Evaluation of the Impact of Platelet Rich Plasma and Hyaluronic Acid on Ketamine-Induced Cystitis in Albino Rats: A Histological and Immunohistochemical Study

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ABSTRACT

Background: ketamine induced cystitis (KIC) is commonly occurred in ketamine abusing individuals. It is associated with damage in the lining of the bladder and lower urinary tract symptoms. Platelets rich plasma (PRP) is attractive therapeutic line in regenerative medicine. Hyaluronic acid (HA) has many beneficial therapeutic effects in treatment of different types of cystitis.

Objective: to evaluate the effect of PRP and HA on ketamine induced cystitis in adult male rats.

Materials and Methods: Forty- four adult male rats were divided into five groups: Group I: control rats. Group II: Ketamine group. Group III: Recovery group. Group IV: Ketamine and PRP treated group. Group V: Ketamine and HA treated group. Specimens from the body of the urinary bladder were processed and examined histologically and immunohistochemically. Morphometrical studies and statistical analysis were conducted.

Results: Groups II and III showed focal areas of degeneration and ulceration with apparently decreased thickness of the urothelium. Cellular infiltration and dilated congested blood vessels of the lamina propria were also observed. A significant increase of the collagen fibers in the lamina propria and between the muscle bundles. Nuclear factor kappa B (NF-κB) immunoreaction was significantly increased but there was a non-significant increase of antiproliferating cell nuclear antigen (PCNA) immunopositive nuclei among urothelium and a significant decrease of desmin, compared to group I. Groups III and IV showed improvement of most of the histological and immunohistochemical changes described before.

Conclusion: The intravesical injection of PRP and HA provide a positive impact on treatment of KIC. HA therapy is a more efficient mean as it provides a better improvement in healing of urothelium and promotes more rapid tissue regeneration.

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Key Words: HA, KIC, PRP.

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INTRODUCTION

The urinary bladder is not a vital organ. Its only and main function is storage of urine followed by periodic emptying with time. The bladder urothelium is particularly quiescent and its turnover rate is low about 0.12%[1].

Cystitis is an inflammatory disease which caused by several etiologic factors, such as bacterial infection, chemicals, exposure to radiation, use of a catheter, or irritating hygiene products, that act through many pathogenic mechanisms[2].

Ketamine is a potent non-competitive receptor antagonist of N-methyl-D-aspartate (NMDA). It is an intravenous anesthetic drug with sedative, analgesic, and anti-depressive effects[3,4].

Ketamine abuse has increased, and its hazardous effects have attracted more and more people’s attention[4,5]. It may cause serious health problems to many systems as brain, cardiovascular and genital systems. Additionally, it can damage the urinary tract integrity and function leading to ketamine induced cystitis (KIC). The main clinical manifestations of KIC are similar to interstitial cystitis (IC), which include frequency, dysuria, hematuria and a reduced capacity of the bladder[6,7].

Various regimens for treatment KIC, mainly symptomatic, have been used, including antibiotics, steroids, non-steroid anti-inflammatory drugs, and anticholinergics. Though, none of the them have an effective and durable results[8].

Platelet-rich plasma (PRP) has become attractive therapeutic option in regenerative medicine for its powerful healing properties[9]. Platelet-rich plasma is an autologous derivative of whole blood that concentrates a large number of platelets in a small volume of plasma to be considered “platelet rich”. PRP is a rich source of several growth factors. It formed of different quantities of plasma, white blood cells, erythrocytes and platelets according to the device and technique applied[10,11].

It has major advantages compared with other methods on clinical purposes as it is a cheap product, easily obtained, and being autologous diminishes the hazards of
rats were injected with one mL syringe through the transurethral tube and each was retained in the bladder for 30 min after aspiration. Then, PRP or hyaluronic acid through the urethra. The lower abdomen was pressed gently to help the urine aspiration. PRP were injected intravesical in a dose of 1 ml/rat/day for two weeks.

**MATERIALS AND METHODS**

**Rats** were arranged into five groups

**Group I (control group)** (n=12): rats were subdivided equally into three subgroups. Subgroup Ia: rats were left without any intervention. Subgroup Ib: rats were injected intraperitoneal daily with 0.5 ml of 0.9% saline for four weeks; Subgroup Ic: rats were injected intraperitoneal daily with the 0.5 ml of 0.9% saline, then daily intravesical injection of 0.5 ml saline for another two weeks.

**Group II (Ketamine group)** (n=8): rats were injected intraperitoneal daily with ketamine for four weeks.

**Group III (Recovery group)** (n=8): rats were injected daily with ketamine for four weeks and then left without treatment for another two weeks.

**Group IV (Ketamine and PRP treated group)** (n=8): rats were injected with ketamine as described in group II; then injected daily intravesical with PRP for another two weeks.

**Group V (Ketamine and HA treated group)** (n=8): rats were injected with ketamine as in group II; then injected daily intravesical with HA for another two weeks.

Rats of subgroup Ib, and group II were sacrificed after four weeks, while those of the other groups were sacrificed after six weeks from the beginning of experiment.

**Preparation of Platelet-Rich Plasma (PRP)**

The PRP was prepared using the double centrifugation tube method. Fresh venous blood was collected from the jugular veins of healthy male rats and treated with sodium citrate anticoagulant. The blood containing tubes were centrifuged at 1700 revolutions per minute (rpm) for about ten min. The blood differentiated into three different density components; the upper one contained plasma and platelets, RBCs were at the bottom, while in-between there was a buffy coat of leucocytes. A cellular plasma was pipetted and transferred into new tube which centrifuged again at 2000 rpm for 15 min. Plasma centrifugation resulted into a supernatant of platelet-poor plasma (PPP) and the platelet pellet. Platelet-poor plasma were removed, and the platelet pellets were suspended in 3-5 ml of plasma to produce the PRP.

**Histological Studies**

Specimens from the body of the urinary bladder were fixed in 10% formalin for 48 hours. Paraffin sections (5-μm-thick) were prepared, processed and stained with hematoxylin & eosin for the histological details of urinary bladder and with Masson’s trichrome for collagen.

**Immunohistochemical Studies**

Other urinary bladder sections were mounted on positively charged slides for immunohistochemical staining.
1. Nuclear factor kappa B (NF-κB) is an indicator of inflammation. The primary antibody used was rabbit polyclonal antibody against NF-κB (Labvision, Fremont, California, USA). NF-κB immunoeexpression is brown nuclear/cytoplasmic staining.

2. Antiproliferating cell nuclear antigen (PCNA), is an indicator of cellular proliferation. The primary monoclonal antibody used was anti-PCNA IgG antibody (Sigma-Aldrich Inc., U.K.). PCNA immunoeexpression appeared as brown nuclear staining.

3. Desmin: the primary antibody, monoclonal anti-Desmin (DE-R-11), (Ventana Medical Systems, Tucson, Arizona, USA). Intermediate filaments of desmin were stained brown (cytoplasmic).

Morphometric Analysis
Morphometric studies were performed by means of a Leica Qwin 500 image analysis computer system (Leica Microsystems Ltd., Cambridge, UK) at the Pathology Department, Faculty of Medicine, Cairo University. Eight slides from different rats of each group (n=8) were evaluated. The mean area percentage of collagen fibers as well as the areas occupied by positive immunoeexpression of NF kappa B, PCNA and desmin were assessed in 10 nonoverlapping fields of each section at ×400 magnification.

Statistical Analysis
All the data collected from the experiment was recorded and analyzed using IBM SPSS Statistics software for Windows, Version 20 (IBM Corp., Armonk, NY, USA). One-way analysis of variance (ANOVA) with Post Hoc LSD test was applied to compare differences among the groups. The data was expressed as t mean (M) value ± standard deviation (SD) and differences were considered significant at $P < 0.01$.

RESULTS

H&E Stain

Group 1: Urinary bladder sections from all control subgroups showed normal histological architecture. Its wall was formed of highly folded mucosa, which was lined by transitional urothelium with underlying lamina propria and smooth muscle bundles. Epithelial lining was formed of superficial umbrella-like cells bulging into the lumen with abundant eosinophilic cytoplasm. The intermediate cells were smaller with ample cytoplasm and oval nuclei.

The sections from groups I, II and III showed some PCNA immunopositive nuclei among urothelium.
Group IV revealed moderate PCNA immunopositive nuclear reaction among urothelium (Figure 7d), while, there were many PCNA immunopositive nuclei among urothelium of group V (Figure 7e).

3- Desmin immunostaining

Group I showed intense brown immunoreaction of desmin intermediate filaments inside the cytoplasm of the smooth muscle fibers (Figure 8a), while desmin immunoreaction was mild in groups II and III (Figures 8b & 8c). Group IV revealed moderated brown immunoreaction of desmin intermediate filaments (Figure 8d) while in group V, the immunoreaction of desmin intermediate filaments was strong brown (Figure 8e).

Morphometric Results

Mean area percentage ± SD of Masson’s trichrome, NF-κ B, PCNA and desmin immunostaining are presented in (Tables and Histograms 1,2,3 and 4). The mean area percentage of Masson’s trichrome staining of collagen fibers and NF-κ B immunoeexpression were significantly increased (P<0.01) in groups II and III compared with group I, while in groups IV and V there was a significant reduction (P<0.01) compared with groups II and III. The mean area percentage of PCNA immunostaining was non significantly increased (P<0.01) in groups II and III compared with group I. Groups IV and V showed significant increase (P<0.01) of PCNA immunostaining compared with groups I&II and III. Mean area percentage of desmin immunostaining was significantly decreased (P<0.01) in groups II and III in comparison to group I. Groups IV and V showed significant increase of desmin immunostaining (P<0.01) compared with groups II and III.
Fig. 2(b): Group II showing vacuolated urothelial cells (↑) with darkly stained nuclei (▲) and karyolitic ones (*). Disrupted lamina propria (LP) containing dilated congested blood vessel (V) is observed (H&E, ×630).

Fig. 2(c): Group III showing an apparently decreased thickness of the urothelium (U) with focal shedding of some cells (↑↑). The lamina propria (LP) shows wide spaces (S), inflammatory cell infiltration (I) and areas of hemorrhage (H) (H&E, ×630).

Fig. 2(d): Group II showing bundles of smooth muscle (M). Some of them are degenerated leaving wide spaces (S). Wavy collagen fibers (W) and dilated congested blood vessels (V) are present in between the bundles (H&E, ×630).

Fig. 2(e): Group III showing degenerated bundles of smooth muscle (M). Note the presence of wide spaces (S) and dilated congested blood vessels (V) in between the muscle bundles (H&E, ×630).

Fig. 3(a): A photomicrograph of a section in the urinary bladder of group IV showing intact urothelium (U). Disrupted lamina propria (LP), inflammatory cell infiltration (I) and dilated congested blood vessels (V) are observed (H&E, ×630).

Fig. 3(b): Showing bundles of smooth muscle (M). Some of them are still degenerated leaving wide spaces (S) (H&E, ×630).
Fig. 4(a): A photomicrograph of a section in the urinary bladder of group V showing apparently normal urothelium (U) and lamina propria (LP) with few inflammatory cell infiltration (I) (H&E, ×630).

Fig. 4(b): Showing more or less normal appearance of the smooth muscle bundles (M) running in different directions (H&E, ×630).

Fig. 5(a): A photomicrograph of a section in the urinary bladder of group I showing normal distribution of the collagen fibers in the lamina propria beneath the epithelium and in between the smooth muscle bundle (↑) (Masson’s trichrome ×400).

Fig. 5(b): Group II showing intense collagen fibers deposition in the lamina propria and in between muscle bundle (↑) (Masson’s trichrome ×400).

Fig. 5(c): Group III showing dense deposition of collagen fibers in the lamina propria and in between muscle bundle (↑) (Masson’s trichrome ×400).

Fig. 5(d): Group IV showing moderate deposition of collagen fibers in the lamina propria and in between muscle bundle (↑) (Masson’s trichrome ×400).
Fig. 5(e): Group V showing deposition of collagen fibers in the lamina propria and in between muscle bundle (↑) (Masson's trichrome ×400).

Fig. 6(a): A photomicrograph of a section in the urinary bladder of group I showing minimal NF-κ B immunoreaction in the urothelium (↑) and lamina propria (▲) (Immunostaining for NF-κ B ×400).

Fig. 6(b): Group II showing strong NF-κ B immunoreaction in the urothelium (↑) and lamina propria (▲) (Immunostaining for NF-κ B ×400).

Fig. 6(c): Group III showing intense NF-κ B immunoreaction in the urothelium (↑) and lamina propria (▲) (Immunostaining for NF-κ B ×400).

Fig. 6(d): Group IV showing moderate NF-κ B immunoreaction in the urothelium (↑) and lamina propria (▲) (Immunostaining for NF-κ B ×400).

Fig. 6(e): Group V showing weak NF-κ B immunoreaction in both the urothelium (↑) and the lamina propria (▲) (Immunostaining for NF-κ B ×400).
Fig. 7(a): A photomicrograph of a section in the urinary bladder of group I showing some PCNA immunopositive nuclei (↑) among urothelium (Immunostaining for PCNA ×400).

Fig. 7(b): Group II showing some PCNA immunopositive nuclei (↑) among urothelium (Immunostaining for PCNA ×400).

Fig. 7(c): Group III showing some PCNA immunopositive nuclei (↑) among urothelium (Immunostaining for PCNA ×400).

Fig. 7(d): Group IV showing many PCNA immunopositive nuclei (↑) among urothelium (Immunostaining for PCNA ×400).

Fig. 7(e): Group V showing many PCNA immunopositive nuclei (↑) among urothelium (Immunostaining for PCNA ×400).

Fig. 8(a): A photomicrograph of a section in the urinary bladder of group I showing intense brown immunoreaction of desmin intermediate filaments (↑) inside the cytoplasm of the smooth muscle fibers (Immunostaining for desmin ×400).
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**Fig. 8(b):** Group II showing mild brown immunoreaction of desmin intermediate filaments (↑) (Immunostaining for desmin ×400).

**Fig. 8(c):** Group III showing mild brown immunoreaction of desmin intermediate filaments (↑) (Immunostaining for desmin ×400).

**Fig. 8(d):** Group IV showing moderated brown immunoreaction in some of desmin intermediate filaments (↑) and intense in others (↑↑). Also, some desmin filaments show weak reaction (▲). (Immunostaining for desmin ×400).

**Fig. 8(e):** Group V showing strong brown immunoreaction in the most of desmin intermediate filaments (↑) (Immunostaining for desmin ×400).

**Table 1:** Comparison of collagen fibers deposition in all groups

<table>
<thead>
<tr>
<th></th>
<th>Group I</th>
<th>Group II</th>
<th>Group III</th>
<th>Group IV</th>
<th>Group V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>12.87%</td>
<td>23.56%</td>
<td>22.91%</td>
<td>17.09%</td>
<td>14.76%</td>
</tr>
<tr>
<td>SD</td>
<td>0.7000</td>
<td>1.1334</td>
<td>0.9961</td>
<td>0.8871</td>
<td>0.7568</td>
</tr>
<tr>
<td>Significance</td>
<td>2,3,4,5</td>
<td>1,4,5</td>
<td>1,4,5</td>
<td>1,2,3,5</td>
<td>1,2,3,4</td>
</tr>
</tbody>
</table>

1=sig. with group I     2=sig. with group II   3=sig. with group III
4=sig. with group IV                                                   5=sig. with group V.

**Table 2:** Comparison of NF-κB immunostaining in all groups

<table>
<thead>
<tr>
<th></th>
<th>Group I</th>
<th>Group II</th>
<th>Group III</th>
<th>Group IV</th>
<th>Group V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>1.19%</td>
<td>5.61%</td>
<td>5.96%</td>
<td>3.47%</td>
<td>1.65%</td>
</tr>
<tr>
<td>SD</td>
<td>0.4075</td>
<td>0.6984</td>
<td>0.4059</td>
<td>0.6803</td>
<td>0.4002</td>
</tr>
<tr>
<td>Significance</td>
<td>2,3,4</td>
<td>1,4,5</td>
<td>1,4,5</td>
<td>1,2,3,5</td>
<td>2,3,4</td>
</tr>
</tbody>
</table>

1=sig. with group I     2=sig. with group II   3=sig. with group III
4=sig. with group IV                                                   5=sig. with group V.

**Table 3:** Comparison of PCNA immunostaining in all groups

<table>
<thead>
<tr>
<th></th>
<th>Group I</th>
<th>Group II</th>
<th>Group III</th>
<th>Group IV</th>
<th>Group V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>0.76%</td>
<td>0.79%</td>
<td>0.70%</td>
<td>1.79%</td>
<td>2.80%</td>
</tr>
<tr>
<td>SD</td>
<td>0.0308</td>
<td>0.02312</td>
<td>0.0595</td>
<td>0.1455</td>
<td>0.1095</td>
</tr>
<tr>
<td>Significance</td>
<td>4,5</td>
<td>4,5</td>
<td>4,5</td>
<td>1,2,3,5</td>
<td>1,2,3,4</td>
</tr>
</tbody>
</table>

1=sig. with group I     2=sig. with group II   3=sig. with group III
4=sig. with group IV                                                   5=sig. with group V.
Table 4: Comparison of desmin immunostaining in all groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean</th>
<th>SD</th>
<th>Significance at P &lt; 0.01</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>26.66%</td>
<td>1.1720</td>
<td>2,3,4</td>
</tr>
<tr>
<td>II</td>
<td>6.94%</td>
<td>1.2928</td>
<td>1,4,5</td>
</tr>
<tr>
<td>III</td>
<td>7.87%</td>
<td>1.6892</td>
<td>1,4,5</td>
</tr>
<tr>
<td>IV</td>
<td>19.25%</td>
<td>1.5084</td>
<td>1,2,3,5</td>
</tr>
<tr>
<td>V</td>
<td>27.64%</td>
<td>1.3477</td>
<td>2,3,4</td>
</tr>
</tbody>
</table>

1=sig. with group I  2=sig. with group II  3=sig. with group III  4=sig. with group IV  5=sig. with group V

DISCUSSION

Ketamine-induced cystitis is a popular disease associated with ketamine administration[22]. Its clinical manifestations include various lower urinary tract symptoms which are identical with that of interstitial cystitis/bladder pain syndrome[18].

The light and immunohistochemical examination of sections from groups II & III showed the same results. They revealed several changes such as focal areas of degeneration and ulceration with apparently decreased thickness of the transitional epithelium. The lamina propria showed inflammatory cell infiltration and dilated congested blood vessels. The smooth muscle bundles were degenerated leaving wide spaces. There was a significant (P < 0.01) increase of collagen fibers deposition and NF-κB immunoreaction, but non-significant increase of PCNA immunopositive nuclear reaction among urothelium. Desmin immunoreaction was significantly decreased (P<0.01), compared to group I.

Our results were in accordance with previous studies[6,7,27,28] who reported that ketamine treatment was associated with a significant reduction in the urothelial thickness, hemorrhage, inflammatory cells infiltration, and interstitial fibrosis of bladder. Other investigators[5,29] have clarified that, in animal models exposed to ketamine, the permeability of the urothelium was increased, which could explain the damage, vacuolization of cells, and widening of the intercellular spaces beyond the superficial layer of urothelium.

The inflammatory mononuclear cell infiltration could be explained by increasing many cytokines as tumor necrosis factor (TNF-α), interleukin (IL-6) and inducible nitric oxide synthase (iNOS) after ketamine administration which latter on stimulating apoptosis and fibrosis[4,5,30].

Nuclear factor-κ B is a member of the protein family which exists in the cytoplasm as an inactive complex bound to inhibitory proteins. Activation of NF-κ B with KIC induces various proinflammatory cascade, as cytokines (TNF-α, IL-1, IL-6, IL-8, and IL-12), chemokines, and thus plays important roles in the immune and inflammation processes[31-34]. The current results agreed with a study[35].
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done previously and reported that the significant increase of NF-κB in both the urothelium and the lamina propria induce bladder toxicity and has vital role in the pathogenesis of cystitis in rats.

Proliferating cell nuclear antigen (PCNA) is a non-histone nuclear protein that used as a standard marker for cellular proliferation activity[98]. Our findings were coincided with previous researches[12,23,37], who stated that urothelium is a very stable type of tissue and has a slow proliferative activity. However, after exposure to an injury, which causes loss of superficial cells, tight junctions are damaged, and the proliferative activity of the basal urothelial cells are increased rapidly.

Desmin is the predominant intermediate filament protein in the visceral smooth muscle[98]. The current work showed marked degeneration of the smooth muscle bundles in the sections of groups II&III. Our results were identical with the findings of prior researches[23,29,38,48] who stated that ketamine can cause damage in the mucosal, submucosal, and muscular layers of the bladder. Also, they suggested that the degeneration and wide spaces between the muscle bundles, are a result of the dilatation in the lymphatic vessels, which are confined mainly in the muscle layer.

Several reporters[4,5,41,42] explained that the most possible mechanism of KIC is mediated through multifactorial process such as metabolization of ketamine by cytochrome p450 in the liver into two principal metabolites (nor ketamine and di-hydronorketamine), which are excreted primarily in urine and to a much lesser extent in the feces. Ketamine and its metabolites accumulate in the urine and induce disruption of the urothelial barrier, bladder irritation and damage. Moreover, ketamine increases the oxidative stress products (ROS), and reactive nitrogen species (RNS) which induces oxidative stress, and in turn triggers cellular changes such as DNA damage, apoptosis, cytokine production and finally inflammation[7,22,41,42].

Group IV of current research revealed some improvement of the histological changes noticed in groups II&III. Moreover, there was a significant (P<0.01) decrease in the collagen fibers deposition. Proliferating cell nuclear antigen immunopositive nuclei and Desmin immunoreaction were significantly (P<0.01) increased, while, NF-κB immunoreaction was significantly decreased compared to group I.

Intravesical administration of PRP was chosen in our study because it can provide more intensive use of the PRP in the bladder and reduce its systemic adverse effects[59].

Our observation confirms the findings of other studies[1,43] suggesting that intravesical instillation of PRP enhances healing of bladder mucosal wound, and repair the defective urothelium. Though direct contact of PRP with the damaged urothelium can initiate cell proliferations above basement membrane and thus protect the urothelium.

Other researches[11,12,44] attributed the ability of PRP to accelerate the epithelial cell regeneration to the biological effects of many growth factors and different cytokines as platelet derived growth factor, vascular endothelial growth factor and transforming growth factor of platelets. These growth factor leads to cell proliferation of epithelial cells, promote tissue angiogenesis, increase blood flow, and improve oxygenation in the site of injury. Proliferating cell nuclear antigen improve inflammation through increasing the intracellular expression of the anti-inflammatory mediators (IL-4, IL-10, and IL-13) and inhibition of NF-κB[9,45,46]. Furthermore, the macrophages and neutrophils recruited by the platelets may also have vital actions in elimination of the inflammation and infection[1].

Different researches[8,11,13,44] found that PRP has an antifibrotic effect as it contains antifibrotic molecules, growth factors, and serum amyloid protein which could ameliorate the fibrosis. Additionally, PRP treatment increases muscle regeneration by stimulating myogenesis[47,48]. Also, it had antiapoptotic effect by downregulating the expression of apoptotic genes as DAPK1 and BIM mRNA and inhibiting p53, Bax, and caspase-3 levels[90].

Group V of the present study showed a marked improvement confirmed by the histological, immunohistochemical, and morphometric results.

Preservation of the histological structures with HA were an identity with previous researchers[17,49], who stated that HA facilitates the process of wound healing, reepithelization and hasten the tissue regeneration rate.

Some researches[17,19,21] mentioned that HA has an immune-modulatory effect through reduction of secretion of the proinflammatory cytokines such as TNF-α, IL-1, and IL-6 and finally lead to limit inflammation. Also, previous authors[19] concluded in their studies on urinary bladder that HA preserved antioxidant enzymes, eventually lead to accelerate epithelial healing of the bladder mucosa and inhibits fibrosis.

Collectively, HA therapeutic effects on urothelial cells could be caused by a combination of multiple mechanisms, such as, coating of the uroepithelium and antioxidant activity against cellular oxidative stress. Also, anti-inflammatory, antiapoptotic, anti-fibrotic and antiproliferation effects were the other different mechanisms[17,19,49].

CONCLUSION

The intravesical injection of PRP and HA provide a positive impact on the treatment of Ketamine-induced cystitis. Hyaluronic acid therapy is a more efficient mean as it provides a better improvement in healing of urothelium and promotes more rapid tissue regeneration. However, further clinical studies on human populations for longer periods are still required to emphasize the results obtained from animal studies.
CONFLICTS OF INTEREST

There are no conflicts of interest.

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الملخص العربي

تقييم تأثير البلازما الغنية بالصفائح الدموية وحمض الهيالورونيك على التهاب المثانة المستحدث بالكيتامين في الجرذان البيضاء: دراسة هستولوجية وهستوكميائية مناعية

نجلة العراقي العزب - عبير مصطفي المحلاوي- علا مصطفي
قسم الأنسجة وبيولوجيا الخلية - كلية الطب - جامعة بنها

يحدث التهاب المثانة الناتج عن الكيتامين بشكل شائع لدى الأفراد الذين يسرفون في استخدام الكيتامين. ويرتبط هذا الالتهاب مع تلف في بطانة المثانة وأعراض المسالك البولية السفلى. تعتبر البلازما الغنية بالصفائح الدموية والحمض الهيالورونيك له العديد من الآثار المفيدة في علاج أنواع مختلفة من التهابات المثانة.

المقدمة:
بحث التهاب المثانة الناتج عن الكيتامين بشكل شائع لدى الأفراد الذين يسرفون في استخدام الكيتامين. ويرتبط هذا الالتهاب مع تلف في بطانة المثانة وأعراض المسالك البولية السفلى. تعتبر البلازما الغنية بالصفائح الدموية والحمض الهيالورونيك له العديد من الآثار المفيدة في علاج أنواع مختلفة من التهابات المثانة.

الهدف من البحث:
تقييم تأثير البلازما الغنية بالصفائح الدموية وحمض الهيالورونيك على التهاب المثانة المستحدث بالكيتامين في ذكور الجرذان البالغين.

الطريقة وخطة العمل:

نتائج البحث:
وقد أظهرت الدراسة للمجموعتين الثانية و الثالثة وجود مساحات بؤرية من التحلل والتقرح مع نقص ظاهري في سمك الظهارة البولية. ولوحظ أيضاً ارتفاع خلوي و احتقان بالأوعية الدموية السفلى في أسطح ظهارة المثانة. وظهرت زيادة ذات دلالة إحصائية في NF-κB كتلات التهاب العضلات، بينما كانت هناك زيادة ذات دلالة إحصائية في NF-κB كتلات التهاب العضلات. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين تكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوكميائية المناعية للنوي ضد بروتين Tكاثر الخلايا (PCNC) في ظهارة المثانة. وقص متزايد في ظهارة المثانة من الصبغات الهستوک