The repeated extracorporeal shock waves and the renal parenchyma injury on normal and diabetic rats

A repetição de ondas de choque extracorpóreas e a lesão do parênquima renal em ratos normais e diabéticos

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ABSTRACT

Purpose: To assess the effect of repeated extracorporeal shock waves (ESW) on renal parenchyma of normal and diabetic rats.

Methods: 40 normal rats (A) and 40 diabetic rats (B) were assigned for ESW (Direx Tripter X1® - 14 KVA) as follow: A1/B1 and A3/B3 no ESW; A2/B2 one ESW (2,000 SW); A4/B4 two ESW (4,000 SW) in an elapsed 14 days. All the animals were sacrificed 3 days after the ESW and samples of renal parenchyma were histologically prepared, stained by H&E. For each animal the frequency of hemorrhage focus (HF) in the subcapsular, interstitial and glomerulus area was calculated (percentage) on 20 randomly histological sections.

Results: No one HF was identified in all normal or diabetic animals without ESW (A1, A3 and B1, B3). In the normal rats the HF frequency was similar to one ESW (subcapsular =15%; interstitial =20% and glomerular =10%) or repeated ESW (subcapsular =25%; interstitial =20%; glomerular=10%). In diabetic rats the occurrence of HF with repeated ESW was more frequent (subcapsular =40%; interstitial =30% and glomerular =10%) than with a single ESW (subcapsular =25%; interstitial =15% and glomerular =15%). Conclusion: A single ESW or a repeated ESW caused a mild and similar damage on renal cortex of normal rats. In diabetic rats the repeated ESW may result in an accumulated damage, especially with focus of hemorrhage in subcapsular and interstitial tissue and glomerulus edema.

Introduction

The formation of stones in the urinary tract stems from a wide range of underlying disorders. There are many advances in genetics, pathophysiology, diagnostic imaging, medical treatment, medical prevention, and surgical intervention of nephrolithiasis. Despite of the advances on medical treatment and in surgical management of nephrolithiasis over the past decade, particularly in endoscope technology and techniques, the shock-wave lithotripsy remains the only non-invasive treatment method for stones. Due to its efficacy and safety, the extracorporeal shock wave lithotripsy (ESWL) became the current procedure for treating renal stone either as monotherapy or combined with other forms of lithotripsy. Shockwave lithotripsy predictably damages renal tissue and transiently reduces function in both kidneys. The effects of shock waves on the renal parenchyma were studied through imaging scans, histopathological tests and plasmatic or urinary functional markers. The morphological and functional changes were transients and recovered in few days in dependence of characteristic of the equipment and the quantity of energy focused on stone and adjacent renal parenchyma. The severity of the renal injury caused by ESWL is related to the number of shock waves administered. Considering the prompt recovery from the initial renal injury, a new lithotripsy session could be performed within a short time period. There is no guideline for time interval reapplication of ESWL, administered 3. Considering the prompt recovery from the ESWL is related to the number of shock waves dependence of characteristic of the equipment and the quantity of energy focused on stone and adjacent renal parenchyma. The severity of the renal injury caused by ESWL is related to the number of shock waves administered. Considering the prompt recovery from the initial renal injury, a new lithotripsy session could be performed within a short time period. There is no guideline for time interval reapplication of ESWL, although experimental essay in dogs reported that shock wave reapplication with a 24-hour interval did not cause although experimental essay in dogs reported that shock wave reapplication with a 24-hour interval did not cause much that risk may be improved by ESW is unknown. The large use of ESWL currently, the poor knowledge about injury caused by re-treatment within a short time interval and the empirical way by which treatment is performed, stimulated the development of the present experimental model. Is there an association with extracorporeal shock waves (ESW) and renal parenchyma injury in diabetic animal? The objective of this work was to assess the effect of repeated ESW on renal parenchyma of normal and diabetic rats.

Methods

The experimental protocol was approved by the Ethics Committee of the Federal University of São Paulo – Escola Paulista de Medicina (UNIFESP – EPM), ratified and developed at Maringá State University (UEM). All the procedures followed, rigorously, the existent regulations about animal experimentation. A hundred male Wistar rats weighing between 250-300g were acclimatized to laboratory conditions for 7 days and fed a standard rat chow and water ad libitum. All the surgical procedures and extracorporeal shock waves (ESW) were done under general anesthesia, using ketamine (60mg.Kg\(^{-1}\)) and xylazine (5mg.Kg\(^{-1}\)). At first, all animals were submitted to laparotomy and a radiopaque device (piece of radiopaque surgical gauze) with 0.2cm of diameter was sutured at the adipose tissue near to the right renal pelvis. The standard position of the X-Ray-marker was confirmed by simple post-anterior radiographic method. After fourteen days of surgical procedure, in order to induce diabetes after overnight fasting, 50 rats (Group A) received a single intravenous dose of saline (1.0 mL.Kg\(^{-1}\)). Other 50 rats (Group B) received a single intravenous dose (45 mg.Kg\(^{-1}\)) of alloxan. Immediately after the injection of alloxan or saline solution all animals were maintained with free access to food. At sixth and third-sixth days after the injection of alloxan or saline solution, always at 4pm, the glycemia value was performed in all animals of both groups. From the hundred animals were selected 40 with normal glycemia for the group A (normal rats) and 40 with persistent glycemia over than 200mg/dl for the group B (diabetic rats). Before the first ESW application the urine of 24 hours was collected for microalbuminuria value (mg/L) determination. For ESW treatment the animals were assigned to one of four groups. The animals of group A1 (n=10) and B1 (n=10) did not receive ESW and were sacrificed at 3\(^{rd}\) day; the groups A2 (n=10) and B2 (n=10) received 2,000 ESW at 1\(^{st}\) day and another 2,000 ESW at 14\(^{th}\) day and were sacrificed at 17\(^{th}\) day; the groups A3 (n=10) and B3 (n=10) did not receive ESW and were sacrificed at 17\(^{th}\) day; the group A4 (n=10) and B4 (n=10) received 2,000 ESW at the 1\(^{st}\) day and were sacrificed at 3\(^{rd}\) day. A Direx Tripter X1\(^{®}\) lithotriptor with 14 KVA of intensity, produced a tension ranging from 500 to 520 Bar and the focus system to be used was obtained by means of acoustic lens and the contact with the animal skin through a bag containing water and gel. Previously to the adjustment of focus system the radiopaque device was localized by fluorescope monitor. The ESW were focused about 0.5cm laterally to the radiopaque device and were applied 2,000 shock waves in each session. According to each group, 72 hours after the ESW, under anesthesia, the animal was submitted to a right nephrectomy and 3 fragments were collected from the area near the radiopaque device (0.5cm) where was...
focused the ESW. The samples of renal parenchyma were fixed in 10% formalin solution, embedded in paraffin wax and 5 µm sections were cut and stained with hematoxilin and eosin (H&E). Histological evaluation was undertaken by an independent pathologist who had no knowledge of the experimental groups from which the specimens were derived. The evaluated parameters were hemorrhagic focus (HF) in the renal cortex (subcapsular, interstitial and glomerular areas). The macroscopic assessment was performed using the following grading scoring: 0 (no changes), 1 (mild = 1 to 7 HF), 2 (moderate = 8 to 15 HF) and 3 (severe = 15 to 20 HF). The abnormalities detected in 20 random sites in every animal plate were subjectively scored and compared with the controls. The rate of occurrence of HF (percentage) was calculated in three plates for each animal. In other 20 random places was measure the thicknesses (µm) of the glomerulus and Bowman capsule, and then calculated the ratio glomerulus/capsule. The statistical analysis was performed using the Mann-Whitney test due to the paired structure of data in order to compare the variation: A1xA2; A3xA4; B3xB4. The Exact Fisher test was performed due to the paired structure of data in order to compare the variation: A1xA2; A3xA4; B1xB2; A2xB2; A4xB4. The significance level was established at 0.05 or 5%.

**Results**

The alloxan was effective to induce the diabetes and all animals from group B (diabetic) developed a functional impairment in microalbuminuria due to a mild diabetic nephropathy. There was a significant increase in microalbuminuria excretion (p < 0.05) in all sub-groups B (B1 to B4) in comparison with the non diabetic control group A (A1 to A4) as shown in Tables 1 and 2. Different superscript letters indicate statistically significant differences with regard to the corresponding control value (Figures 1 and 2). The analysis of 20 areas in the three plates stained by H&E as shown in characteristic photomicrography of Figure 1, allowed to perform the semi-quantitative evaluation and calculated the HF frequency (in percentage) that were plotted at Figure 3 (A, B and C). The frequency of absent subcapsular hemorrhage was quite similar in the normal rats with a single (A4 = 85%) or repeated ESW (A2 = 75%). However, in the diabetic rats the HF (mild and severe) were more frequent mainly after repeated ESW (B2 = 40%) in comparison with repeated ESW (A2 = 25%) in normal rats (Figure 3A). The mild and severe interstitial hemorrhage were low frequent in the normal rats, with one (A4 = 20%) or two ESW (A2 = 20%), that means 80% of HF absence in each group. In the diabetic rats the frequency of interstitial hemorrhage was more frequent with repeated ESW (B2 = 30%) than one ESW (B4 = 15%) (Figure 3B). The frequency of glomerular hemorrhage was the same in normal rats with one ESW (A4 = 10%) or repeated ESW (A2 = 10%). In the diabetic rats the frequency of HF was similar with one ESW (B4 = 10%) or repeated ESW (B2 = 15%) (Figure 3C). Different superscript letters indicate statistically significant differences with regard to the corresponding group value.

**TABLE 1** - Distribution of the microalbuminuria values (µg/L) in 24 hours urine, just before the first ESW. Values are given as mean ± sd and median

<table>
<thead>
<tr>
<th>Group A (non Diabetes)</th>
<th>Group B (Diabetes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1 6.3a 8.2b 5.9c 6.0d</td>
<td>B1 30.3a 30.3b 30.8c 31.7d</td>
</tr>
<tr>
<td>sd 0.3 0.2 0.2 0.3</td>
<td>0.4 0.4 0.3 0.3</td>
</tr>
<tr>
<td>Median 5.6 7.7 5.6 5.6</td>
<td>25.0 25.0 25.0 25.0</td>
</tr>
</tbody>
</table>

Mann-Whitney test (Z crit = 2.12): aA1 < B1 (Z cal = 3.74*); bA2 < B2 (Z cal = 3.66*); cA3 < B3 (Z cal = 3.87*); dA4 < B4 (Z cal = 3.84*)

**TABLE 2** - Distribution of means (sd) and median of the ratio of measures of Bowman capsule and glomerulus diameters (µm) in the groups of diabetics and non diabetics rats with one or two ESW

<table>
<thead>
<tr>
<th>Group A (non Diabetes)</th>
<th>Group B (Diabetes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1 1.42a 1.11e 1.33b 1.17b,f</td>
<td>B1 1.39c 0.57c,e,g 1.36d 1.05d,f,g</td>
</tr>
<tr>
<td>sd 0.2 0.3 0.3 0.2</td>
<td>0.2 0.4 0.3 0.3</td>
</tr>
<tr>
<td>Median 1.37 0.8 1.28 1.10</td>
<td>1.33 0.35 1.30 0.8</td>
</tr>
</tbody>
</table>

Mann-Whitney test (Z crit = 2.12): aA1 > A2 (Z cal = 2.68*); bA3>A4 (Z cal = 2.42*); cB1 > B2 (Z cal = 3.38*); dA3 = B3 (Z cal = 3.75*)

Non diabetes x diabetes

Mann-Whitney test (Z crit = 2.12): A1 = B1 (Z cal = 1.48); eA2>B2 (Z cal = 2.83*); A3 = B3 (Z cal = 1.75); fA4 > B4 (Z cal = 2.97*)

One x Two ESW

A2=A4 (Z cal = 1.14); gB2 < B4 (Z cal = 2.84*)
FIGURE 1 - Photomicrography of hemorrhagic focus (HF). A-subcapsular area and mild hemorrhage (arrow) (H&E-200X). B-interstitial area and mild hemorrhage (arrow) (H&E –400X). C- glomerular area and mild hemorrhage (arrows) on renal cortex. (H&E –400X)

FIGURE 2 - Photomicrography of glomerular area of normal rat (A) and a mild hemorrhage (black arrow) of glomerular area of a diabetic rat and the diameters (µm) of Bowman capsule (yellow arrow) and glomerulus (orange arrow). (H&E -200X)

FIGURE 3 - Plotted occurrence (frequency in percentage) of hemorrhagic focus on subcapsular (A), interstitial (B) and glomerular (C) areas on renal cortex of normal and diabetic rats with one session ESW (A4 / B4) or repeated ESW (A2 / B4)
Discussion

Experimental works using rabbits\textsuperscript{10}, dogs\textsuperscript{11,12}, pigs\textsuperscript{3,13-15} and rats\textsuperscript{3,5,18,16} confirm the transient and focal ESW renal injury. Some factors can produce contradictory results including the sample size, different equipments with different energy principles, number of impulses and focus size\textsuperscript{2}. In our experimental model, the area affected by the shock wave, with the lithotriptor employed in small animal kidney such as rat would have a proportionally larger area (100\%) of the renal parenchyma affected by the shock wave in comparison to humans (10\%), but a proportional dose of energy for lithotripsy (2,000 EWS) as used in humans was used by the authors. Diabetes is a disease that promotes morphological and biochemical changes\textsuperscript{6-7} that worsen the conditions of renal parenchyma and is an etiologic impairment in the follow-up of urolithiasis treatment\textsuperscript{17}. The present experimental model of diabetes induced by alloxan was enough, 30 days after\textsuperscript{2}, to promote a diabetic nephropathy confirmed by the 24 hours microalbuminuria values\textsuperscript{4} (Table 1). Research in lithotripsy started with the effort to characterize acute shock wave damage to the kidney. It is appreciated now that shock wave trauma is primarily a vascular lesion, that injury is dose dependent, and if the hemorrhage is severe may be associated to a permanent loss of functional renal mass\textsuperscript{13,18,19}. Our results confirm the vascular injury due to ESW. As it was expected, it was not identified any hemorrhage focus (HF) in all normal or diabetic animals without ESW (A1, A3 and B1, B3). In the normal rats the HF frequency was similar with one ESW (A4) in the three areas (subcapsular =15\%; interstitial =20\% and glomerular =10\%) or repeated ESW (A2) in the same areas (subcapsular =25\%; interstitial =20\%; glomerular =10\%). A second ESW after fourteen days did not improve the HF frequency. The elapsed time will be enough to promote a complete recovery of the first ESW. Overall, this animal model, based on those parameters of morphological injury showed quite appropriated for the study of the effects of ESW on renal parenchyma. The lack of a consensus concerning the choice of time interval to repeat ESWL is due to enormous variability of particular conditions found in the use of ESWL in humans\textsuperscript{5,7,20} and the limitations of the animals model\textsuperscript{10-21}. Our animal model suggests that the mild vascular injury on the renal parenchyma can be recovered in fourteen days, as is showed by the current reports in the biomedical literature\textsuperscript{16,18,22}. On the other hand, on diabetic rats the occurrence of HF with repeated ESW (A2) was more frequent in the three areas (subcapsular =40\%; interstitial =30\% and glomerular =10\%) than with a single ESW (A4) in the same three areas (subcapsular =25\%; interstitial =15\% and glomerular =15\%). Diabetes is a disease that promotes morphological and biochemical changes that worsen the conditions of renal parenchyma and it is an etiologic impairment in the follow-up of urolithiasis treatment\textsuperscript{2}. Our experimental results suggested that the diabetes can improve the risk of renal injury in the treatment with repeated ESW. The transient functional damage after ESW would be a result of morphological injury to the nephrons\textsuperscript{12,13,21}. Plasmatic and urinary markers such as N-acetyl-beta-glucosaminidase (NAG), beta-galactosidase, gamma-glutamyl-transferase and high molecular weight proteins such as macroglobulins and calbindin d28K assessed renal trauma resulting from shock waves because they are proteins that, under normal conditions, are not filtered by the glomerulus\textsuperscript{2,4}. In our work the ESW caused an injury to the nephrons expressed by the measure of the ratio between the capsule of Bowman diameter and the glomerulus diameter (Table 2). In normal rats occurred a thin peripheral space of 2.0-3.0µm (Figure 2A) that diminished or disappeared in some groups like in diabetic rats with two ESW (Figure 2B). The enlargement of glomerulus was not only due to hemorrhage focus but mainly by the liquid sequestration causing the edema. In the normal rats the edema was similar to one or two ESW and, similar to the HF, can recover in fourteen days. However, in diabetic animals the edema was more intense and worst after the second ESW. These findings suggest once more, that diabetes was a risk factor for reapplication of ESW. Considering the effects of ESW on the frequency of hemorrhage focus and glomerulus impairment suggested by this work, and the well known fact that the damage is dose-dependent, staged ESW should be advisable in a clinical setting of diabetic patient.

Conclusion

A single ESW (2,000 SW) or a repeated ESW (4,000 SW) caused a mild and similar damage on renal cortex of normal rats. In diabetic rats the repeated ESW (4,000 SW) may result in an accumulated damage, especially with focus of hemorrhage in subcapsular and interstitial tissue and glomerulus edema.

References


