The role of ischemic preconditioning at the gracilis muscle of rats in the early phase of reperfusion injury

O papel do precondicionamento isquêmico do músculo grácil de ratos na fase precoce da reperfusão

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ABSTRACT

Purpose: Verify the role of ischemic preconditioning (IPC) in ischemia and reperfusion injury on gracilis muscle of rats.

Methods: Wistar rats (n=30) were distributed in three groups, I/R and IPC groups were subdivided concerning ischemia time. A near-amputation model of the posterior limb was produced by a hip joint level incision, preserving the vascular bundle and the femur bone and ischemia was induced for 2h and 4h, G-I 2h/R (n=6) and G-I 4h/R (n=6), followed by 1h of vascular reperfusion. The preconditioned groups, G-PCI 2h (n=6) and G-PCI 4h (n=6), were preceded by 3 cycles of 5min of ischemia followed by 5min of vascular reperfusion before sustained ischemia. In the Control Group, C-G (n=6) animals were subjected to regional approach. The analysis was done with Light Microscopy (LM).

Results: The levels of fibril fragmentation were progressive in the G-I 2h/R (67% of muscle preservation) and in the G-I 4h/R (0% of muscle preservation). However in the group of the precondition the lesion degree being in level similar to the group controls in the G-I 2h/R (100% of muscle preservation) while at G-I 4h/R occur less protection (67% of muscle preservation). The degree of tissue inflammatory reaction was worst at G-I 4h/R (0% without inflammation signals) than at G-I 2h/R (50% without inflammation signals); while in the precondition group G-IPC-2h (83% without inflammation signals) was better than the G-IPC-4h (67% without inflammation signals). The vascular stasis was absent only in 17% of the G-I 4h/R and in 33% of the G-I 2h/R. In precondition group, however, the vascular stasis was absent in 33% at G-IPC 2h and absent in 50% at G-IPC 4h.

Conclusion: The IPC showed, in an earlier phase, a benefic role at I/R derived injury on gracilis muscle of rats, as proven for the largest preservation of the fibers muscular, smaller inflammatory reaction and smaller vascular stasis.

Key words: Reperfusion Injury. Reperfusion. Skeletal, Muscle. Transplants. Rats.

RESUMO

Objetivo: Verificar o papel do precondicionamento isquêmico (PCI) na lesão de isquemia e reperfusão (I/R) do músculo grácil de ratos. Métodos: Trinta ratos Wistar foram distribuídos em três grupos, sendo G-I/R e G-PCI subdivididos em relação ao tempo de isquemia. Produziu-se modelo de quase-amputação do membro posterior por meio de uma incisão, preservando o feixe vascular e o osso femoral, induzindo-se isquemia por 2h e 4h, G-I 2h/R (n=6) e G-I 4h/R (n=6), seguida por 1h de reperfusão. Os Grupos Precondicionados, G-PCI 2h (n=6) e G-PCI 4h (n=6), foram precedidos por 3 ciclos de 5min de isquemia seguidos de 5min de reperfusão, antes da isquemia sustentada. No Grupo Controle, G-C (n=6) realizou-se apenas a abordagem regional. As avaliações morfológicas foram realizadas por meio de Microscopia Óptica (MO) para avaliação da desestruturação miofibrilar, processo inflamatório e estase venosa no tecido do músculo grácil. Foram atribuídos escores semi-quantitativos às variáveis estudadas. Resultados: Houve preservação miofibrilar de 76% em G-I 2h/R e 0% em G-I 4h/R, G-IPC 2h a preservação foi de 100% e de 67% em G-IPC 4h. A reação inflamatória foi mais acentuada em G-I 4h (100% com inlação) do que em G-I 2h (50%);G-IPC 2h apresentou 82% sem inflamação e G-IPC 4h apresentou 67% sem inflamação. A estase vascular esteve ausente em 17% do G-I 4h e em 33% do G-I 2h; em G-IPC 2h esteve ausente em 33% e em G-IPC 4h em 50%. Conclusão: O PCI mostrou, em uma fase inicial, papel benéfico na lesão de I/R no músculo grácil de ratos, comprovado pela maior preservação das fibras musculares, menor reação inflamatória e menor estase venosa. Descritores: Traumatismo por Reperfusão. Reperfusão. Músculo Esquelético. Transplantes. Ratos.
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Introduction

The use of microsurgical flaps, such as the ones from the gracilis muscle, has brought a revolution in the treatment of complex tissue defects. The gracilis muscle not only allows a quantity of tissue fit for the reconstruction, but it can also exert, through microneurorrhaphy, complementary muscular contractility to the reparation, being beneficial towards ailments such as facial paralysis. However, the microsurgical transference sometimes causes prolonged muscular ischemia, and it means a significant injury to tissue compromising its viability. Some protection strategies have been used for the preservation of the flaps, amongst them the ischemic preconditioning (IPC). The ischemic preconditioning is a surgical alternative that employs short cycles of ischemia followed by reperfusion, previously to a sustained ischemia. The first studies conducted with IPC involved the myocardium. In 1984, Neely and Grotyohann, using the myocardium of rabbits, showed that a short period of ischemia, previously to the sustained cardiac ischemia, produced a protective effect against the cardiac injury of ischemia-reperfusion (I/R). Afterwards, Murry et al., using the myocardium muscle of dogs, could register the protective effect of IPC through a number of experiments, when the term ischemic preconditioning was definitively divulged. From these experiments, the IPC has been used in studies related to different organs and tissues, including in the skeletal muscle. It was observed in the skeletal muscle that, relating with the myocardium, it was necessary more ischemia-reperfusion cycles to obtain a threshold of protection. The maintenance mechanism of the cellular oxidation, the reduction of the production of reactive oxygen specimen (ROS) and the preservation of the integrity and functioning of the cellular membrane, through different mediators, are being studied as probable action mechanisms of the IPC. Although previous studies for the IPC evaluation in skeletal muscles were conducted, the gracilis muscle of rats only recently started to be evaluated for its protection by IPC. Zhang et al. studied the effects of IPC in the late phase of vascular reperfusion of the gracilis muscle of rats demonstrating positive effects in the tissue preservation. The gracilis muscle can have a different behavior from other muscle groups previously analyzed for IPC and there is a lack of studies on the early phase of the ischemia and reperfusion process. This research was developed with the goal to elucidate the cellular alterations of the early phase of ischemia and reperfusion of the gracilis muscle of rats subjected to IPC.

Methods

The present study has been approved by the Committee of Ethics in Research of the Hospital São Paulo – UNIFESP – EPM, under the register number 1095/03, ratified by the Committee of Ethics in Research in Human Beings and Animals of the Lutheran University of Brazil – Rio Grande do Sul – ULBRA-RS, under the register number 2003-006 A. Thirty Wistar male rats, 250 to 350 g, 3 to 4 months old, raised in a conventional animal colony from ULBRA-RS, were kept in controlled conditions of light (clear cycle from 7h to 19h), temperature (22-24°C) and receiving standard food and water ad libitum. The animals were distributed at random in three groups according to the procedure conducted: control group (G-C: n=6), ischemia-reperfusion group (G-I/R: n=12) and ischemic preconditioning group (G-IPC: n=12). The I/R and IPC groups were subdivided taking into consideration the ischemia time (2h and 4h), originating G-I 2h/R (n=6), G-I 4h/R (n=6), G-IPC 2h (n=6) and G-IPC 4h. The animals were kept fasting for solid foods during daytime during a period of three hours prior to the operation procedures and had their weight checked before they entered the procedure room, to calculate the anesthetic dose. They were brought to the operation room in plastic boxes appropriated for transportation and subjected to pre-anesthesia in a chamber saturated with diethyl ether, with the subsequent administration of atropine 0,044mg.Kg⁻¹, through the intraperitoneal way (IP), fulfilling a latency time of 10 minutes until the use of the anesthetic medication. The administered anesthesia was the combination between ketamine chloride (Ketalar®) 50mg.Kg⁻¹ and xylazine chloride (Anasedan®) 10mg.Kg⁻¹ IP. The anesthetic plan was evaluated through the following parameters: respiratory frequency, cardiac frequency, corneal reflex and caudal pinch reflex. The anesthesia complementation was performed with half the initial dose in an average time of 1h30min after induction, being helped by the parameters of the anesthetic plan. They were subjected to epilation around the left posterior limb in the area of the hip articulation in the left posterior limb, followed by antisepsis of the skin with a solution of polivynilpirrolidone (topic PVPI®) at 1%. It was performed the euthanasia of the animal through an intraperitoneal way (IP), fulfilling a latency time of 10 minutes. The anesthesia was the combination between ketamine chloride (Ketalar®) 50mg.Kg⁻¹ and xylazine chloride (Anasedan®) 10mg.Kg⁻¹ IP. The anesthetic plan was evaluated through the following parameters: respiratory frequency, cardiac frequency, corneal reflex and caudal pinch reflex. The anesthesia complementation was performed with half the initial dose in an average time of 1h30min after induction, being helped by the parameters of the anesthetic plan. They were subjected to epilation around the left posterior limb in the area of the hip articulation in the left posterior limb, followed by antisepsis of the skin with a solution of polivynilpirrolidone (topic PVPI®) at 1%. It was used, for the execution of the surgical procedure, a binocular magnifying glass (Heine®) with magnifying capacity of 3,5X. The skin, subcutaneous, and thigh musculature, around the hip articulation, was incised by plans, circumferentially, with a scalpel, preserving the femoral bone and the neurovascular femoral bundle. The femur periosteum was removed, below the hip articulation, by an extension of 0.5 cm. The hemostasis was done by compression. The tissue dissection was done next to the neurovascular bundle with the help of nippers and microsurgical scissors, and later on, with the use of delicate traction of the bundle sheath, it were individualized the arterial, vascular and nervous components. The nervous component was divided for simulating the microsurgical transference. In the G-C, it was performed, after 30min following the vascular isolation, the collection of fragment of the gracilis muscle for analysis under light microscopy. In the G-I/R, after 30min following the vascular isolation, it was proceeded the ischemic interval proposed according to the subgroups, 2h or 4h, followed by 1h of vascular reperfusion. The vascular microclamp (ASSF®), used in the interruption of the blood flow, had pressure graduated for vessels between 0.5 and 1.5mm of external diameter. In the G-IPC, it was applied three ischemia cycles of 5min, intercalated with the same period of arterial reperfusion before the ischemia period, 2h or 4h, and sustained reperfusion for 1h (Figures 1 and 2). Afterwards it was collected, in bottles containing formalin at 10%, a fragment of the gracilis muscle, with an approximate dimension of 1.5 x 1.0 x 0.3 cm, for posterior analysis. After the obtainment of the muscular fragment it was performed the euthanasia of the animal through an
endovenous injection of anesthetic, ketamine chloridate 150mg.Kg⁻¹ and xylazine chloridate 30mg.Kg⁻¹ IV. The analysis under light microscopy was conducted in the Research and Diagnosis Institute of the Santa Casa de Porto Alegre – RS. The fixated material was subjected to dehydration in ethyl alcohol at 70%, 95%, 100%; clearing with Xilol; impregnation with paraffin molten at 60°C; inclusion in rectangular moulds; cut by microtomes (Leica ultracut®UCT RM2025) with thickness of 5μm, discardable plates model 818 size C and coloration by Hematoxiline – Eosine (HE). The analysis was done through a light microscope (Olympus® BX40F40), with zoom between 100 and 1000x in a system of image acquisition through digital system (Samsung aerospace-ICONOS®- Imaging Systems) and computer processing in an equipment with processor with 2.4 Gz speed (Athlon®), with the image diagnosis program (SDI® - System of image diagnosis, developed by Defferari Informatica Ltda®).

It was evaluated the general aspects of the tissue reaction to ischemia and vascular reperfusion, with special attention to fibril fragmentation, to the presence of inflammatory reaction and vascular stasis, being semiquantified in: 0 (lack or discreet presence –25% or less), 1 (moderate presence – 26 to 50%) and 2 (intense – 51% or upper), according to its occurrence. The standard of minor lesion intensity (0) was established in the animals which were less assailed in the G-C and the major degree of considered lesion (2) was based on the animals with the most intense alterations, identified as belonging to G-I 4h/R. The intermediary level of assail (1) was adopted when the lesional characteristics were found between the magnitudes observed in the described groups. It was obtained, thus, the percentage of animals, by analysis subgroups, in relation to the level of intensity of the lesions, looking to establish the influence of the ischemia time in the intensity of the observed lesions. The qui-square test was performed (p<0.05) for all groups.

**Results**

It was verified under light microscopy that, in relation to the control group (Lesion Degree 0: 100%), the G-I 2h/R and G-I 4h/R, in a progressive manner and proportional to the ischemia time, presented fibril fragmentation (Lesion Degree 0: IPC 2h = 100%; IPC 4h = 0%) in relation to the other groups (Figure 3 and 4; Table 1). It was evidenced the biggest level of tissue inflammatory reaction and vascular stasis in the G-I 4h/R (Lesion Degree 0: 0%) in relation to the abdomen. v: femoral vein; G: gracilis muscle. The arrow indicates the position of the vascularization of the gracilis muscle.
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67%) and inflammatory reaction (Lesion Degree 0: IPC 2h = 83%; IPC 4h = 67%), p = 0.038 and 0.041 respectively, (Tables 1 and 2; Figures 4 and 6). There were no benefits of IPC in relation to the vascular findings, comparing to the G-I/R (Table 3; Figures 7 and 8).

### TABLE 1 - Data referent to the presence of fibril fragmentation classified as Degree 0 (lack or discreet presence – 25% or less), Degree 1 (moderate presence – 26 to 50%) and Degree 2 (intense –51% or upper) in the control group (G-C) ischemia/reperfusion group (G-I 2h/R and G-I 4h/R) and IPC group (G-IPC 2h and G-IPC 4h)

<table>
<thead>
<tr>
<th>GROUPS</th>
<th>Degree 0</th>
<th>Degree 1</th>
<th>Degree 2</th>
<th>Percentage of Degree 0</th>
</tr>
</thead>
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<tr>
<td>CONTROL</td>
<td>6</td>
<td>0</td>
<td>0</td>
<td>100</td>
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<tr>
<td>I 2h/R</td>
<td>4</td>
<td>2</td>
<td>0</td>
<td>67</td>
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<tr>
<td>IPC 2h</td>
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<td>IPC 4h</td>
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<td>67</td>
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</tbody>
</table>

For the statistic analysis the lesion degree 1 and 2 were added up forming the degree 1 and the subgroups subjected to 2h and 4h ischemia were placed together as one group, taking into consideration the kind of intervention which was carried out, resulting in the I/R and IPC groups.

### TABLE 2 - Data referent to the presence of tissue inflammatory reaction classified as Degree 0 (lack or discreet presence – up to 25%), Degree 1 (moderate presence – 26 to 50%) and Degree 2 (intense – above 50%), in the control group (G-C), ischemia/reperfusion group (G-I 2h/R and G-I 4h/R) and group IPC (G-IPC 2h and G-IPC 4h)

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<td>100</td>
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<tr>
<td>I 2h/R</td>
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<td>2</td>
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</tr>
<tr>
<td>IPC 2h</td>
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<td>0</td>
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<tr>
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<td>IPC 4h</td>
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<td>2</td>
<td>0</td>
<td>67</td>
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</table>

For the statistic analysis the lesion degree 1 and 2 were added up, forming the degree 1 and the subgroups subjected to 2h and 4h ischemia were placed together as one group, taking into consideration the kind of intervention which was carried out, resulting in the I/R and IPC groups.

### TABLE 3 - Data referent to the presence of vascular stasis classified as Degree 0 (lack or discreet presence – up to 25%), Degree 1 (moderate presence – 26 to 50%) and Degree 2 (intense – above 50%), in the control group (G-C), ischemia/reperfusion group (G-I 2h/R and G-I 4h/R) and group IPC (G-IPC 2h and G-IPC 4h)

<table>
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<tr>
<th>GROUPS</th>
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<th>Degree 1</th>
<th>Degree 2</th>
<th>Percentage of Degree 0</th>
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<tr>
<td>CONTROL</td>
<td>6</td>
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<td>2</td>
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<tr>
<td>IPC 2h</td>
<td>2</td>
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<td>33</td>
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<tr>
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<tr>
<td>IPC 4h</td>
<td>3</td>
<td>3</td>
<td>0</td>
<td>50</td>
</tr>
</tbody>
</table>

For the statistic analysis the lesion degree 1 and 2 were added up, forming the degree 1 and the subgroups subjected to 2h and 4h ischemia were placed together as one group, taking into consideration the kind of intervention which was carried out, resulting in the I/R and IPC groups.
FIGURE 4 - Percentage distribution of the animals as to the intensity of the fibril fragmentation according to the groups. Qui-square Test * (p=0.038)

FIGURE 5 - Photomicrographies showing the intensity of tissue inflammatory reaction under light microscopy: a: Degree 0 (G-C) lack or discreet presence. b: Degree 1 (G-I 2h/R) moderate presence. c: Degree 2 (G-I 4h/R) intense presence. F: muscular fiber. I: Inflammatory infiltrated. V: neutrophilic vasculitis. HE, zoom of 400X

FIGURE 6 - Percentage distribution of the animals as to the intensity of the tissue inflammatory reaction according to the groups. Qui-Square Test * (p=0.041)

FIGURE 7 - Photomicrographies showing the intensity of the vascular stasis under optic microscopy: a: Degree 0 (G-C) normal capillary, preserved endothelium. b: Degree 1 (G-I 2h/R) moderate stasis. c: Degree 2 (G-I 4h/R) accentuated stasis. C: capillary. I: margined neutrophils. Circle in the figure c delimits capillary in detail. HE, zoom of 400X
mediator. Stembergh et al. 11 have verified that the kind of preconditioning has only recently been conducted by between the evaluation of this muscle and the ischemic muscular transferences. However, the combination phenomenon 2. The IPC has a very important action in that can be related to the ischemia-reperfusion tissue transference through microsurgery are submitted 4% to 26% of the final numbers of cases subjected to the and 95% in microsurgical transplants. However, around around 17% of the G-I 4h/R and in 33% of the G-I 2h/R. In precondition group, however, the vascular stasis was absent in 33% at G-IPC 2h and absent in 50% at G-IPC 4h.

So, the findings under light microscopy allowed the verification of proportionality between the ischemia time and the tissue injury, and by other hand allowed the benefits of precondition procedure in protecting the skeletal muscle against the ischemia-reperfusion phenomenon. It is important to highlight the inflammatory findings observed at G-I 4h/R. Although the goal of the study is the early phase of vascular reperfusion, the presence of tissue inflammatory reaction can be an indication of an evolution to an irreversible lesion like necrosis. The morphologic preservation of the musculature of G-IPC 4h, together with the lack of inflammatory reactions, showed a bigger potential of tissue recuperation in this group. The fact is confirmed by a study conducted by Kruel et al. 13, who have verified irreversible alterations in the muscle of dogs subjected by a study conducted by Kruel et al. 13, who have verified irreversible alterations in the muscle of dogs subjected to more than 2h of sustained ischemia. Labbe et al. 14 observed 30,3% of necrosis of the gracilis muscle of dogs subjected to 4h sustained ischemia, in contrast with 2% in the group subjected to 2h of ischemia.

It is identified in the G-I 4h/R a tendency to the accumulation of neutrophils in endovascular and perivascular sites. The findings were compatible with important endothelial lesions. It was not identified in the G-IPC 4h such occurrences, suggesting the attenuation of vascular lesions.

The IPC was able to diminish the adhesion and migration of leucocytes in a study conducted by Akimitsu et al. 15. The adhesion and migration of leucocytes are related to the non-vascular reperfusion phenomenon and the failure of muscular transplants, having a connection with I/R lesions. The mechanism of IPC was not yet clear but it, with no doughty, protected the skeletal muscle from injury of ischemia/reperfusion events.

**Conclusion**

The ischemic preconditioning promoted the protection of the gracilis muscle of rats against the ischemia and vascular reperfusion lesion, in the early phase of reperfusion, according to the morphologic evaluation at light microscopy.

This research showed that levels of fibril fragmentation (Table 1 and Figures 4 and 5) were progressive in the G-I 2h/R (67% of muscle preservation) and in the G-I4 h/R (0% of muscle preservation). However in the group of the precondition the lesion degree being in level similar to the group controls in the G-I 2h/R (100% of muscle preservation) while at G-I 4h/r occur less protection (67% of muscle preservation). The degree of tissue inflammatory reaction (Table 2 and Figures 6 and 7) was worst at G-I 4h/R (0% without inflammation signals) than at G-I 2h/R (50% without inflammation signals); while in the precondition group G-IPC-2h (83% without inflammation signals) was better than the G-IPC-4h (67% without inflammation signals). The vascular stasis (Table 3 and Figures 8 and 9) was absent only in 17% of the G-I 4h/R and in 33% of the G-I 2h/R. In precondition group, however, the vascular stasis was absent in 33% at G-IPC 2h and absent in 50% at G-IPC 4h.

The ischemic preconditioning promoted the

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**Discussion**

It is usually accepted a success level between 90 and 95% in microsurgical transplants. However, around 4% to 26% of the final numbers of cases subjected to the tissue transference through microsurgery are submitted to a surgical review because of vascular interferences that can be related to the ischemia-reperfusion phenomenon 2. The IPC has a very important action in the attenuation of this kind of lesion and has the advantage of its possibility of use in the current surgical period, with no need of additional surgical procedures, and with no collateral effects from the use of drugs applied to raise the tissue viability 10.

The gracilis muscle has been used as an object of study in various animal experiment models related to muscular transferences. However, the combination between the evaluation of this muscle and the ischemic preconditioning has only recently been conducted by Zhang et al. 9, analyzing the nitric oxide as a possible IPC mediator. Stembergh et al. 11 have verified that the kind of fiber and the location of the muscular groups are not directly related to the lesion degree caused by the ischemia and reperfusion phenomenon. There is, therefore, a special interest in the mapping of the behavior of the gracilis muscle facing the ischemic preconditioning in the ischemia/reperfusion lesion.

The tissue transferences subject the muscle to sustained vascular privation with subsequent reperfusion, with the possible occurrence of a significant lesion. The lesions can be reversible or irreversible according to the duration of the stimulus. Once established the irreversible lesion of the cell, the cellular death manifests itself through necrosis or apoptosis. The necrosis is the most common kind of cellular death after exogenous stimulus, such as ischemia; it manifests itself through intense tumefaction or cell rupture, denaturation and coagulation of cytoplasm proteins and the degradation of cellular organelles. There is a preponderant inflammatory component. The apoptosis happens when the cell dies due to the activation of a program of cell removal without the activation of an inflammatory process. The cellular elimination is based on DNA and the cellular fragmentation. The condemned cells are removed with minimal damage to the surrounding tissue 12.

**FIGURE 8** - Percentage distribution of the animals as to the intensity of the vascular stasis according to the groups. Qui-Square Test: findings were non-significant
References


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