**Naja nigricollis** venom altered reproductive and neurological functions via modulation of pro-inflammatory cytokines and oxidative damage in male rats

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**ABSTRACT**

Reproductive and neurological anomalies are often characterized by malfunctioning of reproductive and nervous organs sometimes attributed to systemic toxins. However, limited information is available on the impact of snake venom toxins on male reproductive and nervous system. This study investigated the toxicological effects of *Naja nigricollis* venom on male reproductive and neural functions in rat model. Twenty male Wistar rats weighing between 195 and 230 g were divided randomly into two groups of ten rats each. Group 1 served as normal control while rats in group 2 were envenomed with a single intraperitoneal injection of 0.25 mg/kg (LD$_{12.5}$) of *N. nigricollis* venom on first and twenty fifth day within the period of fifty days experiment. The venom significantly decreased sperm counts, motile cells and volume combined with increased sperm abnormalities. The venom induced hormonal imbalances in the envenomed group as levels of testosterone, luteinizing and follicle stimulating hormones depreciated compared to the control. Oxidative stress biomarkers: malondialdehyde significantly increased parallels with depletion of glutathione level and catalase activities in testis, epididymis and brain of envenomed rats. Furthermore, *N. nigricollis* venom up-regulated tumor necrosis factor-alpha (TNF-α) and interleukin1-beta (IL-1β) production in testis, epididymis and brain of envenomed rats compared to the control. Also, various histological alterations were noticed in tissues of testis, epididymis and brain of envenomed rats. Findings indicated that *N. nigricollis* venom is capable of inducing reproductive and neurological dysfunction in envenomed victims.

1. Introduction

Venomous snakes are species that produces venom in their venom glands and these group of snakes have been classified into diverse families and sub families [1]. However, report abounds that fatalities resulting from snake envenoming are largely due to bites from species that belongs to Viperidae (Vipers) and Elapidae (Elapids) families [1]. The predatory behaviours of vipers and elapids differs, the former typically ambush their prey by injecting an haemotoxic venom which contains mainly large proteins that majorly causes haemostatic disturbances, while the latter where cobras belongs are active predators with venoms predominantly composed of neurotoxic and cytotoxic proteins and peptides that typically attacks the central nervous system [2].

Cobras are known to inflict multiple-organ failure after venom injection which often results to death in case of severe envenomation [3]. Symptoms that manifests after envenoming by cobra are local pain, severe swelling, blistering, necrosis and other non-specific effects [4]. Biochemically, cobra venom are mixtures of proteins, peptides, enzymes, cytotoxins, coagulants and neurotoxins exhibiting toxic actions in living systems [5]. The neurotoxins in cobra venom blocks the transmission of nerve impulses to the muscles consequently affecting the nervous system, heart and lungs thereby leading to organ failure.

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malignantly and ultimately death as a result of complete failure of respiratory function [6].

*Naja nigricollis* commonly called black-necked spitting cobra is a venomous snake species belonging to a widely distributed snake family known as the Elapidae [7]. The specie is one of the most medically important elapids and their envenoming have resulted to quite a number of morbidity and mortality among rural dwellers in Nigeria [8]. Incidentally, *N. nigricollis* is believed to be the most urbanized specie, as many victims of their bites resides in urban areas in Africa [9].

The only treatment of snake envenoming is the use of serum anti-venom which are ineffective against local tissue damage and cannot reverse the damages to organ systems [10]. Also, most rural dwellers use traditional methods for snake bite treatment due to inaccessibility to a cooling container at maintained temperature of 4°C. For these reasons, not only do the antivenoms and some of these treatments may only ameliorates and not effectively and completely halt further actions of venom toxins on vital organs of the body. Furthermore, most antivenoms do not effectively neutralize snake venom due to toxins variations as they are produced using the snake species of a different location and imported to a different country for use. As a result of this, snake envenomed victims may encounter other physiological damages to their organ system functions due to the venom toxins which may not be fatal. Consequently, there is a paramount need to investigate the effect of venom toxins at tissue level of important reproductive and nervous organs of human body after envenoming.

Investigators of toxicology had since launch researches on numerous effects of snake venoms in cells, tissues and organ systems of animals with outstanding reports [11–13] but limited information is available on their reproductive and neural toxicity. Although, snake venom toxins from *Daboia russelli* have been documented to induced atrophy in testis of mice [14] while *Crotalus durissus* ssp. (rattlesnake) venom could cause an increase in sperm morphology abnormality [15]. Also, toxins present in *Bothrops jararaca* have been reported to inhibit spermatozoa production [16]. Despite the available information, the impact of *N. nigricollis* envenoming on male reproductive and neural functions in biological system remain unreported. Whereas, diverse pathophysiology attributed to venom toxins of *N. nigricollis* venom have been studied and information on their toxicities in different biological systems are available [1,17,18]. Hence, the present study was aimed to unravel the potential damage caused by *N. nigricollis* envenoming on male reproductive and nervous organs in male albino rats.

2. Materials and methods

2.1. Chemicals and kits

The rats enzyme-Linked Immunosorbent Assay (ELISA) Kits for Testosterone, FSH and LH assays were procured from Bio-Intec Diag nostic Ltd., Ken House, London, UK while enzyme-linked immunosorbent assay (ELISA) kits manufactured by PeproTech, Inc. London, UK were used for TNF-a and IL-1β assays. Other reagents and chemicals were of good grade and purchased from Sigma-Aldrich, Inc, USA.

2.2. Venom collection

A lyophilized venom of *N. nigricollis* was obtained from herpetarium of the Department of Veterinary Physiology and Pharmacology, Amadu Bello University, Zaria, Nigeria. The venom sample was transported via a cooling container at maintained temperature of 4°C to the Animal Physiology Laboratory, Department of Zoology, University of Ibadan, Nigeria. The lyophilized venom was stored at 4°C until required.

2.3. Experimental animals

Male albino Wistar rats (weighing between 195 and 230 g; 14 weeks old; n = 10) were procured from the Animal House of the Department of Zoology, University of Ibadan, Ibadan, Nigeria. The animals were placed in a transparent plastic cages on a 12 h light/dark cycle and allowed free access to standard pelleted rat chow and tap water *ad libitum*. The experimental protocol was conducted in accordance with the regulations of the local ethics committee in animal care and use in Research of the University of Ibadan, Nigeria. The animal experiment complied with the National Research Council’s publication on guide for the care and use of laboratory animals [19].

2.3.1. Experimental groups

The rats were randomly distributed into two groups of ten rats (n = 10) each. Group 1 were injected with 0.2 ml of saline and served as normal control while group 2 were envenomed with 0.2 ml of *N. nigricollis* venom and served as the venom group.

2.3.2. Envenoming procedures

The lethal dose concentration (LD₅₀) of *N. nigricollis* venom based on our previous study was 1.0 mg [17]. In this current study, rats were envenomed with a single intraperitoneal injection of 0.25 mg/kg⁻¹ (LD₁2.₅) of *N. nigricollis* after the venom dose was reconstituted in 10 ml of normal saline. The animals were envenomed with 0.2 ml of the venom on Day 1 with a repeated dose on Day 25 of the experiment at 8:00 a.m. each day. All rats were observed for a period of 50 days post envenomation and mortalities were recorded. The exposure duration of 50 days was selected based on the completion of rat spermatogenesis [20]. Animals were monitored for clinical signs of toxicity including food and water consumption behavioral patterns and mortality.

2.3.3. Determination of body weight gain

The body weight of animals was measured before venom injection on day 1 and prior to day 50 before they were sacrificed as terminal weight. The body weight gain was calculated using the formula:

\[
\text{Body weight gain} = \frac{\text{Terminal weight of rats} - \text{Initial weight of rats}}{\text{Initial weight of rats}} \times 100.
\]

2.3.4. Blood sample and organs collection

At post-envenomation, blood from the experimental rats was collected using heparinised capillary tubes into plain bottles through retro-orbital sinus punctuation and centrifuged at 380 g for 10 min to obtain serum for hormonal assays. Immediately after blood collection, the rats were sacrificed by cervical dislocation following the protocol of Rowett, [21]. The cauda epididymis was surgically removed and processed for sperm parameters analysis. The testes and brain were also removed and weighed. Parts of tissues of the epididymids, testes and brain were used for biochemical assays while the remaining parts were preserved in Bouin’s fluid and 10% formalin respectively before been processed for histological analysis. The relative testes and brain weight were determined using the formula:

\[
\text{Relative organ weight} = \frac{\text{Organ weight}}{\text{Terminal body weight}} \times 100.
\]

2.4. Determination of sperm parameters

2.4.1. Sperm counts

For sperm count, a 1:10 serial dilution of the sperm suspension was used with the aid of an improved Neubauer hemocytometer using the protocol of previously described [22]. After dilution, exactly 100 μl of sperm suspension was loaded onto the hemocytometer to count the spermatozoa under a light microscope at a 400 × magnification. Sperm count values were multiplied by the dilution factor and recorded as millions per milliliter (10⁹/ml). To evaluate the sperm volume, the epididymis was immersed in 5 ml normal saline in a measuring cylinder and the volume displaced was taken as the volume of the epididymis [23].
2.4.2. Sperm motility

The cauda epididymis was placed individually in a Petri dish and minced in normal saline (1 ml) to form the sperm suspension, and 10 μl were placed in triplicate on microscopic slides and observed for motility under the light microscope at a magnification of x400. Sperm motility was assessed by classifying 200 spermatozoa into two categories, motile and immotile spermatozoa. Three sperm classes were categorised as motile spermatozoa: rapid progressive, slow progressive and non-progressive spermatozoa [24].

2.4.3. Sperm morphology

The assessment of sperm morphological abnormalities was carried out following the protocol of Wyrobek et al. [25]. 450 μL of the sperm suspension was mixed with 50 μL of 1% aqueous eosin Y for 10 min. The stained sperm suspension was used to make a thin smear on a pre-cleaned grease-free microscopic slide. Prepared slides were allowed to air dry and abnormalities were observed in 250 spermatozoa with four replicates in each rat at a magnification of × 1000.

2.5. Hormonal assays

The levels of free sera sex hormones were measured for testosterone, luteinizing hormone (LH) and follicle stimulating hormone (FSH) using Bio-Inteco Enzyme-Linked Immunosorbent Assay (ELISA) Kit (Inteco Diagnostic, UK). The optical density was taken at 450 nm with a microplate reader.

2.6. Antioxidant parameters in testis, epididymis and brain tissues

Lipid peroxidation end product; malondialdehyde (MDA), was measured as thiobarbituric acid reactive substance (TBARS) as described [26]. Catalase activity (CAT) was measured as described [27], while levels of reduced glutathione (GSH) were assayed according to the method of Ellman [28].

2.7. Measurement of pro-inflammatory cytokines production in testis, epididymis and brain tissues

2.7.1. Tissue preparation

Frozen testis, epididymis and brain tissue samples were homogenized in a 1.5 ml RIPA buffer (25 mM TrisHCl, 150 mM NaCl, 1% NP-40, 1% sodium deoxycholate, 0.1% SDS pH = 7.6) supplemented with Protease inhibitors at 4 °C. The homogenate was incubated on ice for 30 min and then centrifuged at 10,000×g for 30 min at 4 °C. Following centrifugation, the supernatants were transferred to labelled Eppendorf and stored at −80 °C for cytokine measurement.

2.7.2. Measurement of tumor necrosis factor-alpha (TNF-α) and Interleukin1-Beta (IL-1β) production

Quantitative measurement of the level of cytokines was performed using Mini Enzyme Linked Immunosorbent Assay (ELISA) Development Kits (Peprotech). Well plates were set up according to the manufacturer’s instructions and read using an ELISA plate reader at 405 nm with 650 nm as the correction wavelength. Concentrations (pg/mL) of testis, epididymis and brain TNF-α and IL-1β were estimated [29].

2.8. Histopathological evaluation

The testis, epididymis and brain tissues from the control and envenomed groups were examined for histological studies using conventional techniques of paraffin-wax sectioning and haematoxylin-eosin staining [30].

2.9. Data analysis

Data were presented as mean ± Standard Error of mean (SEM) and analyzed using T-Test to compare the significant (P < 0.05) differences between the experimental and control group. Independent Sample test was used for comparison. All statistical analyses were done using GraphPad Prism version 7.0.

3. Results

3.1. Clinical signs of toxicity

Rats in the envenomed group showed some clinical signs of toxicity including: dizziness, low appetite for food with increase in water consumption and mortality. Mortalities were not observed in the normal control rats while only one death occurred on 20th day of the experiment in the envenomed group (Table 1). The body weight gain of normal control rats was significantly (P < 0.05) higher compared to the envenomed rats. Testicular and brain weights of envenomed rats significantly (P < 0.05) decreased compared to the control rats while the same trend was observed in the organo-somatic index of the experimental rats (Fig. 1).

3.1.1. The effects of N. nigricollis venom on sperm parameters

N. nigricollis envenomation induced a significant decrease (p < 0.05) in the percentage of motile spermatozoa compared to the control rats. Consequently, a significant increase (p < 0.05) in immotile spermatozoa was noticed in the envenomed rats compared to the control (Table 2). Sperm volume and count was significantly (p < 0.05) decreased in envenomed rats compared to the control (Table 2).

3.1.2. The effects of N. nigricollis venom on sperm morphology of envenomed rats

The percentage of sperm abnormalities showed a significant (p < 0.05) increase in the envenomed rats compared to the control. However, the sperm abnormalities as revealed in control rats have no unique abnormal spermatozoa compared to the sperm abnormalities noticed in envenomed rats (Table 3). High sperm abnormalities caused by the venom as noticed were short hook, banana shape and amorphous head respectively while unique sperm abnormalities observed were double tail, double head and Long and sickled hook (Table 3).

3.2. The effects of N. nigricollis venom on reproductive hormones

The serum concentration of luteinizing hormone (LH) and testosterone (TEST) of envenomed rats significantly (P < 0.05) reduced compared to the control. However, follicle stimulating hormones (FSH) of normal control was statistically (P < 0.05) not different compared to envenomed group however, there was a marked reduction in FSH levels of envenomed group compared to control (Table 4).

3.3. Effects of N. nigricollis venom on catalase enzyme activities of testis, epididymis and brain tissues

The activities of catalase (CAT) in epididymis and brain tissues of the control were significantly (P < 0.05) not different compared to envenomed rats, however, a marked decrease in CAT activities was observed in envenomed rats compared to the control. Whereas, the venom

Table 1

| Groups    | Envenomation | Day 1 | Day 20 | Day 50 | Mortality (%) |
|-----------|--------------|------|--------|--------|---------------|
| Control   | –            | –    | –      | –      | 0.00          |
| Envenomed | –            | 1    | –      | –      | 10.00         |

Number of rats per group (n = 10). Data are represented as mean ± SE (n = 5). Values in the same column with different superscript are considered significant (p < 0.05).
Table 2 
Effects of *N. nigricollis* venom on sperm parameters of envenomed rats.

| Groups          | Sperm Motility (%) | Sperm Volume (mL) | Sperm Count(10⁶/mL) |
|-----------------|-------------------|------------------|---------------------|
|                 | Motile            | Immotile         |                     |
| Control         | 79.06 ± 1.55      | 20.94 ± 1.06*    | 8.28 ± 1.09*        |
| Envenomed       | 10.55 ± 1.00      | 89.45 ± 2.06b    | 4.59 ± 0.72b        |

Data are expressed as MEAN ± S.E, (n = 5). Values in the same column with different superscript are considered significant (p < 0.05).

Table 3 
Sperm morphological abnormality counts in rats envenomed with *N. nigricollis* venom.

| Sperm Parameters | Control | Envenomed |
|------------------|---------|-----------|
| Amorphous head (AM) | 5.67 ± 1.45* | 57.85 ± 0.79b |
| Banana shape (BS) | 6.42 ± 1.32b | 45.23 ± 1.41b |
| Double tails (DT) | 0.00 ± 0.00a | 29.10 ± 1.14a |
| Folded Sperm (FS) | 1.24 ± 0.54a | 43.90 ± 1.02b |
| Knobbed head (KH) | 2.67 ± 0.63a | 34.37 ± 1.00b |
| Long and sickled hook (LSH) | 0.00 ± 0.00a | 27.52 ± 0.50a |
| Double head (DH) | 0.00 ± 0.00a | 27.25 ± 0.70a |
| Short hook (SH) | 7.02 ± 1.14a | 47.32 ± 3.67b |
| Wrong tail attachment (WTA) | 3.62 ± 0.70a | 36.53 ± 1.67b |
| Pin head (PH) | 4.18 ± 0.87a | 28.42 ± 1.14a |
| No hook (NH) | 5.56 ± 1.00a | 54.22 ± 1.30b |
| Wrong-angled hook (WAH) | 3.37 ± 0.48b | 30.97 ± 1.16b |
| Total abnormal cells | 39.75 ± 2.30a | 462.68 ± 4.60b |
| Percentage abnormalities | 3.98 ± 1.33a | 46.27 ± 1.41b |

Data are represented as mean ± SE (n = 5). Mean ± S.E in the same column having similar superscript are not significantly different at P < 0.05. Mean ± S.E are fractions of the 1000 sperm cells assessed.

Table 4 
Levels of the reproductive hormones in the blood of the rats envenomed with *N. nigricollis* venom.

| Groups          | LH (ng/ml) | FSH (ng/ml) | TEST (ng/ml) |
|-----------------|------------|-------------|--------------|
| Control         | 50.33 ± 2.02* | 455.33 ± 49.46* | 2.33 ± 0.10* |
| Envenomed       | 42.33 ± 0.88b | 336.33 ± 54.81b | 1.25 ± 0.20b |

Data are expressed as MEAN ± S.E, (n = 5). Values in the same column with different superscript are considered significant (p < 0.05).

LH: Luteinizing Hormone, FSH: Follicle Stimulating Hormone, TEST: Testosterone Hormone.
compared to the control (Fig. 5).

3.4. Effects of N. nigricollis venom on tumor necrosis factor-alpha (TNF-α) production in envenomed rats

The interleukin1-Beta (IL-1β) responses attributed to the venom action in epididymis and brain tissues was significantly (P < 0.05) not different to control, although the responses was more intense in envenomed group compared to control. Whereas, IL-1β production in the testis tissues of the envenomed rats significantly (P < 0.05) increase compared to the control (Fig. 5).

3.4.1. Effects of N. nigricollis venom on Interleukin1-Beta (IL-1β) production in envenomed rats

Production of tumor necrosis factor-alpha (TNF-α) as a result of venom injection significantly (P < 0.05) increased in the epididymis and brain tissues compared the control however, response to TNF-α in testis tissues of envenomed rats was statistically (P < 0.05) not different to the control although, TNF-α production was higher in the envenomed rats (Fig. 6).

3.5. Histopathological assessment of the testis, epididymis and brain tissues

The examinations of the testis, epididymis and brain revealed various histological defects. The tests of control rats showed no observable lesion as several uniformly sized, closely packed seminiferous tubules with regular outlines were observed and contained countless spermatogenic cells with spermatocytes and round spermatids indicating a healthy reproductive status in males. However, the tests of the envenomed group revealed a tubular atrophy, depletion of spermatogenic cells, distorted germinal epithelium and necrosis of the spermatogenic cells (Fig. 7). On the other hand, lesion was not observed in the epididymis of control rats however, a prominent mucosal epithelium with little luminal content was noticed in epididymis of envenomed group (Fig. 8). Furthermore, a fused neuronal degeneration, aseiaemic neuronal necrosis and gliosis was observed in brain tissues of envenomed rats while there was no observable lesion in brain tissues of the control (Fig. 9).

4. Discussion

Snake venom toxins are known to initiate wide spectrum of biological activities and cause different metabolic disorders by altering cellular inclusions and enzymatic activities of different organs and systems of the body following snakebite envenoming [31,32]. This current study unraveled reproductive and neural toxicities due to N. nigricollis envenoming. The post envenoming behaviour of the envenomed animals, weight loss due to loss of appetite and resultant to a mortality may be attributed to toxins present in the venom as earlier observed by Adeyi et al. [17]. Findings from this study showed that the venom toxins induced significant decrease in testicular weight and organo-somatic index of the envenomed rat which could arise from decrease in the number of epithelial spermatogenic cells nesting the seminiferous tubules, through testicular ischemia and necrosis, leading to loss of testicular weight [33]. Significant decrease in relative testicular weight in this study is an evidence of testicular toxicity as earlier observed in other animal studies due to the effects of toxicants [33,34].

Cells of the body are composed of membrane receptors which are sensitive to the action of snake venoms but often targeted by venom components. Sperm cells are types targeted by animal venoms and sensitivity of sperm to venom components has been reported as their cells contain a notorious rich repertoire of ion channels and membrane receptors, making them a priori susceptible interesting targets for the
action of venom components [35,36]. Consequently, the observed significant reduction in motile sperm, count, and volume in envenomed rats can be attributed to the action of *N. nigricollis* venom toxins through initiating impairment of spermatogenesis in the testes. The toxic enzymes present in *N. nigricollis* venom are cytotoxic majorly phospholipase A2 (PLA2), an enzyme that hydrolyzes the sn-2 fatty acid acyl ester bond of phosphoglycerides to free fatty acid and lysophospholipids specifically with plasma membranes and catalyzes the breakdown of phospholipids within their structure [2,37]. Therefore, PLA2 breakdowns plasma membranes of sperms and contributes to decreased sperm counts and motility. Interestingly, inhibition of male sperm motility and acrosome reaction by action of venom phospholipases A2 have been

Fig. 7. Histological examination of the testis Group 1 (Control): No lesion, Group 2 (Envenomed): Tubular atrophy, distortion of germinal epithelium and necrosis of the spermatogenic cells. HE x400.

HE x400.

Fig. 8. Histological examination of the epididymis Group 1 (Control): No lesion, Group 2 (Envenomed): The mucosal epithelium are prominent with little luminal content.

HE x400.

Fig. 9. Histological examination of the brain Group 1 (Control): No lesion, Group 2 (Envenomed): Ischaemic neuronal necrosis and gliosis.
It is known that sperm motility is a key requirement for fertilization to occur and cases of infertility are attributed to defective immotile sperm or altered acrosome reaction arising from sperm cells abnormalities [39]. In this study, a significant increase in abnormal sperm cells induced by the venom was noticed which is in tandem with previous studies [15]. Although, the mechanism of sperm abnormality induction was not yet ascertained but this may be due to excessive free radicals particularly Hydroxyl (OH·) generated by the venom toxins interacting with the DNA in the sperm heads. Also, OH· generated by the venom may interact with biomolecules such as the DNA, proteins and lipids to induce oxidative damage [40] however, further investigation is required to ascertain the exact mechanism.

The administration of N. nigricollis venom-induced harmful reproductive effects on hormone profiles as evidenced by the decrease in levels of testosterone and FSH which are essential hormones for activating the process of spermatogenesis. FSH binds to Sertoli cells and stimulates testicular fluid production while LH is produced by gonadotrophic cells to stimulate Leydig cells to produce testosterone which in turn plays an important role in the development of male reproductive tissues such as testis [41,42]. The decline in the concentration of testosterone post-venom injection indicates harmful changes in the Leydig interstitial cells of testes, which are responsible for testosterone biosynthesis and secretion [43]. Furthermore, the reduction in sperm count as earlier noticed could be attributed to the decline in the concentration of testosterone through induction of Leydig cell damage that disrupts testosterone synthesis.

Oxidative stress may be a result of excessive reactive oxygen species generation or failure of the cellular antioxidant system [11]. Reports have documented that increased concentration of polyunsaturated fatty acids (docosahexaenoic acid) and low expression levels of antioxidants make spermatozoa highly susceptible to ROS attacks resulting in lipid peroxidation [44]. In this current study, N. nigricollis venom toxins caused up-regulation of lipid peroxidation levels in testis, epididymis and brain tissues. However, there was low responses in levels of GSH and catalase activity in testis, epididymis and brain tissues indicating the venom caused an elevation of oxidative stress indicators such as lipid peroxidation and damaged the oxidant system. Glutathione plays an important role in system detoxification process and found mainly in the cell cytosol as they are known to be cell’s natural antioxidant that neutralizes free radicals formed in cells [11] while catalase subsequently helps in detoxification of H2O2 to water as protective mechanism against lipid peroxidation [45]. Malfunctioning of the oxidant system as observed in this study aligned with previous interpretation of the consequences GSH deficiency in the body system as a result of oxidative damage and elevated lipid peroxidation which in turn leading to cell damage [11,46,47].

Pro-inflammatory cytokines are produced by testicular somatic and spermatogenic cells, both under normal conditions to control immune cell function within the testis, stimulate and maintain spermatogenesis or in response to inflammatory stimuli [48]. Also, excess production of pro-inflammatory cytokines may result in physiological damages as they are known to provoke inflammation and infections responses thereby contributing to transformation from acute to chronic inflammation and development of autoimmune reaction [49]. In this study, IL1 and TNF produced in reproductive organs of the envenomed rats was more evident compared to control an indication of possible reproductive organs inflammation due to cytokines imbalances which may contribute to infertility [50]. Up-regulation of pro-inflammatory cytokines by N. nigricollis venom as observed in this study aligned with our previous studies using cobra venom [50].

The observed reduction in sperm parameters was consistent with the histopathological examination of the testis and epididymis that showed significant harmful tissues changes with tubular atrophy and distorted germinal epithelium occurring in testicular tissue while a prominent mucosal epithelium with little luminal content was noticed in epididymal tissue which could possibly be due to venom induced oxidative stress. In addition, degenerated reproductive organ tissue ascribed to snake venom toxins have been documented [14].

Access of toxins to the nervous tissues enhances the possibility of a direct neurotoxic effect as neurotoxicity ultimately depends on the ability of the substance to bind to neural tissue targets and interfere with functional or structural integrity [51]. The assessment of the venom toxicities on the brain of envenomed rats was based on mechanism of action exhibited by N. nigricollis venom as neurotoxins in cobra venom are known to attack the central nervous system [6], disrupt its functions, facilitates nervous dysfunction and shutdown resulting to tetany and ultimately death.

The mammalian nervous system has functional design features that predispose it to chemical perturbation. Consequently, neurological dysfunction is among the most common of the toxic responses of humans to toxicants and brain is a major target organ for some toxins [51]. This may inform the observed increase in oxidative stress markers resulting from excessive generation of reactive oxygen species (ROS) and up-regulation of pro-inflammatory cytokines production as revealed in the brain tissues of envenomed rats. In addition, toxins in N. nigricollis venom induced significant decrease in the brain weight and brain index compared to the control an indication the toxins have a direct interaction with the brain tissues as evident with various histological alterations noticed on brain tissues of envenomed rats which was in consonance with findings from our previous studies on cobra venom [50]. The alterations noticed on histological and biochemical composition of brain tissues after envenomation is an indication of brain toxicity due to direct effects of the venom toxins on the brain. This may occur as a result of damage to the blood brain barrier and consequently disrupt the functions of the nervous system.

It should be noted that reproductive functions are ultimately regulated by the hypothalamus-pituitary-gonadal (HPG) axis of the nervous system mainly the brain [52]. The role of the brain as the primary regulator of neuroendocrine and reproductive function through brain’s responsiveness to hormonal feedback [53] makes it an important organ to consider when studying reproductive and nervous system-disrupting toxicants. The hypothalamus, located at the base of the brain, is especially vital for regulation of endocrine function as it serves as an interface between the nervous and endocrine systems [53] including the reproductive system. Consequently, alterations of hormone levels along with effects on the reproductive system caused by exposure to toxicants could manifest due to disruptive structure and function of the brain. However, the mechanism of N. nigricollis venom toxins altering the reproductive and neurological functions in the envenomed rats may not yet be ascertained in this present study, but results obtained is worthy of attention and provides caution to effectively treat snakebite envenomed victims using a standard antivenom to avoid systemic toxicity which may manifest post-envenoming if proper treatment is not administered.

5. Conclusion

This study demonstrated that N. nigricollis venom is capable of exhibiting serious toxic effects on nervous and male reproductive system as the venom alters biochemical compositions of the brain, decreased reproductive hormones levels and increased sperm abnormalities combined with reduced motile sperm and count. These observed effects or in combination are major factors reportedly responsible for nervous and reproductive dysfunctions which will have an important implication to public health most especially infertility challenges. The information obtained from this study may provide a better knowledge of other clinical implications of ineffective treatment of snakebite envenomed patients most especially in rural areas and further drive focus on development and implementation of a new therapeutic strategy for the treatment of snakebite envenoming.
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