Effect of Ethanol Extract of *Moringa oleifera* Leaf on Nissl Granules, Immunohistochemistry of Inferior Colliculus and Oxidative System Following Quinine Toxicity in Wistar Rats

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Abstract: The metabolite of Quinine, 3-hydroxyquinine is the active substance and more toxic than quinine hence, it can exert pro-oxidant effect in the body. In this study, the effect of ethanol leaf extract of *Moringa oleifera* was evaluated against quinine toxicity on nissl granules, immunohistochemistry of inferior colliculus and oxidative system of Wistar rats. Fifty Wistar rats weighing 180 – 200g were allotted into 10 groups of 5 rats each. Group 1 served as control while Groups 2 to 10 were the treatment groups. Groups 2 – 4 received 10, 20 and 30 mg/kg body weight of quinine hypochloride injection intramuscularly eight hourly for 7 days. Groups 5 – 7 were orally administered 250, 500 and 750 mg/kg body weight of *M. oleifera* leaf extract daily for 7 days respectively. Group 8 received 250mg/kg of leaf extract orally and intramuscular (IM) 10mg/kg of quinine; Group 9 was treated with oral 500mg/kg of the extract and IM 20mg/kg of quinine while Group 10 received 750mg/kg of the extract orally and IM 30mg/kg of quinine for 7 days. Results revealed that quinine significantly (p < 0.05) decreased the activities of anti-oxidative enzymes (superoxide dismutase and catalase) and increased lipid peroxidation (malondialdehyde, MDA) in treated rats. However, *Moringa oleifera* leaf extract showed significant increase in the activities of anti-oxidant enzymes and decreased lipid peroxidation. Histomorphological and immunohistochemical observations revealed that quinine caused neuronal distortions, nuclear degenerations, and depletion of Nissl granules, vascular congestions and vacuolations and increased astrogliosis in the inferior colliculus. Incorporation of the extract to the quinine treated groups revealed neuronal regenerations, increased staining intensity and density of the Nissl granules, reduced astrogliosis and restoration in the inferior colliculuscyto-architectures. In conclusion, *Moringa oleifera* leaf extract can be used to ameliorate the adverse effects of quinine toxicity on inferior colliculus.

Keywords: Quinine, *Moringa oleifera*, Inferior Colliculus, Nissl Granules, Oxidative Enzymes

1. Introduction

Non-tolerability and non-compliance of patients to medications may be as a result of the adverse effects of the drugs on them [1]. This could also be one of the causes that may result in drug resistance by the offending organisms to the drugs. Adverse effects of drugs could be caused by reactive oxygen species which can lead to oxidative damage of body tissues [2, 1]. Quinine is recommended to be used as an antimalarial drug especially when artesunate is unavailable [3]. It is safe even in the first trimester of pregnancy [4]. Although quinine occurs naturally in the bark
of the cinchona tree, it has been synthesized in the laboratory [4]. Quinine is rapidly absorbed both orally and parenterally, reaching peak concentration within 1 - 3 hours. It is distributed throughout the body fluid and is highly protein bound mainly to alpha (α) -1 acid glycoprotein and the binding capacity in plasma is concentration dependent, and the levels of alpha-1 acid glycoprotein [4]. Quinine is incorporated in minute quantities as a flavor component in tonic water and bitter lemon. It is used in treating lupus erythematosus and arthritis. In addition to its poor tolerability and poor compliance with complex dose regimes [5], quinine has several setbacks. It is known for its ototoxicity and neurotoxicity [6]. The side effects commonly encountered at therapeutic concentration and referred to as cinchonism include mild form of tinnitus, slight impairment of hearing, headache and nausea. More marked adverse effects are vertigo, vomiting, abdominal pains, marked auditory loss and visual symptoms including loss of vision [4]. Permanent blindness with standard dose of quinine has been documented [3].

In recent years, several plant extracts and other natural products have been tested for their antioxidant properties. Furthermore, the use of antioxidant supplements can reverse or minimize the oxidative damage to host caused by the free radicals released by the use of antimalarial drugs [7]. Most of plants used for medicinal purposes have been correlated to their possession of antioxidant activity [8, 9]. Antioxidants quench, scavenge and suppress the formation of reactive oxygen species and free radicals or oppose their actions [10-12]. *Moringa oleifera* is an economically important tree and vegetable. Preliminary evidence suggest that it has an antioxidant and anti-inflammatory potency [7, 13]. *Moringa oleifera* is an herbal plant with immense medicinal value because of its antioxidant properties [14, 15]. The plant is also a rich source of vitamins and antioxidants. It contains good amount of proteins, minerals, vitamin A, vitamin B complex, essential amino acids and high content of vitamin E [16].

In view of the antioxidant properties of *Moringa oleifera* leaf and its richness in minerals and vitamins, this research was aimed at examining its effect on nissl substances, histomorphology of inferior colliculus, some oxidative enzymes, lipid peroxidation following quinine toxicity in adult Wistar rats.

### 2. Materials and Method

#### 2.1. Animal Care and Protocols

Fifty (50) male Wistar rats weighing 180 – 200g used in this study were obtained from the Animal House of the Faculty of Basic Medical Science (FBMS), University of Uyo, Uyo, Nigeria. Approval for the study was obtained from the Faculty Animal Care and Use Research and Ethical Committee and the research was strictly conducted according to the international guidelines for the care and use of Laboratory animals of the Faculty Animal Care and Use Committee. The animals were randomly grouped into 10 (ten) groups with 5 rats in each group and were fed with growers mash (Pfizer Nigeria Ltd) and allowed drinking water *ad libitum*.

#### 2.2. Preparation of Plant Material and Drug

The *Moringa oleifera* leaf samples of the same species and varieties were collected from a local farm in Uyo, Akwa Ibom State, washed and transported under hygienic condition. The leaves were identified and authenticated by a taxonomist in the Department of Botany and Ecological Studies, Faculty of Science, University of Uyo, Uyo with the Voucher No. A (50) i. *Moringa oleifera* leaf extract concentrate was prepared by ethanol extraction using Percolation Method described by United States Pharmacopoeia Convention Inc (2000). The leaves were carefully washed in clean water and dried under shade for about 7 days. The air-dried leaves were powdered and macerated in 70% ethanol. After mixing thoroughly in a suitable round bottom flask for 72 hours at room temperature, the extract was filtered with Whatman filter paper size No. 1 and the filtrate was then concentrated in water bath to dryness at 40°C. The concentrated extract was preserved in refrigerator at the temperature of -4°C until commencement of the research.

#### 2.3. Determination of Acute Toxicity and Drug Preparation

Acute toxicity testing [lethal dose (LD₅₀)] of the extract was determined using modified Lorke’s method (1983) [17], in two stages using Swiss albino mice, age 6 to 8 weeks. The mice were obtained from the animal house of the Faculty of Basic Medical Science (FBMS), University of Uyo, Uyo. The extract was administered orally. In the first stage, the animals (mice) received 1000, 2000, 3000, 4000, and 5000mg/kg of the ethanol extract of *moringa oleifera* leaves. In the second stage, the animals (mice) received 2300, 2400, 2500, 2600, and 2700mg/kg of the ethanolic extract of *moringa oleifera* leaves. All the experimental mice were observed for physical signs of acute toxicity such as decreased respiration, limbed body and death for 12 hours and the number of animals that died within 24 hours recorded. According to modified Lorke’s method [17], the (LD₅₀) was calculated which was 2500mg/kg.

The stock solution was prepared and calculated by dissolving 2500mg/kg of the extract in 20mls of distilled water to get the stock solution of 125mg/ml. The therapeutic dose was calculated using 10% of the lethal dose as 250mg/kg body weight. The volume of the stock solution used was calculated using Tedong equation

$$V = \frac{(DXP)}{C}$$

Where;

- **D** = group dose used (mg/kg)
- **P** = animal body weight (kg)
- **C** = Conc. of the stock solution
- **V** = volume of the experimental drug (ml)

Injectable quinine hydrochloride was obtained from
The experimental animals were sorted randomly into ten groups (Groups 1 – 10) with five rats in each Group. Group 1 was taken as the control group and was administered with distilled water. Groups 2, 3 and 4 were given 10 mg/kg, 20 mg/kg and 30 mg/kg of quinine hypochloride injections intramuscularly respectively eight (8) hourly for seven days. Groups 5, 6 and 7 were administered 250, 500 and 750 mg/kg of extract of *M. oleifera* leaf orally, respectively for 7 days. Groups 8, 9 and 10 received concomitantly, 250 mg/kg of *M. oleifera* leaf extract daily and 10 mg/kg of quinine 8 hourly, 500 mg/kg of *M. oleifera* leaf extract daily and 20 mg/kg body weight of quinine 8hourly and 750 mg/kg of *M. oleifera* leaf extract daily and 30 mg/kg of quinine 8 hourly for 7 days respectively. The administration of quinine and ethanol leaf extract of *M. oleifera* lasted for 7 days.

At the end of the experiment, the animals were sacrificed under chloroform inhalational anaesthesia then the blood samples collected by intracardiac puncture for assay of antioxidant enzymes activity while the experimental animals were perfused with phosphate buffer solution (PBS) and the brain tissues harvested, processed and stained for histological studies. A section of the brain was homogenized and the brain homogenate used for evaluation of indices of lipid peroxidation.

### 2.4. Experimental Design

The experimental animals were sorted randomly into ten groups (Groups 1 – 10) with five rats in each Group. Group 1 was taken as the control group and was administered with distilled water. Groups 2, 3 and 4 were given 10 mg/kg, 20 mg/kg and 30 mg/kg of quinine hypochloride injections intramuscularly respectively eight (8) hourly for seven days. Groups 5, 6 and 7 were administered 250, 500 and 750 mg/kg of extract of *M. oleifera* leaf orally, respectively for 7 days. Groups 8, 9 and 10 received concomitantly, 250 mg/kg of *M. oleifera* leaf extract daily and 10 mg/kg of quinine 8 hourly, 500 mg/kg of *M. oleifera* leaf extract daily and 20 mg/kg body weight of quinine 8hourly and 750 mg/kg of *M. oleifera* leaf extract daily and 30 mg/kg of quinine 8 hourly for 7 days respectively. The administration of quinine and ethanol leaf extract of *M. oleifera* lasted for 7 days.

At the end of the experiment, the animals were sacrificed under chloroform inhalational anaesthesia then the blood samples collected by intracardiac puncture for assay of antioxidant enzymes activity while the experimental animals were perfused with phosphate buffer solution (PBS) and the brain tissues harvested, processed and stained for histological studies. A section of the brain was homogenized and the brain homogenate used for evaluation of indices of lipid peroxidation.

### 2.5. Determination of Antioxidant Enzyme Activity and Lipid Peroxidation

Quantitative determination of whole-cell catalase activity was accomplished using a protocols described by Herbert [18] and Beers and Sizer [19]. SOD activity was assayed according to the method by Kakkar, Das and Viswanathan, [20]. Thiobarbituric acid reactive substance, the last product in lipid peroxidation pathways was estimated by the methods of Nichaus and Samuelsson [21].

### 2.6. Histopathological Analysis

The experimental animals were perfused with phosphate buffer solution (PBS); shortly after the perfusion, brain tissues were harvested and fixed immediately in neutral buffered formalin for histomorphological evaluation. Haematoxylin and Eosin method was used for general histologic evaluation, Cresyl Fast Violet Method [22] for Nissl Granules and GFAP Immunohistochemistry Protocol were used to evaluate the reactive astrocytes of the inferior colliculus of the experimental animal.

### 2.7. Statistical Analysis

The data obtained were subjected to analysis of variance (ANOVA) at 5% level of significance to establish the statistical differences between means of each group. Duncan’s New Multiple Range Test was used to compare the means as described by Steel and Torrie [23].

### 3. Results

#### 3.1. Effect of Quinine and Ethanol Leaf Extract of *Moringa oleifera* on the Histomorphology of Inferior Colliculus

The general histology of the inferior colliculus (H and E Method) is presented in Figure 1. Photomicrograph HE01 revealed Inferior colliculus of control group with normal area of neuronal cell body, and neuronal density. The nerve fibres are not thickened. Photomicrograph HE02, HE03 and HE04 represent Inferior colliculus of Group 2 (10 mg/kg bw of quinine), Group 3 (20 mg/kg body weight of quinine) and Group 4 (30 mg/kg body weight of quinine). Observed features include neuronal cell degeneration, reduced neuronal density and thickened nerve fibers. The histology of Groups 5 (250 mg/kg of *M. oleifera*), 6 (500 mg/kg of *M. oleifera*) and 7 (750 mg/kg of *M. oleifera*) are represented in photomicrograph HE05, HE06 and HE07. Normal neuronal cell, neuronal density and nerve fibres were observed. The photomicrograph of Group 8 (HE08) treated concomitantly with 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* revealed normal neuronal cell and neuronal density, the nerve fibres are not thickened. Photomicrograph HE09, the histology of Group 9 (20 mg/kg of quinine and 500 mg/kg of *M. oleifera* concomitantly), revealed slight neuronal cell degeneration, slight hypertrophy of neuronal cells and slightly thickened irregular nerve fibers and reduced neuronal density. The histology of Group 10 (HE10) treated concomitantly with 30 mg/kg of quinine and 750 mg/kg of *M. oleifera* showed features such as slight neuronal cell degeneration, thickened irregular nerve fibers and reduced neuronal density.

#### 3.2. Effect of Quinine and Ethanol Leaf Extract of *Moringa oleifera* on the Immunohistochemistry of Inferior Colliculus

The results of the immunohistochemistry studies on the inferior colliculus in the present study, using CFV and GFAP staining methods are presented in Figures 2 and 3 respectively. The photomicrograph labelled NS01 representing Group 1 (control) revealed neuron cell body, with prominent nucleus and densely packed nissl granules within normal cellular cytoarchitecture. The inferior colliculus of Groups 2 (10 mg/kg body weight of quinine – NS02), 3 (20 mg/kg body weight of quinine – NS03) and 4 (30 mg/kg body weight of quinine – NS03) revealed cell body shrinkage of neurons, Pyknotic Nucleus shrinkage, Karyolitic Nucleus and Nissl granules depletion. Administration of ethanol leaf extract of *M. oleifera* to Groups 5 (250 mg/kg body weight of *M. oleifera* – NS05), 6 (500 mg/kg body weight – NS06) and 7 (750 mg/kg body weight – NS07) showed Inferior colliculus with cell body of...
neurons, containing Nucleus and densely packed Nissl granules within cytoplasmic content. Concomitant administration of 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* in group 8 (NS08) revealed Inferior colliculus with cell body of neurons, containing Nucleus and reduced concentration of densely packed Nissl granules within cytoplasmic content. However, the cytoarchitecture of Group 9 labelled NS09 (20 mg/kg of quinine and 500 mg/kg of *M. oleifera*) revealed Inferior colliculus with cell body shrinkage of neurons, containing Karyolitic Nucleus and nissl granules depletion within cytoplasmic content. Similarly, the cytoarchitecture of Group 10 labelled NS10 (30 mg/kg of quinine and 750 mg/kg of *M. oleifera*) revealed Inferior colliculus with cell body shrinkage of neurons, containing Karyolitic Nucleus and nissl granules depletion within cytoplasmic content.

Photomicrographs AS01 – AS10 represent the immunohistochemistry (GFAP Method) of the inferior colliculus of experimental animals in Group 1 – 10 respectively and they show astrocytes reactivity. Normal cellular architecture and no astrocytes reactivity were observed in the photomicrograph of the control group labelled AS01. Administration of 10, 20 and 30 mg/kg bw of quinine to Groups 2, 3 and 4 respectively revealed severe reactive astrocytes (AS) with thick process. Vacuolation in the stroma (arrows) was also observed in Group 4. Administration of 250, 500 and 750 mg/kgbw of *M. oleifera* to Groups 6, 7 and 8 respectively revealed Inferior colliculus with normal cellular profile. Normal cellular profile was also observed in Group 8 following treatment with 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* concomitantly. The inferior colliculus of Group 9 treated concomitantly with 20 mg/kg of quinine and 500 mg/kg of *M. oleifera* showed reactive astrocytes and neuronal cell. Astrocytes with thick processes were observed in the inferior colliculus of animals treated concomitantly with 30 mg/kg of quinine and 750 mg/kg of *M. oleifera*.

### 3.3. Effect of Ethanol Leaf Extract of Moringa oleifera and Quinine on Oxidative Stress Indices

The effects of ethanol leaf extract of *Moringa oleifera* and quinine on the MDA level in adult Wistar rats is presented in Figure 4. The results showed that rats in Groups 2, 3 and 4 which were treated with 10mg/kg, 20mg/kg and 30mg/kg of quinine respectively had significant increase (p < 0.01) in MDA level compared with control group. *Moringa oleifera* leaf extract inhibited the amount of MDA generated (and thus lipid peroxidation) significantly (p < 0.01) as observed in Groups 5, 6 and 7 rats. Thus, the decrease in the MDA level on administration of the leaf extract indicates the role of the extracts as an antioxidant. The results revealed MDA level was high in Groups 2, 3 and 4 on administration of 10, 20 and 30 mg/kg of quinine respectively but low in Groups 5, 6 and 7 on administration of 250, 500 and 750 mg/kg of *M. oleifera* leaf extract respectively. The MDA level in Group 8 (10 mg/kg of quinine and 250 mg/kg of *M. oleifera* leaf extract concomitantly) revealed no significant difference (p > 0.01) when compared with control group. The MDA level in the group treated with 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* leaf extract concomitantly significantly decreased (p < 0.01) compared with Group 2 which received 10 mg/kg of quinine. Groups 9 (20 mg/kg of quinine and 500 mg/kg of *M. oleifera* leaf extract concomitantly) and Group 10 (30 mg/kg of quinine and 750 mg/kg of *M. oleifera* leaf extract concomitantly) showed a significant increased (p < 0.01) in MDA level when compared with control. Comparing Groups 9 and 10 with Groups 3 (20 mg/kg of quinine) and 4 (30 mg/kg of quinine) respectively showed a significant decrease (p < 0.01) in MDA level in Groups 9 and 10.

The effects of ethanol extract of *M. oleifera* leaf extract and quinine on antioxidant enzymes are shown in Figures 5 and 6. The results revealed that quinine significantly (p < 0.01) decreased superoxide dismutase (SOD) and catalase (CAT) activities in Groups 2, 3 and 4 rats treated with 10, 20 and 30 mg/kg of quinine respectively when compared to the control group. All doses of ethanol extract of *M. oleifera* leaf significantly increased SOD and CAT activities when compared with the control group (p < 0.01) The antioxidant enzymatic activities (SOD and CAT) were low in Groups 2, 3 and 4 on administration of quinine as observed but high in Groups 5, 6 and 7 on administration of *Moringa oleifera* leaf extracts. There was no significant difference (p > 0.05) in SOD and CAT activities of Group 8 when compared with the control group but significantly increased (p < 0.01) compared with Group 2. The SOD and CAT activities in Group 2 were significant decreased (p<0.01) when compared with the control group. There was significant increase in the activity of SOD and CAT in Groups 9 and 10 when compared with Groups 3 and 4 respectively.
Figure 1. Photomicrographs of the inferior colliculus of experimental animals showing neuronal cells and cellular cyto-architecture following the administration of quinine and ethanol leaf extract of Moringa oleifera. Neuronal cell (NC); Nerve Fibres (NF); Neuronal Cell Degeneration (NCD). (H and E Method, Mag X400).

Photomicrographs HE01 – HE10 represent histology of Groups 1 – 10 respectively. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of *M. oleifera*; Group 6 – 500 mg/kg of *M. oleifera*; Group 7 – 750 mg/kg of *M. oleifera*; Group 8 – 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of *M. oleifera* concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of *M. oleifera* concomitantly.
Figure 2. Photomicrographs of the inferior colliculus of experimental animals showing nissl substances and cellular cytoarchitecture following the administration of quinine and ethanol extract of Moringa oleifera. Pyknotic Nucleus shrinkage (pN); Karyolitic Nucleus (kN); Nissl granules depletion (Ngd); neuron cell body (CB); Nissl granules (Ng); cell body shrinkage (CBs); Nucleus (N). (CFV Histological Method, Mag X400).

Photomicrographs NS01 – NS10 represent histology of Groups 1 – 10 respectively. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of M. oleifera; Group 6 – 500 mg/kg of M. oleifera; Group 7 – 750 mg/kg of M. oleifera; Group 8 – 10 mg/kg of quinine and 250 mg/kg of M. oleifera concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of M. oleifera concomitantly.
Figure 3. Photomicrographs of the Immunohistochemistry of the inferior colliculus of experimental animals showing astrocyte reactivity following the administration of quinine and ethanol extract of Moringa oleifera. Astrocytes (AS); neuronal cell (NC). (GFAP Immunohistochemistry Method; Mag. X400).

Photomicrographs AS01 – AS10 represent histology of Groups 1 – 10 respectively. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of M. oleifera; Group 6 – 500 mg/kg of M. oleifera; Group 7 – 750 mg/kg of M. oleifera; Group 8 – 10 mg/kg of quinine and 250 mg/kg of M. oleifera concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of M. oleifera concomitantly.

Figure 4. The effect of ethanol leaf extract of M. oleifera and quinine on lipid peroxidation (LPO) of albino Wistar rats. Values presented Mean± SEM. *=significantly different when compared to control group at p < 0.01. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of M. oleifera; Group 6 – 500 mg/kg of M. oleifera; Group 7 – 750 mg/kg of M. oleifera; Group 8 – 10 mg/kg of quinine and 250 mg/kg of M. oleifera concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of M. oleifera concomitantly.
Figure 5. The effect of ethanol leaf extract of M. oleifera and quinine on peroxide dismutase activity (SOD) of albino Wistar rats. Values presented Mean±SEM. *=significantly different when compared to control group at p < 0.01. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of M. oleifera; Group 6 – 500 mg/kg of M. oleifera; Group 7 – 750 mg/kg of M. oleifera; Group 8 – 10 mg/kg of quinine and 250 mg/kg of M. oleifera concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of M. oleifera concomitantly.

Figure 6. The effect of ethanol leaf extract of M. oleifera and quinine on catalase activity (CAT) of albino Wistar rats. Values presented Mean±SEM. *=significantly different when compared to control group at p < 0.01. Group 1 – Control; Group 2 – 10 mg/kg of quinine; Group 3 – 20 mg/kg of quinine; Group 4 – 30 mg/kg of quinine; Group 5 – 250 mg/kg of M. oleifera; Group 6 – 500 mg/kg of M. oleifera; Group 7 – 750 mg/kg of M. oleifera; Group 8 – 10 mg/kg of quinine and 250 mg/kg of M. oleifera concomitantly; Group 9 – 20 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly; Group 10 – 30 mg/kg of quinine and 750 mg/kg of M. oleifera concomitantly.

4. Discussion

Medicinal plants can be used in different form, as raw materials for the extraction of active compounds or for the extraction of abundant but inactive constituents which can be transformed by partial synthesis into active compounds, as drugs or extracts or traditional preparations [16]. Medicinal Plants are therapeutics resource much used by the population of the world specifically for health care [24]. World-wide interest in the use of medicinal and aromatic plants is increasing. Beneficial effects of plant based medicines and other plant based products are being rediscovered. Different parts of Moringa oleifera have been used for the treatment of inflammation, infectious diseases, cardiovascular, gastrointestinal, haematological and hepatorenal disorders [25, 26]. Quinine is an alkaloid derived from the bark of the cinchona tree that is used in the treatment of malaria. Several toxic effect have been observed following quinine administration and they include tinnitus, slight impairment of hearing, headache and nausea. The metabolite of quinine (3-hydroxyquine) is of higher toxicity and it is also likely that 3-hydroxyquinine contributes to adverse effects, the toxicity observed or felt during quinine administration [27]. Based on the aforementioned, the ameliorative effect of ethanol leaf extract of Moringa oleifera on the oxidative stress and inferior colliculus of albino rats intoxicated with quinine was evaluated in the present study.

The present study demonstrated that quinine exhibited a deleterious effect on the inferior colliculus of rats as observed in quinine treated groups (Groups 2, 3 and 4). From the result, the changes of astrocytic phenotype were mild, moderate and
severe as shown in Groups 2, 3 and 4 treated with 10, 20 and 30 mg/kg bw of quinine respectively. The changes are based on the capacity of matured cells to undergo a switch in response to an insult and it essentially characterized by up-regulation of intermediate filament protein [28]. Quinine might actually be the key to initiate internal process that optimizes neuronal cells and conditions them to prepare for damaging effect. In this study, quinine could have acted as toxins to the cells of the inferior colliculus, affecting their cellular integrity and causing a defect in membrane permeability and cell volume homeostasis. Quinine is known to cross membranes by simple diffusion, thus getting access to the cells [4]. This property of quinine could have been one of the causes of the degenerative changes observed in this study. In cellular necrosis, the rate of progression depends on the severity of environmental insults.

The prime candidates for inducing the massive cell destruction observed in neuro-degeneration are neurotoxins [11]. These may be substances present in small amounts in the environment, or even naturally occurring chemicals such as glutamate used by the brain as transmitter substances [13]. The latter, when present at a critical level, can be toxic to the brain cells they normally excite [13, 29]. The decrease in neuronal density of quinine treated group as reported in this study may have been as a result of cell death caused by the toxic effect of quinine [4]. A study by Ajibade et al., [30] corroborates the observed neuronal degeneration in association with loss of Nissl bodies and reduced staining intensity of the nissl substances in the inferior colliculus of quinine treated rats in the present study. Ajibade et al., [30] had reported similar cyto-architectural changes in the cerebral cortex following the administration of quinine to adult Wistar rats. Furthermore, loss of Nissl substance and shrinkage of the nucleus in dogs have been reported following intramuscular administration of artemether [31]. Injury to axons or neuronal exhaustion, resulting from strong or prolonged stimuli, causes a reduction in the number of Nissl bodies. This alteration, which is called chromatolysis, occurs simultaneously with nuclear migration to the periphery of the perikaryon [13], and consequently the RNA level is reduced. Chemical and toxic substances affect the Nissl substance thereby influencing their metabolic activity [13]. Similarly, it has been reported that neuronal degeneration causes a reduction in Nissl bodies [1]. Neuronal degeneration in quinine-treated rats has already been reported by [30].

Consequently, reduced staining intensity of the Nissl substance in the inferior colliculus of the treated rats may be due to neuronal pathology, following the neurotoxic effect of quinine. It is possible that quinine had an irreversible effect on the Nissl substances of the inferior colliculus, bringing about such microstructural changes in the neurons which manifested as degeneration and loss of Nissl substances with reduced staining intensity of the Nissl substances in the inferior colliculus of the quinine-treated rats. Degeneration and loss of Nissl substances may consequently affect the synthesis of both structural protein and protein for transport in correlation with neuronal functions. The reduced neurogenesis and increased cell death can all cause or be the result of neuro-degenerative changes [2].

However the GFAP immunoreactivity in quinine treated group suggests that quinine showed earlier astrocytes response. Quinine induced degeneration in the inferior colliculus result in diffused astrogliosis. Astroglial persist throughout the period of quinine administration. Thus astrocytes reactivity was up-regulated during quinine administration. Astrocytes reactivity is heterogeneous with differential phenotypes dependent upon the inducing stimulus [32]. Although quinine specific mechanism of action resulting in demyelination is still unclear, quinine has been shown to induce a stress response in oligodendrocytes [32].

*Moringa oleifera* leaf extract did not produce any histologically observed deleterious effect on the inferior colliculus. A positive effect of *M. oleifera* leaf extract on other brain tissues such as cerebral cortex has been reported [33]. The polyphenolic compound of the leaf extracts might have played a vital role. It has been reported that the plant polyphenols provide protection against neurodegenerative changes [34]. Based on the effect of polyphenolic compound just mentioned, it was also possible that the neuroprotective of the plant extract was associated with these compounds.

In the present study, the effect of quinine and *Moringa oleifera* leaf extract concurrent administration on the inferior colliculus of rats in the group administered 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* leaf extract concomitantly revealed normal features. This is an indication that the structural damage that occur in the inferior colliculus of rats which received 10 mg/kg of quinine was absent following administration of the extract. This points to the evidence that *M. oleifera* was able to restore to normal, the structural damage caused by low dose of quinine on the inferior colliculus. Anti-toxicity effect of *M. oleifera* leaf extract on cerebral cortex has been reported [35]. *M. oleifera* leaf extracts exhibited the ability to provide neurite protection, in other words it protects against structural damage. This is reflected by preservation to or close to normal in terms of morphological structures as observed in 10 mg/kg of quinine and 250 mg/kg of *M. oleifera* concomitantly treated group. The presence of neurogenic agent such as *M. oleifera* leaf extracts could ameliorate the process by providing a positive stimulation while damping the neurodegenerative effect of quinine and prevent erratic uncoordinated stimulation.

Observations in Groups 9 (20 mg/kg of quinine and 500 mg/kg of *M. oleifera* concomitantly) and 10 (30 mg/kg of quinine and 750 mg/kg of *M. oleifera* concomitantly) revealed that structural damage occur on the inferior colliculus of rats in these groups. Comparison between treated groups revealed normal inferior colliculus morphology in Group 8 (10 mg/kg of quinine and 250 mg/kg of *M. oleifera* concomitantly) compared with Group 2 (10 mg/kg of quinine) which showed degenerative changes. Inferior colliculus of animals in Group 3 (20 mg/kg of quinine) revealed a moderate degeneration compared with Group 9 (20 mg/kg of quinine and 500 mg/kg of *M. oleifera* concomitantly) which showed mild degenerative changes.
Inferior colliculus of animals in Group 4 (30 mg/kg of quinine) showed a severe neurodegeneration compared with Group 10 (30 mg/kg of quinine and 500 mg/kg of M. oleifera concomitantly) which revealed a mild neurodegeneration. These are indications of the neuroprotective effect of Moringa oleifera leaf extract, thus damping the neurodegenerative effect of quinine dose dependent.

Reactive oxygen species (ROS) are related to oxidative stress and many scientific reports have shown that excessive production of reactive oxygen species (ROS) can further aggravate oxidative stress and have complicated ROS in number of disorder and disease processes [36, 37]. The equilibrium between the reactive oxygen stress (ROS) and antioxidant enzymes including superoxide dismutase (SOD) and catalase (CAT) are crucial and could be an important mechanism for preventing damage by oxidative stress. Antioxidants appear to act against disease processes by increasing the level of endogenous antioxidant enzyme such as SOD and CAT and decreasing toxic byproducts of lipid peroxidation such as malondialdehyde [38]. Malondialdehyde (MDA) is one of the end products of polyunsaturated fatty acid peroxidation and it is a good indicator of the degree of lipid peroxidation which is related to quinine-induced tissue damage [12].

The reduction in SOD and CAT activities with increased in the level of MDA in the pathogenesis of quinine intoxication as observed in Group 2 (10 mg/kg of quinine), 3 (20 mg/kg of quinine), 9 (20 mg/kg of quinine and 500 mg/kg of Moringa oleifera concomitantly) and 10 (30 mg/kg of quinine and 750 mg/kg of Moringa oleifera concomitantly) indicate oxidative stress. The decreased enzymatic activity in turn enhanced the excess reactive oxygen species (ROS) resulting in increased in oxidative damage reflected in the elevation of MDA leading to neurodegeneration. The increase in MDA level could be due to a reduction in antioxidant defense. A decrease in SOD production can be attributed to enhanced superoxide generation and utilization of this enzyme during reactive metabolites detoxification. The antioxidant enzyme SOD is the first line of defense against reactive oxygen species, converting them to toxic hydrogen peroxide in living tissue. Catalase (CAT) further carries out the detoxification of hydrogen peroxide to molecules of oxygen and water. Similar changes in the activities of antioxidant enzymes have been reported by Verma et al., [39] and Wu et al., [11] following administration of drugs.

Moringa oleifera leaf extracts enhanced SOD and CAT activity and decreased MDA level. This is consistent with recent report that M. oleifera increased the antioxidant enzymatic activities and decreased lipid peroxidation in Swiss albino mice [31]. These suggest that the extract may act as radical chain terminator and transforms free radical species to stable non-reactive product [39]. The highly protective effect of Moringa oleifera against neurotoxicity induced drugs has previously been studied [34]. This points to a conclusive evidence that the protective effect of M. oleifera leaf extract on SOD and MDA concentration may be attributed to the presence of phyto constituent particularly flavonoids and polyphenols that scavenge free radicals, activate the antioxidant enzymes and inhibit oxidation [33]. Based on these results, it may be suggested that the therapeutic potentials of M. oleifera leaf extract are dependent on the antioxidant mechanism.

5. Conclusion

In conclusion, the present study has shown that quinine induced neurodegeneration manifesting as altered cytoarchitecture of inferior colliculus, depletion of nissl granules and increased astrocytes reactivity in the experimental animal. Quinine dose dependent toxicity of the inferior colliculus may be through the mechanism of free radicals generation leading to oxidative stress. The product of lipid peroxidation increased in a dose dependent manner coupled with decreased activity of the antioxidant enzymes with quinine administration. M. oleifera leaf extract however, showed a protective effect against quinine induced neuronal toxicity in albino Wistar rats. Restoration of normal cytoarchitecture of the inferior colliculus was observed following the administration of M. oleifera. Ethanol leaf extract of M. oleifera further increased the activities of superoxide dismutase and catalase while inhibiting the generation of free radicals as reflected in the reduced concentration of the product of lipid peroxidation. The administration of high doses of quinine should be discouraged and Moringa oleifera may be administered as an adjuvant during quinine therapy to ameliorate its toxicity.

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