Qualitative and Quantitative Study of the Changes in the Ultrastructure of Mammalian Adrenal Cortex Caused by the Venezuelan Tigra Mariposa (Bothrops venezuelensis) Snake Venom

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Abstract

The damage of the adrenal gland by snake venoms needs to be clarified. Lethality (LD₅₀) of Bothrops venezuelensis (Bv) venom was established by intraperitoneally mice injections. Preparation of specimens for transmission electron microscopy samples from cortex adrenal gland biopsies at 3, 6, and 24 h was processed. The quantitative description by the principal component analysis (PCA) of the adrenal gland was as follows: thickening of the capillary endothelium, area of the capillary lumen, cell nucleus area, enlargement of the perinuclear space, number of mitochondria, area of the mitochondria, number of mitochondrial cristae, number of cristae per mitochondrial unit, and tubular diameter of the smooth endoplasmic reticulum (SER). Sections of the adrenal cortex, after 3 h postinjection with Bv venom showed in the cortical cells: mitochondria with tubular cristae and slightly swollen SER cisternae, nucleus with variable heterochromatin content, irregular edges, and swollen nuclear envelope. After 6 h, cells with swollen nucleus envelope, electron dense lipids and mitochondria with loss of their cristae were observed. Myelin figures, close to the microvilli of the cortical cell, multivesicular bodies, swollen profiles of the SER, and electron dense lipid drops were noticed. After 24 h, thickening of the endothelial wall, fenestrae and projections into the capillary lumen, loss of the mitochondrial cristae, destruction of the capillary and the plasma membrane of the cortical cell, multivesicular body, SER loss, and an enlargement of the perinuclear space were detected. In the quantitative PCA, there were significant changes after the venom treatments.

Keywords: Adrenal gland, Bothrops venezuelensis, mitochondria, principal component analysis, smooth endoplasmic reticulum, ultrastructure, venom

INTRODUCTION

Hemostasis disorders and organ damages by proteolytic action are the main signs of envenoming by Bothrops snake venoms.[1-6] The snake venom proteases have been related to the pathogenicity of the adrenal gland injuries.[7-16] Bothrops venezuelensis (Bv) venom effects[1] were carried out covering the first 24 h, obtaining samples from the adrenal gland cortex of injected mice at 3, 6, and 24 h.

In the present work, a group of subcellular changes were showed, which could explain the pathophysiological alterations concerning the adrenal gland caused by Bv venom.

MATERIALS AND METHODS

Snakes

The “tigra mariposa” Bv snakes were obtained from the Waraira Repano mountains, at North of Caracas city (Bolivarian Republic of Venezuela), a typical subtropical rain forest, with average temperatures of 25°C, annual rainfall

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of 1500–2000 mm, and atmospheric humidity superior than 80%. Six both sexes’ snakes were located and maintained at the Serpentarium of the Tropical Medicine Institute of the Universidad Central de Venezuela (Caracas, Venezuela).

**Venom collection**

Venom was achieved by permitting the snakes to bite into a Para-film® stretched over a disposable plastic cup. The venom samples were centrifuged (500×g for 15 min) and filtered through 0.45 μm filter under positive pressure. The venom was frozen at −80°C, lyophilized, and maintained at −30°C until used. Sublethal (5.0 mg/kg) samples of reconstituted venom were prepared for the experimental injection of mice.

**Mice**

Ultrastructural adrenal gland studies were performed using NIH mice (18–22 g) purchased from the vivarium, of the National Institute of Hygiene “Rafael Rangel,” Caracas, Venezuela. Animals were supplied with water and food ad libitum until used.

**Ethical statement**

Qualified staff controlled all the experimental methods relating to the use of live animals. Applicable regulations as well as institutional guidelines, according to the protocols approved by the Institute of Anatomy Ethical Committee of the Universidad Central de Venezuela, on March 7, 2018, under assurance number 07-03-18. The research was carried out in accordance with the U. K. Animals (Scientific Procedures) Act, 1986 and associated guidelines, EU Directive 2010/63/EU for animal experiments.

**Lethal activity**

Lethality of *Bv* venom was established by intraperitoneally injections into mice, and the LD₉₀ calculation was done, according to the method of Spearman-Kärber. Six groups containing five mice were injected (0.2 mL) intraperitoneally, with different venom concentrations (4.0–7.0 mg/kg) diluted in phosphate-buffered saline (PBS) solution. An equivalent volume of PBS was injected as a negative control group. The animals were observed for 48 h after injections.

**Preparation of specimens for electron microscopy**

Three adult male NIH strain mice (normal control group) were injected intravenously (IV) in the tail vein with 0.1 mL of PBS solution. Nine mice (experimental group) were injected IV with 5.0 mg/kg of *Bv* in 100 μL of PBS. After 3, 6, and 24 h, three mice from each group were ready for adrenal gland biopsies. The fragments were instantly obtained, from control and experimental mice, after to be euthanized by cervical dislocation. The samples were without delay in situ fixed with 3% glutaraldehyde and then with 1% OsO₄ (both fixatives diluted in 320 mM phosphate buffer saline, pH 7.4), dehydrated in ethanol, and embedded in LX-112 resin (Ladd Research Inc.). Ultrathin sections were stained with uranyl acetate and lead citrate. At that point, observed with an FEI, Tecnai Spirit 12G2 model transmission electron microscope, with an accelerating voltage of 100 kV.

**Digitization of the image**

For the descriptive study of the biological systems from a micrograph, the magnification used for their registration was considered, as well as the magnification factor, with which the revealed physical image was obtained. Using the “mouse,” we could delineate points, the length, the angle, the area, etc., in the image taken directly from the TEM so that the object in the image could be established in real time. Thus, the images acquired directly from the electron microscope were used to the quantitative study. The structures measured with the TEM were processed with the computer to analyze and classify the data using statistical programs. They were included and maintained the size according to the original magnification, with which it was achieved. The images obtained by the FEI Quanta 250 FEG equipment were stored digitally in a computer for further study.

**Calculated usage of the results**

**ImageJ**

From the micrographs acquired in the thin and thick sections, morphometric magnitudes were carried out through the IMAGEJ program.

**Principal component analysis**

The principal component analysis (PCA) consisted of expressing a set of variables in a set of linear combinations of factors, which are not correlated among them. This process permitted to symbolize the original data (individuals and variables) in a space of fewer dimensions than the original space. The assessed variables for the quantitative description of the adrenal gland were thickening of the capillary endothelium, area of the capillary lumen, cell nucleus area, swelling of the perinuclear space, number of the mitochondria, area of the mitochondria, number of mitochondrial cristae, number of cristae per mitochondrial unit, and tubular diameter of the smooth endoplasmic reticulum (SER).

**Statistical analysis**

ANOVA is a statistical procedure that uses the analysis of variance, which allows examining, if more than two groups diverge significantly among themselves, in terms of their means and variances. Classically, the ANOVA is used to associate a probability, with the assumption that the mean of a group of scores is different from the mean of another group of scores. Through the comparison, it was attempted to test a hypothesis of difference between more than two groups.

**Smallest statistically detectable difference test**

It was carried out comparing the statistical values obtained by *Bv* venom treatment with those obtained in the normal control group treatment. The letters a, b, c, d, e, f, and g are derived from a posteriori smallest statistically detectable difference (SDD) test.

**Results**

**Lethal activity of Bothrops venezuelensis venom**

The lethal activity of *Bv* venom expressed as LD₅₀ was 6.4 mg/kg.
Adrenal gland changes induced by Bothrops venezuelensis venom analyzed by transmission electron microscopy

In the adrenal gland cortex normal control, the mitochondria were observed with their tubular cristae, central nuclei with the normal nuclear envelope, associated with heterochromatin, rough endoplasmic reticulum (RER) short cisternae, tubular profiles of the SER, Golgi (Go) reticular apparatus, lipofuscin granules in different stages of growth (triangle), capillary microvilli (oval), and a lipid drop (Lip) [Figure 1].

The adrenal gland cortex after 3 h of Bv venom injection showed loss of the continuity of the endothelial wall, projections into the capillary lumen, and remains of the endothelial wall (the plasma membrane was not distinguished). There was a larger space between the limit of the endothelial cell and the cortical cell. In the cortical cell, the round and slightly elongated profiles corresponded to the SER tubules. Swollen mitochondria with scarce cristae were observed [Figure 2a].

In the cortical cell, a nucleus of irregular contours with abundant heterochromatin and the enlargement of the perinuclear space was noticed. There was a Golgi apparatus, clearly swollen mitochondria, very swollen cisternae of RER, and an autophagic vacuole, with mitochondrial debris [Figure 2b].

Circular profiles of different diameters corresponded to cross-sections of SER. Mitochondria with tubular cristae were also observed [Figure 2c].

Lipid droplets of different electron densities and swollen mitochondria were noticed. Adjacent to the mitochondria, a lipofuscin granule was seen. Circular profiles and small tubules of SER were observed. There were also profiles of larger diameter [Figure 2d].

The adrenal gland cortex after 6 h of Bv venom injection showed a fibroblast with an elongated nucleus and an enlargement of the perinuclear space, which lies between the plasma membrane of the cortical cell and the capillary wall where the fenestra was noticed. Swollen cisternae of the RER were detected. The separation of the plasma membrane from the cortical cell and the fibroblast was seen. A mitochondrion with tubular ridges and a lipid drop was also observed [Figure 3a].

A pseudoinclusion of the cytoplasm was seen, surrounded by a nuclear envelope and the nuclear side; besides heterochromatin was noticed [Figure 3b].

A very electron dense lipid drop and a myelin figure were seen. The microvilli of the cortical cell and the mitochondria were

![Figure 1: The adrenal gland cortex normal control. The mitochondria are observed with the tubular cristae (star), central nuclei (Nu) with normal perinuclear space (arrow) associated with heterochromatin, short cisternae of the rough endoplasmic reticulum, tubular profiles of the smooth endoplasmic reticulum, Golgi (Go), lipofuscin granules in different stages of growth (triangle), capillary microvilli (oval), and a lipid drop (Lip). Micromark = 1 μm](#)

![Figure 2: Adrenal gland cortex after 3 h of Bothrops venezuelensis venom injection. (a) An erythrocyte is observed within the capillary (E), loss of the continuity of the endothelial wall (oval), projections of the capillary lumen and remains of the endothelial wall (the plasma membrane is not distinguished) (circle). The longer oval indicates a larger space between the limit of the endothelial cell and the cortical cell. In the cortical cell, the round and slightly elongated profiles correspond to the smooth endoplasmic reticulum tubular. Swollen mitochondria with scarce cristae (stars) are observed. Micromark = 1 μm. (b) In the cortical cell, a nucleus (Nu) of irregular contours with abundant heterochromatin and dilated perinuclear space (arrowhead) is noticed. There is a Golgi apparatus (Go), a clearly swollen mitochondria (star), a very swollen cisternae of rough endoplasmic reticulum (RER) and an autophagic vacuole with mitochondrial debris (triangle). Micromark = 1 μm. (c) The circular profiles of different diameters correspond to cross-sections of smooth endoplasmic reticulum. Mitochondria with tubular cristae (star) are observed. Micromark = 1 μm. (d) Lipid droplets (Lip) of different electron density are seen; swollen mitochondria (star) where a lipofuscin granule is observed adjacent to the mitochondria. Circular profiles and small tubules of smooth endoplasmic reticulum are seen). There are also profiles of larger diameter. An erythrocyte is located (E). Micromark = 1 μm](#)
appreciated. The profiles of different diameter of SER were observed [Figure 3c].

Mitochondria with swollen cristae, a lipid droplet, profiles of different diameter of SER, and a multivesicular body (MVB) were also seen [Figure 3d].

The adrenal gland cortex after 24 h of Bv venom injection, the endothelial wall showed some fenestrae, and there was an area of wall discontinuity. Several lipid drops, profiles of SER, as well as areas lacking of them were noticed. Various swollen mitochondria were seen [Figure 4a].

The destruction of a capillary was observed where the endothelial wall disappeared. A MVB, loss of the SER, and a lysosome were observed [Figure 4b].

The nucleus exhibited some areas of enlargement of the perinuclear space [Figure 4c]. It was observed a nucleus without heterochromatin and an electron dense body that could be the nucleolus, but with its characteristic structure lost (fibrillar center, dense fibrillar component, and granular component). The structure of the perinuclear space was not distinguished, and in the cytoplasm, there was degeneration, with swollen profiles of SER. Mitochondria with swollen cristae were observed [Figure 4d].

The quantitative principal component analyses under Bothrops venezuelensis venom activity and smallest statistically detectable difference test

In the statistical values obtained by Bv venom treatment, compared with those obtained in the control treatment, there was a significant difference between the results [Table 1]. The letters a, b, c, d, e, f, and g are derived, from a posteriori SDD test, in which the groups, that have significant differences amid their averages [Table 1].

Table 2 shows that principal component (PCA)-1 [X axis of Figure 5] describes 90.89% of the information of the variable system and (PCA)-2 [y axis of Figure 5] 4.9% of this
Table 1: Quantitative analysis of principal component analyses, under Bothrops venezuelensis venom actions

| B. v venom (time in hours) | Thickening of the capillary endothelium (µm) | Capillary lumen (µm²) | Nucleus area (µm²) | Thickening of perinuclear space (µm) | Number of mitochondria (µm²) | Area of the mitochondria (µm²) | Number of mitochondria cristae | Diameter of the cisternae of the SER (µm) |
|---------------------------|---------------------------------------------|-----------------------|--------------------|-------------------------------------|----------------------------|-----------------------------|-------------------------------|-----------------------------------|
|                           | Mean ± SD                                   |                       |                    |                                     |                            |                             |                                |                                    |
| 3                         | 0.29±0.017                                 | d                     | 8.68±0.10          | d                                   | 16.08±0.31                 | ab                          | 1.55±0.080                    | 14.88±0.19                       |
| 6                         | 0.47±0.012                                 | c                     | 9.98±0.14          | b                                   | 12.53±0.48                 | c                           | 1.35±0.092                    | 16.33±0.17                       |
| 24                        | 0.81±0.014                                 | a                     | 11.10±0.17         | a                                   | 8.33±0.43                  | dce                         | 1.08±0.115                    | 19.18±0.18                       |

To evaluate the effect of thickening of the capillary endothelium, capillary lumen, nucleus area, thickening of perinuclear space, number of mitochondria, area of the mitochondria, number of mitochondrial cristae, and diameter of the cisternae of the smooth endoplasmic reticulum, an ANOVA of two factors at 95% confidence was developed in which it was found that there were significant differences for changes in the treatments (factor 1) \( P=0.00 \) and significant difference for times (factor 2) \( P=0.00 \), with an interaction of \( P=0.00 \). The table shows the trends of the means and standard deviations for the two treatments over time, obtained during the study of the above parameters. The letters a, b, c, d, e, f, and g are derived from a posteriori SDD, in which the groups that have significant differences, in alphabetical order among their means, and they were given with different letters. The group with the higher mean was defined with the letter a, and when its average significantly decreased, they were assigned and denoted by different letters (a to g)\(^2\). ANOVA: Analysis of variance, SDD: Smallest statistically detectable difference; SD: standard deviation; B. v: Bothrops venezuelensis, SER: Smooth endoplasmic reticulum

Table 2: Principal components analyses

| Percentage | Accumulated percentage | Percentage | Accumulated percentage | Percentage | Accumulated percentage |
|------------|------------------------|------------|------------------------|------------|------------------------|
| 99.89      | 99.89                  | 95.84      | 95.84                  | 94.90      | 94.90                  |
| 0.496      | 0.496                  | 0.436      | 0.436                  | 0.436      | 0.436                  |

The PCA-1 (X axis of Figure 5) describes 90.89% of the information synthesized in the eight variables. The PCA-2 (Y axis of Figure 5) 9.99% of this information. The figure as a whole expresses 95.85% of the variables (accumulated percentage) which is considered a satisfactory representation. PCA: Principal component analysis.
At 6 and 24 h [Tables 1-3 and Figures 5 and 6a], the current study demonstrated that Bv venom significantly induced an increase in the thickening of the capillary endothelium, being more intense at 24 h (0.81 μm) (normal = 0.31 μm).

The changes in the capillary lumen appeared 6 h after the venom injection, increasing to 9.98 μm² (normal = 8.48 μm²). At 24 h, an increase up to 11.10 μm² (control = 8.78 μm²) was observed [Tables 1-3 and Figures 5 and 6b].

The nucleus area only had a significant decrease at 24 h (8.33 μm²) (normal = 17.28 μm²) after venom injection, although at 6 h, it had a discrete decrease (12.53 μm²) (control = 16.93 μm²) [Tables 1-3 and Figures 5 and 6c].

The perinuclear space at 3 h remains within the parameters of normal control but increased slightly at 6 h (16.33 μm) (control = 14.10 μm), but at 24 h (19.18 μm) (control = 14.0 μm), the rise is higher [Tables 1-3 and Figures 5 and 6d].

The number of mitochondria remained within the parameters of normal control (1.78 mit/analyzed fields), until 24 h (1.08 mit/analyzed fields) when there was a decrease in their number [Tables 1-3 and Figures 5 and 7a].

The mitochondrial area showed at 6 (0.30 μm²) (normal control = 0.13 μm²) and 24 h (0.37 μm²) (normal control = 0.14 μm²) postinjection of venom that their area increased [Tables 1-3 and Figures 5 and 7b].

Merely, at 24 h, the number of cristae sensibly decreased (8.98 cristae for mitochondria (crist/mit) (normal control 26.62 crist/mit), maintaining normal values at 3 and 6 h [Tables 1-3 and Figures 5 and 7c].

The diameters of the cisternae of the SER showed an increase, at 3 h after the injection of the venom (0.047 μm) (normal

Table 3: Principal component variables

| PCV        | Axis 1 | Axis 2 |
|------------|--------|--------|
| AMit       | 0.325  | −0.106 |
| NMitC      | −0.328 | −0.032 |
| NMit       | −0.319 | 0.175  |
| NA         | −0.322 | −0.208 |
| DSERC      | 0.325  | −0.075 |
| TCE        | 0.315  | 0.12   |
| CL         | 0.301  | 0.494  |
| TPS        | 0.316  | 0.388  |

The information shows the principal component variables that were correlated. It was observed that there was a high positive correlation between the area of the mitochondria, the diameter of the cisternae of the smooth endoplasmic reticulum, thickening of the capillary endothelium, capillary lumen, and the perinuclear space in Axis 1 and with a correlation positive, between the number of mitochondria, thickening of the capillary endothelium, capillary lumen, and the perinuclear space in Axis 2. PCV: Principal component variables, AMit: Area of mitochondria, NMitC: Number of cristae, NMit: Number of the Mitochondria, NA: Nucleus Area, DSERC: Diameter of cisternae of the smooth endoplasmic reticulum, TCE: Thickening of the capillary endothelium, CL: Capillary lumen, TPS: Thickening perinuclear space.
control = 0.011 μm). At 6 h, an intense increase was detected (0.147 μm) (normal control = 0.009 μm) and at 24 h, a higher increase (0.209 μm) (normal control = 0.007 μm) in the diameter of the cisternae was observed [Tables 1-3 and Figures 5 and 7d].

**DISCUSSION**

The studies in transmission electron microscopy of several organs have allowed describing the alterations caused by the bothropic venoms; however, there are not records on similar studies in the adrenal gland. Therefore, it was decided to study the alterations in this gland, produced by the venom of *Bv*. The adrenal gland is an organ of capital importance for the correct homeostasis equilibrium in mammals throughout the synthesis of mineralocorticoids and glucocorticoids hormones.[21] The endocrine adrenal gland controlling the intensities of cortisol and other vital stress-associated hormones is an essential organ of the hypothalamic–pituitary–adrenal axis (HPA) designated as the body’s “stress system.”[21,22]

The participation of pituitary–adrenal axis (HPA) in the acute phase of circulatory shock has been confirmed by histopathological validation.[23,24] The adrenal gland cortex is responsible for the production of cortisol, aldosterone, and androgens, which control the electrolyte balance, blood pressure, lipid and glycogen metabolism, and estrogenic biosynthesis[23] is controlled by neuroendocrine hormones, which are originated from the pituitary gland, under the control of the brain hypothalamic region, as well as by the renin-angiotensin system.[20] The biochemical or ultrastructural alterations induced by the toxins of the *Bv* venom can be generated indirectly through the previous system or directly through the adrenal gland. In the present study, it is shown that adrenal cortex injuries induced by *Bv* venom have diverse toxin-pharmacological effects, resulting in important adrenal gland subcellular damages. It could be speculated that by their enzymatic character, mainly the phospholipases and metalloproteases, may be involved in this damage.

In an early period, 3 h after *Bv* venom injection, the adrenal cortex showed swollen SER of cortical cells. The endoplasmic reticulum (ER) damage is caused by disturbances in it structure and function, mainly triggered by glucose deprivation and depletion of Ca²⁺ stores; also produced by the exposure to free radicals, and the accumulation of unfolded or misfolded proteins.[27] This lead to the initiation of a cell stress response, such as the unfolded protein response. As it is known, surface and secreted proteins are synthesized, fold, and assemble before being transported throughout the ER.[27]

In the nucleus, an ultrastructural damage with variable heterochromatin content, irregular edges, and swollen perinuclear space were seen. The nuclear envelope has physical connections to chromatin, via nuclear lamins (Class V intermediate filaments), interacting with membrane-associated

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**Figure 7:** (a) number of mitochondria fluctuations, (b) mitochondria area variations, (c) number of mitochondria cristae, and (d) diameter of the smooth endoplasmic reticulum cisternae after 3, 6, and 24 h of *Bothrops venezuelensis* venom injection were evaluated by an analysis of variance of two factors at 95% confidence. It was found that there were statistically significant differences for changes in treatments (factor 1) \( P = 0.00 \) and significant difference for times (factor 2) \( P = 0.00 \), with an interaction of \( P = 0.00 \). The figure shows the tendencies of the means by the two treatments over time. The values are showed in Table 1.
proteins and forming the nuclear lamina on the interior of
the nuclear envelope that links straight to chromatin[29] and to
cromatin-binding nuclear membrane proteins. The shape of a
nucleus is determined by the nuclear lamina, which is closely
related with the inner nuclear membrane, and the cytoskeleton.
Nevertheless, the way linking the differentiation condition
of a cell and the shape variations of its nucleus are not well
comprehended. The nuclear-size escalation has been related
with nuclear lamina CxxM motif proteins, such as lamins and
cugelkern gen.[20,30]

Nuclear envelope is a theorized regulator, of the effects of
cellular influences on chromatin or gene expression.[31,32] The
exact mechanisms governing the edema of the perinuclear
space caused by the toxins of Bv are unknown; it is possible
that the phospholipases, serine proteases, and metalloproteases
may be participating in this damage. A swollen SER cisternae
and a pathological autophagic vacuole, with swelling of the
mitochondrial cristae, as well as an increase in the swelling
of the SER, probably were also produced by the action of
phospholipase A_{2} (PLA_{2}).[33] Contrariwise, as it is known,
under physiological conditions, autophagy is considered as
a normal event, which is required for maintaining a healthy
mitochondrial compartment, by the elimination of old and
damaged mitochondria.[34]

After 6 h postinoculation with Bv venom, the adrenal cortex
changes observed at 3 h (swollen SER and nucleus with swollen
nuclear envelope) persisted, but now, other ultrastructural
changes such as electrondense lipids and mitochondria with
swollen cristae as well as a pseudooinclusion in the nucleus
with associated heterochromatin were noticed. The myelin
figure may be resulting from the dissociation of lipoproteins
phosphatidic groups, which stimulate the incorporation and
intercalation of water between the lamellar stacks of the
membranes.[35]

After 24 h postinoculation with Bv venom, the adrenal cortex
showed a disseminated damage of the capillary vascular
system, with a MVB presence. Hemorrhagic metalloproteinases
and toxic PLA_{2} of Bv venom are highly toxic hemorrhagic
proteases, degrading endothelia and cleaving plasma proteins,
in a relatively specific and nonspecific manner.[3,36]

At 24 h, mitochondrial damage was still evident, with
disappearance of cristae and SER. The lesions observed
at 3 h were maintained and aggravated after 24 h. This
enzymatic cocktail of Bv venom proteases together with the
phospholipases was undoubtedly the cause of these subcellular
alterations. The (ER) has a crucial role in preserving cellular
homeostasis, and the MVB development occurs during
ER stress, which is induced by the inhibition of ER stress
transducers inositol, which requires Inositol-requiring
enzyme 1 (IRE1) and PKR-like ER kinase (PERK).[37]

Once the qualitative alterations were established, these results
lead us to conclude that there was a statistically significant
difference, among the treatments since the value of \( \alpha \) was
less than the significance (\( \alpha \)) specified. These values were
unlikely to be fortuitous but rather were alterations caused by
the Bv venom treatment. It was important to demonstrate that
these alterations at quantitative level allowed us obtaining
information at the different periods studied.

The quantitative analysis showed the morphometric analyses,
which was carried out in 100 randomly selected electron
micrographs, from each adrenal gland cortex, corroborating
whether the action of the venoms had statistical significance.

This quantitative study of PCA, under Bv venom activity on
thickening of the capillary endothelium, capillary lumen,
nucleus area, thickening of nuclear envelope, number
of mitochondria, area of the mitochondria, number
of mitochondrial cristae, and diameter of tubular elements
of the SER were established. The application of the previous
method showed good reproducibility and can be used for the
demonstration of alterations of these subcellular structures by
the venom action.

The current study demonstrated that Bv venom, at 6 and
24 h [Table 1], significantly induces an increase in the
thickening of the capillary endothelium. It is known that the
endothelial cell synthesizes supplies and releases different
molecules accomplishing autocrine, paracrine, and endocrine
functions. The endothelial membrane has different receptors
for physical, chemical, hormonal, and immunological signals,
which integrate it into the psychoneuroimmunoendocrine
complex. Therefore, the endothelium thickening can produces
an alteration, both in the excretion and in its capacity of
receptors mentioned above. The highest exposure of the
endothelia is probably due to its high dissemination throughout
the vascular system of the adrenal gland. Thus, venom activity
on various ligands could induce a variety of physiopathological
and toxicological reactions, causing several ostensible adrenal
toxicities. PLA_{2}s from Bothrops venoms are the leading
constituents, liable for cellular damage during the hydrolysis
of membrane phospholipids.[38] These abnormalities observed
in the capillaries, such as thickening of the endothelium and
the presence of prolongations toward its lumen, constitute
alterations that occur during bothropic envenomation, and
whose results have been reported by a number of authors.[39,40]

The present study also showed at 6 and 24 h [Table 1] that
Bv venom induced an increase in the capillary lumen. The
relative capillary lumen widening was the first morphologic
change observed. Despite the fact that relative capillary area
is decreased by the edema, the true capillary lumen was increased.
This may be produced by the elevation of blood pressure effect
in a blocked capillary when a new flow of blood is streaming
from the artery (which could be caused by coagulant proteins
of venom).[31]

This work showed that at 6 h, Bv venom induced a discrete
decrease in the nucleus area [Table 1]. However, at 24 h, the
venom caused a significant nuclear decrease. The processes of
achievement of Bothrops venom nucleus toxicity are not yet
explained. Nevertheless, this decrease in nuclear area has been
defined as a cellular response to lesions made by pathogens, in
a process known as hypofunction, which consists of a decrease in metabolic activity, and occurs in cases of atrophy. Many venom snake toxins are able to stimulate the production of free radicals, contributing to the inflammatory activities. These mediators are closely associated to the oxidative stress. Nevertheless, nuclear damage by the apoptosis generation may not be discarded. Many authors have described the apoptotic action of different classes of toxins, for instance, metalloproteases and PLAs.

This study showed that at 6 h, \( Bv \) venom induced a minor increase in the thickening of the nuclear envelope [Table 1]. However, at 24 h, the increase was higher. The nuclear envelope is constituted of an external nuclear membrane that is continuous to the ER and the pore components. It is structurally and functionally different of the inner nuclear membrane. Thus, it is formed of specific integral membrane proteins and has a clear connection with the underlying lamina. Once again, it is thought that the thickening of this membrane could be caused by the action of the proteases and the phospholipases present in the \( Bv \) venom. Thus, revisions on the mechanism of venom action indicated that the majority of the detected toxic effects possibly result from specific actions of this venom, on the target cell membrane and breaking its integrity, which induces instabilities in cell homeostasis.

The present study also demonstrated that the time of venom exposition (24 h) decreases the average number of mitochondria per cell [Table 1]. Mitochondria frequently change their morphology since they are extremely dynamic organelles. The changes occur to adequate the bioenergetics necessities of the cell, particularly in a state of environmental or metabolic pressures. However, they do not only change their morphology; additionally, they decrease their number under stress situations. In general, the mitochondrial compartment dynamic forces are organized by two contrasting routes, such as mitochondrial fusion and fission. Both increase the production of energy Adenosine triphosphate (ATP) but decrease the number of mitochondria.

Several studies have shown that various toxicological factors inhibit protein synthesis (e.g., cycloheximide), which can induce a high mitochondrial fusion, termed as stress-induced mitochondrial hyperfusion. The mitochondria once fused are further metabolically competent and can deal with augmented energy request through stress circumstances. This persistent stress eventually directs to mitochondrial disintegration and to apoptosis. Both mitochondrial fusion and fission processes can be affected by proteolysis and posttranslational modifications, caused in the current situation by the \( Bv \) venom proteases. Finally, autophagy, which also decreases the number of mitochondria, is essential for preserving a healthy mitochondrial segment, by removal of old and damaged mitochondria.

\( Bv \) venom-induced modifications in the morphology of mitochondria and the mitochondrial matrix come to be cleared due to the swelling. The mitochondrial area was increased at 6 and 24 h postinjection of venom. The normal membrane structure is the requirement for retaining mitochondrial oxidative phosphorylation and ATP formation. Mitochondrial swelling dysfunction directs to reduced oxidative phosphorylation and increased permeability. In the adrenal gland, this could cause a release of proapoptotic factors from the mitochondria, which triggers events for mitochondrial apoptotic signaling. These ultrastructural adrenal cortex changes have been described in previous studies caused by the toxin action of \( A. m. \) venom. Other micrograph examinations in cells of the cortex adrenal revealed a destruction of mitochondrial cristae exposed to \( Bv \) venom. The number of cristae sensibly decreased as early as 6 h of venom treatment and extended progressively until 24 h of experiment; its persistence evidences a general incursion of the intracellular membrane systems. After 6 h of venom injection, the number of mitochondrial cristae began to decrease, observing an electron density of the mitochondrial matrices, which can initiate a process of autophagy. Possibly, this deterioration can be due to a cytotoxic Viperidae snake venom effects, which could explain, why these alterations were observed in the case of \( Bv \) venom. There have been exceptions, particularly in rapidly metabolically active tissues, in which the mitochondria can be large or small and have or lack of cristae. These facts agree with the number of mitochondrial cristae per mitochondrion, and the number of the mitochondria per unit area was apparently unrelated. These mitochondrial alterations coincide with modifications reported in mouse hepatocytes, experimentally envenomed with rattlesnake venom. The manifestation of membrane-degrading enzymes such as venom phospholipases and proteases is advised. The alteration of mitochondrial cristae structure or number is closely related with severe mitochondrial dysfunction. The mitochondria display a fast prompt fall in cytochrome oxidase and a progressively lessening of succinic dehydrogenase. Mitochondria generate more than 90% of the energy required to maintain organ function. When they break down, less energy is produced inside the cell; it is damaged and can still die.

Examining the cisternae of the SER diameters, a light increase was observed at 3 h from venom injection, at 6 h, it was maintained, but at 24 h, an intense increment in the cisternae diameter was noticed. Given the large presence of electro dense lipids observed in the results, we can assume that there was disorganization in the activity of the SER, and based on this assumption, a more detailed study must be focused. SER covers approximately 80% of the adrenal gland and also participates in the process of conversion of pregnenolone to corticosteroids, another reason for the importance of its observation during toxin venom accident. Lloyd et al. indicated that the swelling of the cisternae of the ER was not necessarily related to the increased activity. It should be remembered that the toxins present in the snake venoms could imply a process of final mitochondrial degeneration, as was seen previously in the mitochondrial alterations; therefore, this would compromise the production of energy (ATP), adapting
the active transport of ions from the cell membrane and from the sodium–potassium pump; the impact of this is the entry of sodium and water into the cell.[34]

**Conclusion**

Our experimental work showed a diverse range of effects caused by the *Bv* venom activities. The *Bv* venom acted on subcellular organelles changing the morphology or number of capillaries, endothelia, nucleus, mitochondria, and SER. It appears that these changes were principally caused by the venom proteases and phospholipases and also other components of the venom such as peptides, several enzymes, nonenzymatic toxins, and minerals. All these results advise that adrenal cortex lesions may be significant in the etiopathogenesis of *Bv* snake envenoming. To our knowledge, this is the first report on ultrastructural adrenal gland damages caused by *Bothrops* venom.

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**Conflicts of interest**

There are no conflicts of interest.

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