Quercetin Attenuates Manganese-Induced Neurotoxicity in Mice Hippocampus by Regulation of Oxidative Stress and Tumour Necrosis factor Alpha Expression

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Abstract

The hippocampus is involved in learning and memory processes, an integral component of cognitive function. The aim of this study is to assess the efficacy of quercetin on manganese-induced neurotoxicity in the hippocampus of the adult mice.

In this study, 40 adult mice of average weight of 18 –29g were randomly distributed into five groups of eight each. The brain was harvested and the region of the hippocampus was grossed for histological and immunohistochemical analysis.

The results revealed a significant increased level of oxidative stress markers of manganese treated mice when compared with the normal control and quercetin treated animals (p<0.05). Immunohistochemical analysis also showed a decrease expression of Tumour necrosis factor alpha (TNFα) with quercetin treated animals when compared with manganese treated animals indicating its neuroprotective function. In addition, quercetin treated animals all had an improved working spatial memory in Y-maze test. The histological results also revealed a degeneration of pyramidal cells with a characteristic pyknotic activities at the granular layer of the hippocampus leading to neuronal integrity damage following chronic exposure to manganese but normal architectural design was however maintained with quercetin.

Conclusively, exposure to manganese in excess may have adverse effect on extensive neuronal degeneration that could affect the learning, memory and possibly spatial navigation ability of mice and quercetin attenuates this induced neurotoxicity via inhibition of oxidative stress and reduction of TNF expression.
Introduction

Neurotoxicity refers to a kind of toxicity by which different agents such as biological, chemical, or physical could cause an adverse effect on the anatomy of the central and peripheral neurons [1]. It could be as a result of exposure to a substance known as neurotoxin to alter the normal structure of the nervous system and this might result into damage to nervous tissue which could perhaps destroy neurons [1]. Exposure to other heavy metals and chemicals such as pesticides in the environment may lead to neurotoxicity [2]. Neurodegeneration is the continuous process of losing the structure or function of neurons which can lead to death of neurons and these have a way to affect person’s movement and cognitive function [3]. Neurodegenerative diseases are complex type of diseases that involves neurodegeneration in different areas of the brain. These diseases are characterized by a progressive degeneration of the both structure and function of the central nervous system. Most therapy adopted for neurodegenerative diseases such as Parkinson’s diseases and Alzheimer’s disease have been reported not to be completely efficient in the management of the disorders thereby leading into a state of progressive degeneration and death of neurons.

Manganese is an essential trace element present naturally in foods such soybeans, leafy vegetables, coffee and available as a dietary supplement and also a cofactor for many enzymes [4, 5]. Evidence have shown that 1-100mcg/L of water contains small amounts of manganese [6]. Evidence has not shown that high dietary manganese intakes could cause manganese toxicity [7]. However, manganese toxicity has been reported in an individual working as welders and miners or those exposed to high amounts of manganese as result of chronic inhalation of manganese dust [8]. Manganese toxicity is known to affects the central nervous system resulting into impaired fine motor and cognitive functions especially those related to memory and learning [8]. However, the neural mechanism of alteration in hippocampus by manganese to affect cognition is still unclear, hence the reason for the study to understand the efficacy of quercetin on manganese-induced neurotoxicity.

Quercetin is grouped as one of the six subclasses of flavonoid compounds found in fruits and vegetables [9]. Evidences revealed that quercetin have a therapeutic potential for the prevention of different diseases such as cardiovascular disease and neurodegenerative diseases [9, 10]. However, quercetin has been shown to affect its antioxidant and anti-inflammatory properties in several areas of animal models as well as in humans by upregulating the signalling pathways via inhibition of oxidative stress markers and apoptotic factor expression involved [9, 10]. For this present study, quercetin has been used on animal model to investigate its antioxidant activities on the cell death induced by excessive consumption of the manganese.

Materials and Methods

Experimental Animals

40 adult male BALB/c mice weighing 18-29g were purchased from Tolex Animal holding, Ekiti State, Nigeria. Approval was sought and obtained from the ABUAD Ethical committee on animal use. The animals were housed in well-ventilated cages and maintained under standard laboratory condition of temperature, humidity and light. The animals were allowed to acclimatize for two weeks and fed with rodent meal (ADEHEZ Rodent Feed®, Ilorin, Nigeria) and drinking water was made available ad libitum. After the acclimatization period, they were reweighed and
randomly assigned to the five groups of eight (8) mice each. All animal experiment for the study complied with the ARRIVE guidelines [11] and was carried out in accordance National Institutes of Health guide for the care and use of Laboratory animals.

Other materials used for this study include: Quercetin were purchased from Medchem express, Manganese (II) chloride (MnCl$_2$·4H$_2$O) tetrahydrate, Sodium hydrogen Phospahte (Na$_2$HPO$_4$), Potassium chloride (KCL), Potassium dihydrogen phosphate (KH$_2$PO$_4$), Ketamine and all consumable used were purchased from Kermel laboratory (Colmar, France). The water used was double-distilled and other reagents were of analytical grade.

Preparation of Quercetin and Manganese
Quercetin was purchased from Medchem express while Manganese chloride was purchased from Kermel laboratory (Colmar, France). 10% DMSO was prepared by diluting 100ml of DMSO and 900ml of phosphate buffer solution (PBS). 400mg of Quercetin was then dissolved in 33.6ml of 10% dimethyl sulphur oxide (DMSO). The Manganese chloride was dissolved in distilled water;

Experimental Design
The mice were randomly divided into five (5) groups of 8 mice each. The first group represent the control group (C) that received normal saline for the duration of 14 days. The second group (Mn) received manganese (40mg/kg) for the 7 days [12, 13]. The third group (Q) received quercetin (50mg/kg) for 7 days. The fourth group (POST) was pre-treated with manganese (40mg/kg) followed by quercetin (50mg/kg) administration [14]. The fifth group (CONC) received both manganese and quercetin concurrently for 7 days. (Mn: 40mg/kg; Q:50mg/kg). All administration was done by intraperitoneal route.

Animal Sacrifices
The mice that were set out for histological analysis were sacrificed by anaesthetized using ketamine followed by transectional perfusion fixation (using 0.9% sterile saline followed by 4% paraformaldehyde) while the animals set out for biochemical analysis were sacrificed through cervical dislocation. The brain was harvested and the region of hippocampus was grossed for histological and immunohistochemical analysis.

Behavioural Analysis
The behavioural test was carried out with proper illumination and sound control in the behavioural room of Afe Babalola University. The behavioural test was conducted after the last administration. The behavioural test carried out were the Y-maze and the NOR (novel object recognition) test to assess learning and memory function, an integral component of cognitive behaviour in the experimental animals and the tests were recorded and properly documented.

Y-Maze Test
Y-maze test was carried out to investigate the spatial working memory of the animals based on the procedure used by Adeniyi et al., (15). The mice were placed in the middle of the Y-box and allowed their arm decision for 5 mins. The correct alternation was (ABC, ACB, BAC, BCA, CAB, CBA) between the arms (ABC) was video recorded and analysed later to determine the memory index which is the percentage of normal alternation. Animals with lower percentage alternation are likely to have lower spatial working memory index and vice versa.

Novel Object Recognition Test
The Novel object recognition test (NOR) was also carried out to evaluate cognition, particularly recognition memory and the test was done in a controlled arena. The objects used were similar in height and volume, but were different in appearance (colour). The frequency of times the animals recognized the new objects were video recorded with tally method.

Biochemical Analysis
Biochemical analysis was done using the whole brain tissue homogenate to assay the activities of malondialdehyde and superoxide dismutase to quantitatively evaluate oxidative stress.

Photomicrography
Microscopy was conducted with a 5.1-megapixel MV550 research microscope connected to a computer and images captured and analysed.

Statistical Analysis
Statistical analysis was done using one-way analysis of variance (ANOVA) and differences between
groups were evaluated using Newman-keuls for post hoc tests with the aid of GraphPad Prism V.5.0. The data statistical analysis was represented in graphs and bar charts with standard error of the mean (mean ± SEM) and significant level was set at *P< 0.05.

**Results**

**Morphological Changes**

There was a significant morphological change observed across the experimental groups. Quercetin (Q) and normal saline (C) treated mice revealed some level of positive changes in average body weight (26 to 32 g) but manganese treated mice experience a kind of decline in body weight (24 to 19g). The post treatment group (intervention) showed little or no changes in body weight. There has been no significant change in fur colour and colour of the eyes in all the groups.

**Biochemical Assay**

**Lipid Peroxidation (MDA) and Superoxide Dismutase (SOD)**

Statistical changes in the markers of oxidative stress in terms of Malondialdehyde (MDA) and SOD activities were observed (**P<0.001**) from figure 1 and 2. Manganese-treated mice showed a significant increase of Malondialdehyde activities (MDA), which is an evidence of oxidative stress and this was invariably decrease with the quercetin administration as shown in Figure (1) when compared with the control group. Also, Activities of superoxide dismutase were significantly increased by quercetin-treated mice when compared with other experimental groups (Figure 2).

**Behavioural Studies**

Memory index was assessed in the experimental animals using NOR and Y-maze test.

**Novel Object Recognition (NOR) Test and Y-Maze Test**

These tests were carried out to assess the spatial working memory of the animals. The memory index for NOR4 was calculated using: Memory Index (%) = (T Novel) / (T Novel + T Familiar) x 100% (Figure 3). Quercetin improve spatial working memory with an evidence of statistical increase in memory index during novel object recognition test of all experimental groups that received quercetin when compared with the manganese-treated and normal control group in Figure 3 (**P<0.05, ***P<0.01 and ****P<0.001**). The percentage alternation as indices for memory index in Y-Maze test was calculated as: % Alternation = No. of right decisions / No. of total arm entries-2 ×100. This was indicated by higher alternation for the quercetin treated mice compared to control mice, although, manganese treated mice showed a decrease percentage alternation but no statistically significant changes were observed (Figure 4).

**Histological Analysis**

The hippocampus is divided structurally into hippocampus proper (cornu ammonis) and dentate gyrus with 3 cell layers and a remarkable distribution of densely packed pyramidal and other cells involved in encoding, organizing and storing memories. From the histological analysis in figure 5, a distinct 3 marked layers of hippocampus with a normal distribution of pyramidal cells and a well stained nuclei were observed in normal control and quercetin treated group and the integrity of these neurons were also seen to be preserved around the dentate gyrus (DG) and cornu ammonis (CA1-C3) with their principal cell layers known as granular and pyramidal layer respectively. As shown in Figure 5 with group that received Mn, the layers of the hippocampus there were shown to have histological alteration (Mn-treated mice). The pyramidal layer of the hippocampus was observed with extensive vacuolations of the cells and pyknotic appearance as compared to the control group, thus indicating neuronal cell death in the CA1 and CA2 region than CA3. The cells of the CA4 region, the hilus of DG and granular cell layer of DG, were least affected while the group that were received quercetin after manganese administration (intervention group) revealed mild neuronal death and co-treated animals (concurrent groups) revealed a very minimal vacuolation in normal appearance of the 3 distinct layers of hippocampus and neuronal distributions.

**Immunohistochemistry (TNF-α)**

Immunohistochemical results of hippocampal (dentate gyrus) section evaluating the ameliorative roles of quercetin following manganese-induced neurotoxicity. Normal control and quercetin treated mice revealed little or no expression of tumour necrosis factor alpha (TNF-α) in the dentate gyrus and cornu
Figure 1. Concentration of MDA (µ/ml) in the brain tissue. There is a significant change observed when control and quercetin treated mice were compared with other experimental mice with manganese. Quercetin protects neurotoxicity by inhibition of markers of oxidative stress via reduction of lipid peroxidation. Control (C); Manganese (Mn); Quercetin (Q); Intervention (POST); Concurrent (CONC); Malondialdehyde (MDA) *P-value < 0.0001.

Figure 2. Concentration of SOD (µ/ml) in the brain tissue homogenate. Activities of superoxide dismutase were significantly increased by quercetin treated mice when compared with other experimental groups. Control (C); Manganese (Mn); Quercetin (Q); Intervention (POST); Concurrent (CONC); Superoxide Dismutase (SOD) **P<0.01, ***P< 0.001.
Figure 3. Memory index function with Novel object recognition test. Quercetin improve memory index when compared with other experimental groups. Control (C); Manganese (Mn); Quercetin (Q); Intervention (POST); Concurrent (CONC); *P<0.05, **P<0.001, ***P<0.0001.

Figure 4. Percentage alternation as an indices of memory index in Y-maze test. Percentage alternation changes were observed across the groups but not significant based on the group comparison. Control (C); Manganese (Mn); Quercetin (Q); Intervention (POST); Concurrent (CONC); ns: not significant.
Figure 5. Photomicrograph of the dentate gyrus (DG) section in control (C) group, manganese (Mn) group, Quercetin (Q) group, post treatment (PT) group and concurrent (PT) group. Stained with H&E. Mg x800
ammonis (CA1-CA3) but this was more pronounced in the Manganese-treated mice. However, other experimental animals that received quercetin recorded a minimal expression indicating the potential neuroprotective effect of quercetin during or after cell death. Fig 6.

Discussion

The present study demonstrated the efficacy of quercetin on manganese-induced neurotoxicity in mice hippocampus. The weights of all the animals were taken into the consideration before grouping and experimentation. The normal control and quercetin treated mice showed an increase in weight gain but manganese-treated mice expressed a decline in size and weight indicating that quercetin enhances feeding efficiency. The post treated animals clearly showed a little or no changes in weight and size. However, there were no significant changes in fur and eye colour in all the groups throughout the experiment and this was supported by research done by Porwal et al. [5, 29].

Oxidative stress increases the levels of reactive oxygen species (ROS) and give rise to lipid peroxidation that could have a deleterious effect on nervous tissues. An imbalance between oxidants and antioxidants form the underlying basis of oxidative stress leading to many pathophysiological conditions in the body, some of these include neurodegenerative diseases such as Parkinson's disease and Alzheimer's disease [6,11,12]. Neurodegenerative diseases can also occur as result of a faulty metabolic process leading to a release of ROS and lipid peroxidation that affect cell membrane and other organelles to operate normally in terms of function [9]. Also, excess manganese exposure in mice has been reported to induce neurotoxicity by generating reactive oxygen species [13, 14] and these free radicals cause oxidative stress that result in brain neuronal damage [10,14]. Malondialdehyde level is commonly known as a marker of oxidative stress. In the biochemical assay, malondialdehyde (MDA) concentration was done in this study to evaluate the level of lipid peroxidation as a marker of oxidative stress in cellular activities. Lipid peroxidation is considered as a major pathway of cell injuries in aerobic mechanism subjected to oxidative stress and the reaction sequences of lipid peroxidation increases the number of free radicals in the cells which leading to a cell death [10]. Significant increase in oxidative stress in all the mice treated with manganese were observed and vice versa with mice treated with quercetin either with intoxicant or quercetin alone (Figure 1). This make us to confirmed indeed that oxidative stress via lipid peroxidation may cause cellular and molecular damage with increases reactive oxygen species (ROS) which further leads to tissue destruction where fragmentation of DNA material from the nuclei are evident [12, 13, 15].

Behavioural studies carried out at the end of drug administration using the novel object recognition and Y-maze test showed that manganese significantly affects the performance of mice and intervention of quercetin improves the spatial memory. Animals administered with manganese alone showed significantly reduction in memory index with longer duration to recognize the novel object compared to familial object which might as a result of hippocampal damage. This indicates poorer learning and memory capacity (Figure 4). This supports earlier evidence that showed that excess Manganese and other neurotoxin negatively affects learning and memory by damaging hippocampal formation [16]. Results from Y-maze analysis (figure 4) also showed that with the normal control animals being the reference point, the quercetin treated animals had an improved spatial working memory in terms of percentage alternation, a function of memory index when compared with manganese-induced animals. This result implies that quercetin improve memory and learning while exposure to high amount of manganese could affect cognitive function and this was supported by results obtained by Shiva et al., Costa et al, and Jian et al., [17, 18,19]. The novel recognition and Y-maze are used to study the psychological and neural mechanisms of spatial learning and memory. The present behavioural studies have shown that Mn negatively affects learning and memory and this indicates an injury to the hippocampus which is involved in spatial memory but quercetin positively improves the spatial learning and memory [20]. The hippocampus of mice is now known to have some pyramidal neurons that help the animals navigate better through their environment as the firing of these cells indicates a
Figure 6. TNF-α expression in there were no significant changes in fur and eye colour in all the groups throughout the experiment mice exposed to manganese chloride and quercetin. Control (C); Manganese (Mn); Quercetin (Q); Intervention (POST); Concurrent (CONC). The deep/dark brown precipitates indicate the expression of TNF-α. DG: dentate gyrus, CA: Cornu ammonis
specific location in the environment of the mice. Extensive destruction of hippocampal pyramidal cells by excess Mn consumption would have affected the ability of exposed mice to correctly navigate in the NOR and y-maze tests and these could also have contributed to their poor cognitive performances. Mice with damaged cells of the dentate gyrus and cornu ammonis as a result of manganese treatment showed a decrease in spatial memory and cognitive function during neurobehavioural test with novel object recognition and y-maze test and this was significantly increase with quercetin (Figure 3 & 4). Histological findings also showed that normal histoarchitectural design of the hippocampus were preserved in the control group with a normal distribution of neuronal cells seen in the pyramidal cell layer of the hippocampus proper (cornu ammonis;CA1-CA3) and granular cell layer of the dentate gyrus of the hippocampus which could indicates normal spatial memory of the mice in these group. However, the manganese-treated group (Mn-treated mice) revealed higher destruction of cells with pyknotic nuclei in the CA1 and CA2 regions than the CA3 region of the pyramidal cell layer. The cells of the hilus of dentate gyrus were least affected by the effect of excess manganese (Figure 5). Pyknosis refers to the condensation of nuclear materials as a result of a cell necrosis or apoptosis [22, 23]. Excess manganese exposure become neurotoxin and these can extensively damage the hippocampal neurons with a characteristic pyknotic nuclei and intact densely dark nuclear fragments to influence learning, memory and possibly spatial navigation ability of mice [13, 23]. Therefore, histological results of this study clearly support the reports that excess Mn exposure extensively destroys hippocampal neurons and induces neurotoxicity in the brain [21]. Excess Mn consumption produces an extensive neuronal degeneration in the hippocampus leading to learning and memory loss [21]. Figure 5 also showed an evidence of quercetin treated mice in adult neurogenesis where there was a regeneration and repair of cells of hippocampus from damage caused by oxidative stress of the induced-manganese mice.

Tumour necrosis factor alpha (TNF alpha) represent an inflammatory cytokine produced by macrophages during inflammation involving signalling events within cells leading to programmed cell death [24, 27]. In the present study, the normal control and quercetin-treated mice according to figure 6 showed no expression of TNF-α while the manganese-induced mice presented the highest expression of TNF-α which implies that high amount of manganese exposure in the brain may lead to neuro-inflammation and neurodegeneration that could be protected by the antioxidant effect of the quercetin via inhibition of pro-inflammatory cytokine expression [25, 26, 27].

**Conclusion**

In conclusion, the efficacy of quercetin on Mn-induced mice on the hippocampus in this present investigation showed an improvement of the endogenous antioxidant defence mechanism by way of inhibition of reactive oxygen species generation and suppression of inflammation as well as reduction of expression of tumour necrosis alpha. Hence, quercetin may have beneficial health effects in Mn-exposed individuals on cognitive function.

**References**

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