *A. australis* hector, LD₅₀ of the venom resulted in myocardial necrosis and degeneration. Neutrophil leucocyte and mononuclear cell infiltration in the interstitium of myocardium were noted within the first 24h period in mice as well. Suppurative bronchopneumonia, focal haemorrhagia, and fibrin deposits are found in the lungs of the mice. Cardiac edema, myocardial haemorrhagia, degeneration and necrosis were also detected with envenomation attributed to *Centruroides sculpturatus* venom (34). In another study that experimented with LD₅₀ doses of *Mesobuthus eupeus* envenomation on rabbits, several important consequences progressed including myocardial necrosis, focal haemorrhage, thrombus formation and inflammation in myocardium and the endocardium surfaces of the heart and lungs (35). In our study, more serious fatal findings, such as hyperemia, edema, and haemorrhage, were evident in the lungs. Degeneration and necrosis in the myocardiocytes occurred within the rabbits in group I envenomated with *L. abdullahbayrami*. However, BALT hyperplasia and microhemorrhage were encountered in the lungs and focal mononuclear cell infiltration in heart in the rabbits envenomated with *A. crassicauda*. *Leirus abdullahbayrami* triggered a more acute reaction especially in regards to the cardiopulmonary system within the
first 24h as compared to another type of venoms documented in several previous studies. Therefore, the results obtained from these study groups were found to parallel the findings of previous studies which also examined the effects that occur within the first 24h following a scorpion sting.

Some deaths in humans and animals were related to the effect of the venom on the kidneys. The scorpion toxin caused hyperemia and degenerative-necrotic changes in the tubules and glomeruli. This situation resulted in acute renal failure (2, 33, 34, 36, 37). Even an envenomation attributed to a *Hemiscorpius lepturus* scorpion can sometimes result in a uremic syndrome in a child (38). In an acute envenomation from *M. eupeus*, congestion in the vessels and glomeruli of the envenomated animals has been reported. In our study, group I animals envenomated with *L. abdullahbayrami* showed similar important damages, including vascular congestion and degenerative changes in tubules, as compared to group II animals envenomated by the venom of the *A. crassicauda*.

On the other hand, certain studies conducted in association with particularly both acute as well as sub-acute envenomation have proven that venom of the *M. eupeus* species can promptly effect animals within 30min up to three to five hours later by resulting in congestion within the central and portal vein of the liver and in interstitial vessels (35). In sub-acute envenomation caused by the *C. sculpturatus's* venom, the liver of rats was affected within two hours and up to five days after the envenomation. In the livers of envenomated animals, congestion of central veins, hydropic degeneration, and single cell necrosis of hepatocytes was also observed (34, 38). In our study, the findings were observed in both experimental groups. However, mononuclear cell infiltrations in the portal field were also noted in all of the animals of group II. The last finding is a different finding as compared to previous reports. Apart from the findings within the vital organs, milder findings such as congestion were encountered in both the spleen and the pancreas (36). In our study, no other prominent findings were evident. However, focal follicular necrosis and haemorrhagia were both noted in all of the animals in group I. The venom of the *L. abdullahbayrami* might be more effective since it triggered an inflammatory reaction.

In relation to the gastrointestinal system, edema and mildly necrosis have occurred in the guts of these envenomed animals. On the other hand, no inflammatory changes were evident even though there were severe microscopical changes noted, such as gastrointestinal haemorrhagia, necrosis, and inflammation (33, 36). In our study, in group I, inflammatory reactions was evident in some of the animals from both groups. Additionally, aggregate lymphoid follicles were found hyperplasic in three animals of group I. There were no findings in the stomach of any of the members from group I, which was in contrast to the focal periarteriolar haemorrhagia found in one of the animals from group II. These findings correlated with the previous findings (36). In our study, the most significant findings in almost all organs were mostly hyperemia, focal haemorrhagia, and mononuclear cell infiltrations as was indicated in several previous studies. Such results have been reported as a result of inflammatory responses under influence of the variety of cytokines as reported in previous reports (18, 24, 25). In our study, the acute findings belonging to *L. abdullahbayrami* envenomation were found to be more severe than the *A. crassicauda* envenomation because it caused much more of an inflammatory reaction within the first 24h. In this sense, the newly identified *L. abdullahbayrami* venom had more of an effect than the *A. crassicauda* venom.

Additionally, many studies have reported on this inflammatory responses following envenomation. Among them, leucocytosis has been mentioned as occurring in the hours following envenomation (14, 18, 25, 39, 40). In the present study, mononuclear cell infiltrations
were more evident in several organs, especially among the *Leirus* sp. group. The *Leirus* envenomation resulted in an ongoing or final stage of acute inflammatory reaction. On the other hand, the main affected organs included the lungs, liver, intestine, and pancreas, respectively (22-26). However, some important findings in the kidneys and the central nervous systems were also evident in our study, which was less likely to be mentioned in prior documents except for the most recent document (27). The inflammation might be a long-term effect from the two types of venom. Particularly, another reason for multiple organ disfai...
Important Scorpions in North West of Iran. J Arthropod Borne Dis. 11(3): 371–382.

4. Pourkhalili K, Kim E, Mashayekhy NR, Kamyab M, Hoseiny SM, Evazy R, Zare Mirakabadi A, Seyedian R (2015) Cardiotoxic and Arrhythmogenic Effects of Hemiscorpius lepturus Scorpion Venom in Rats. J Arthropod Borne Dis. 9 (2): 215–225.

5. Ozkan O, Adiguzel S, Yakiştirar S, Cesaretli Y, Karaer KZ (2006) Androctonus crassicauda (Oliver 1807) scorpionism in Sanliurfa Province region of Turkey. Turkiye Parazitol Derg. 30(3): 239–245.

6. Seiter M, Koc H, Ullrich A, Yagmur EA (2016) The case history of a toxic sting of a Leiurus abdullahbayrami scorpion in Turkey. Arachnologische Mitteilungen. 51: 64–66.

7. Dokur M, Dogan M, Yagmur EA (2017) Scorpion-related cardiomyopathy and acute pulmonary edema in a child who is stung by Leiurus abdullahbayrami. Turk J Emerg Med. 17(3): 104–108.

8. Ismail M, Abdel-Rahman AA, El-Asmar MF (1975) Pharmacological studies of scorpion (Androctonus amoreuxi Aud. and Sav.) venom. Toxicon. 13: 253–259.

9. Ismail M, Abdoulah ME, Morad AM, Ageel AM (1980) Pharmacokinetics of 125I-labelled venom from the scorpion Androctonus amoreuxi (Aud. and Sav.). Toxicon. 18: 301–308.

10. Radmanesh M (1990) Androctonus crassicauda sting and its clinical study in Iran. J Trop Med Hyg. 93(5): 323–326.

11. Adiguzel S (2010) In vivo and in vitro effects of scorpion venoms in Turkey: A mini-review. J Venom Anim Toxins incl Trop Dis. 16(2): 198–211.

12. Konca C, Tekin M, Genc Y, Turgut M (2015) Epidemiological and Clinical Characteristics and Outcomes of Scorpion Envenomation in Hospitalized Chil-

dren in Adiyaman, Turkey. Iran J Pediatr. 25(1): e199.

13. Caglar A, Köse H, Babayigit A, Oner T, Duman M (2015) Predictive factors for determining the clinical severity of pediatric scorpion envenomation cases in Southeastern Turkey. Wilderness Environ Med. 26: 451–458.

14. Bertazzi DT, Assis-Pandochi AI, Azzolini AE, Talhaferro VL, Lazzarini M, Arantes EC (2003) Effect of Tityus serrulatus scorpion venom and its major toxin, TsTX-I, on the complement system in vivo. Toxicon. 41: 501–508.

15. Petreicevich VL (2010) Scorpion venom and the inflammatory response. MEDIATORS INFLAMM. 2010: 1–10.

16. Barbourche MR, Haguiga H, Nouira S, Krifi MN, Abroug F, Bouchoucha S (1996) Inflammatory cytokines and scorpionic analysis of serological levels in 46 Tunisian patients. Toxicon. 34: 156–157.

17. Meki A, Mohey El-Dean ZM (1998) Serum interleukin 1β, interleukin-6, nitric oxide and α1-antitrypsin in scorpion envenomed children. Toxicon. 36: 1851–1859.

18. Magalhaes MM, Pereira ME, Amaral CF, Rezende NA, Campolina D, Bucaretchi F, Gazzinelli RT, Cunha-Melo JR (1999) Serum levels of cytokines in patients envenomed by Tityus serrulatus scorpion sting. Toxicon. 37: 1155–1164.

19. Fabiano G, Pezzolla, A, Filograna MA, Ferrarese F (2008) Traumatic shock-physiopathologic aspects. G Chir. 29 (1-2): 51–57.

20. Krifi MN, Marrakchi N, El Ayeb M, Dellagi K (1998) Effect of some variables on the in vivo determination of scorpion and viper venom toxicities. Biologicals. 26(4): 277–288.

21. Mirakabadi AZ, Jalali A, Jahromi AE, Vatanpour H, Akbary A (2006) Biochemical changes and manifestations of
Venom produced by *Odonthobuthus doriae* venom in rabbits. J Venom Anim Toxins Incl Trop Dis. 12(1): 67–77.

22. Ozkan O, Carhan A (2008) The neutralizing capacity of *Androctonus cassicusida* antivenom against *Mesobuthus eupeus* scorpion venom. Toxicon. 52(2): 375–379.

23. Freire-Maia L, De Matos IM (1993) Heparin or a PAF antagonist (BN-52021) prevents the acute pulmonary edema induced by *Tityus serrulatus* scorpion in the rat. Toxicon. 31: 1207–1220.

24. Novoa E, D'Suze G, Winter M, Crespo A, Tortoledo MA, Marcano H, Friedman E, Sevik C, Zabrer J, Sanchez de Leon R (2003) The indirect effect of *Tityus discrepans* on rabbit pulmonary vasculature. Respir Physiol Neurobiol. 134(1): 33–41.

25. D'Suze G, Salazar V, Diaz P, Sevcik C, Azpurua H, Bracho N (2004) Histopathological changes and inflammatory response induced by *Tityus discrepans* scorpion venom in rams. Toxicon. 44: 851–860.

26. Bakir B, Ozkan O, Alcigir ME, Vural SA (2012) Effects of *Androctonus crassicauda* scorpion venom on the heart tissue. J Anim Vet Adv. 11(14): 2594–2599.

27. Afifi SH, El-Kashef R, Seddek, AS, Salem, DA (2016) Light and transmission electron microscopical changes associated with *Leiurus quinquestratus* venom in rabbits. Mac Vet Rev. 39(1): 51–57.

28. Ozkan O, Yagmur EA, Ark M (2011) A newly described scorpion species, *Leiurus abdullahbayrami* (Scorpion: Buthidae), and the lethal potency and in vivo effects of its venom. J Venom Anim Toxins incl Trop Dis. 17(4): 414–421.

29. Ozkan O, Yagmur EA (2017) Neutralization capacity of monovalent antivenom against existing lethal scorpions in the Turkish Scorpiofauna. Iran J Pharm Res. 16 (2): 653–660.

30. Duru M, Karakuş A, Yengil E, Sahan M, Zeren C, Kekek Z, Yagız AE, Yaman FN (2013) Demographic, clinic and laboratory characteristics of cases presented to emergency department with scorpion sting: 345 cases over a 4-years period. Acta Medica Mediterranea. 29: 611–616.

31. Karakuş A, Arica V, Çeçik T, Tutanç M, Zeren C, Arica ŞG (2013) Death due to scorpion sting: Child case report. J Kartal TR, 24(1): 50–53.

32. Balık S (2010) Güneydoğu Anadolu Bölgesindeki Çocuklarda Akrep Sokma-larının Retrospektif İncelenmesi. Yayınlanmamış yüksek lisans tezi. T.C. Dicle Üniversitesi Tip Fakültesi. p. 55. Available at: http://kutup.dicle.edu.tr/ekitap/0038768.pdf.

33. Heidarpour M, Ennaifer E, Ahari H, Srairi-Abid N, Borchani L, Khalili G, Amini H, Anvar AA, Boubaker S, El-Ayeb M, Shahbazzadeh D (2012) Histopathological changes induced by *Hemiscorpius lepturus* scorpion venom in mice. Toxicon. 59: 373–378.

34. Bertke EM, Atkins JH (1961) Effect of *Centruroides sculpturatus* venom on rat tissue: a histopathologic study. Toxicon. 2: 205–209.

35. Zayerzadeh E, Zare MA, Koohi MK (2011) Biochemical and histopathological study of *Mesobuthus eupeus* scorpion venom in the experimental rabbits. Arch Razi Inst. 66(2): 133-138.

36. Dehghani R, Khamechian T, Vazirianzadeh B, Vatandoost H, Moravvej SA (2012) Toxic effects of scorpion, *Hemiscorpius lepturus* (Hemiscorpiidae) venom on mice. J Anim Plant Sci. 22 (3): 593–596.

37. Valavi E, Ansari MJ (2008) Hemolytic uremic syndrome following *Hemiscorpius lepturus* (scorpion) sting. Indian J

http://jad.tums.ac.ir
Published Online: April 27, 2019
38. Correa MM, Sampaio SV, Lopes RA, Mancuso LC, Cunha OA, Franco JJ, Giglio JR (1997) Biochemical and histopathological alterations induced in rats by Tityus serrulatus scorpion venom and its major neurotoxin tityustoxin-1. Toxicon. 35(7): 1053–1067.

39. Sofer S, Gueron M, White RM, Lifshitz M, Apte RN (1996) Interleukin-6 release following scorpion sting in children. Toxicon. 34: 389–392.

40. Hodhod S, Shoukry S, El-Shimi IR, El-Asmar MF (1977) Effect of scorpion Androctonus amoreuxi (Aud. and Sav) venom on blood and liver of rat. The 1st Ain Shams Medical Congress. 1977, Egypt. 1: 535–560.

41. Ismail M (1995) Review article: the scorpion envenoming syndrome. Toxicon. 33: 825–858.

42. Possani LD, Becerril B, Delpierre M, Tytgat J (1999) Scorpion toxins specific for sodium channels. Eur J Biochem. 264: 287–300.