Anxiolytic and curative effect of *Solanum macrocarpon* leaves extract on acetaminophen induced brain injury in adult Wistar rats

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**Abstract**

**Background of study:** Herbal plants such as *Solanum macrocarpon* may have free radical scavenging activity and is used in the treatment of brain tissue hence this study was aimed at investigating the effect of aqueous extract of *solanum macrocarpon* leaf on acetaminophen induced brain injury.

**Materials and Method:** Twenty- four adult male wistar rats with average age of 120-215 grams were purchased and used for this experiment. The rats were divided into six (6) groups of four (4) rats each. After 14-days of acclimatization, the rats were grouped into control group (group A) and experimental groups (group B, C, D, E and F). The control group were given normal saline and feed. Group B rats were given 0.5ml of acetaminophen with 200mg/kg of aqueous extract. Group C received 0.5 ml of acetaminophen with 400mg/kg of aqueous leaf extract. Group D received 0.5ml of acetaminophen with 600mg/kg of aqueous leaf extract while Group E received 0.5ml of acetaminophen only. Group F were given 0.5ml of acetaminophen with 200mg/kg of aqueous extract. Group C received 0.5ml of acetaminophen with 400mg/kg of aqueous leaf extract. Group D received 0.5ml of acetaminophen with 600mg/kg of aqueous leaf extract. Group E received 0.5ml of acetaminophen only. Group F were given standard drug, vitamin E orally for 10 days. The animal sacrificed on the 20th day, and the organ was harvested for assaying brain function and histopathological test. Brain tissues were excised and fixed in formal saline then processed using the Hematoxylin and Eosin method.

**Result:** Administrations of aqueous leaf extract of *S. macrocarpon* shows moderate healing with mild condensed neurons (CN), mild microcystic space (MS) and some plasma cell (PC) secondary to acetaminophen induced hypercholesterolemic rats (Sodipo et al., 2009c, 2011b) [18, 15, 19]. Generally, the plant has been used as an indigenous medicine for the treatment of several ailments such as asthma, allergic rhinitis, nasal catarrh, skin infections, rheumatic disease and swollen joint pains, gastro-esophageal reflux disease, constipation, dyspepsia and also in weight reduction (Dalziel, 1937). The aqueous fruit extract of the plant, has also been reported to exhibit lipid lowering activities as well as renal and hepatoprotective effects in diet-induced hypercholesterolemic rats (Sodipo et al., 2009a,b; Sodipo et al., 2009c, 2011b) [18, 15, 19], high phenolic and flavonoid content, thus possessing a potent antioxidant activity which can offer good protection against oxidative damage to some body tissues (Adewale et al., 2014) [12, 20, 41].

**Conclusion:** This present study proved that the aqueous extract of *Solanum macrocarpon* leaf exhibited a dose dependent significant antioxidant property on the brain against acetaminophen induced inflammation on the experiment rats. It is however recommended that further investigations should be carried on the efficacy and safety of use this extract on the brain and other tissues before recommending its use for certain medicinal.

**Keywords:** Anxiolytic, *Solanum macrocarpon*, acetaminophen induced brain, Wistar rats

**Introduction**

*Solanum macrocarpon* also known as the African eggplant or Gboma is a plant that belongs to the Solanaceae family and is popularly consumed as edible tropical perennial plant that is closely related to the eggplant. (Oboh et al., 2005) [6, 10]. Studies have shown that the leaves are rich in protein, fat, crude fiber, calcium, zinc (Oboh et al., 2005) [6, 10] and have gastric medicinal benefits (Bukenya-Ziraba and Bonsu, 2004) [7]. The leaves of *Solanum macrocarpon* also aid in treatment of constipation, ulcers, tooth ache and is also a snake bite antidote (Oladiran, 1989) [11]. It is used in treatment of skin diseases, infections and sores (Edijala et al., 2005) [9]. Generally, the plant has been used as an indigenous medicine for the treatment of several ailments such as asthma, allergic rhinitis, nasal catarrh, skin infections, rheumatic disease and swollen joint pains, gastro-esophageal reflux disease, constipation, dyspepsia and also in weight reduction (Dalziel, 1937). The aqueous fruit extract of the plant, has also been reported to exhibit lipid lowering activities as well as renal and hepatoprotective effects in diet-induced hypercholesterolemic rats (Sodipo et al., 2009a,b; Sodipo et al., 2009c, 2011b) [18, 15, 19], high phenolic and flavonoid content, thus possessing a potent antioxidant activity which can offer good protection against oxidative damage to some body tissues (Adewale et al., 2014) [12, 20, 41]. The protective effect of the antioxidant compounds in *Solanum macrocarpon* leaves have demonstrated high electron donating capacity which initiates chain termination in the lipid peroxidation mechanism and transforming reactive free-radical species into more stable nonreactive products, hence can offer good protection against oxidative damage to body cells.
particularly liver and brain (Olusola et al., 2014)[12,20]. Drug abuse has been shown to cause structural and functional disorders in prefrontal cortex and invariably impaired cognitive control (Charle et al., 2014). Paracetamol first made in 1877, (Mangus et al., 2005)[25] also known as acetaminophen /APAP is enlisted on the World Health Organization’s List of Essential Medicines, (World Health Organization Model list of Essential Medicines 2019)[27] and available as a generic medication with trade names including Tylenol and Panadol, among others. (Hamilton et al., 2013).

Paracetamol is generally safe at recommended doses (Russell et al., 2003)[22] and frequently used to treat pain and fever. (American Society of Health –System pharmacist et al., 2016)[21]. The recommended maximum daily dose for an adult is four grams (Lewis et al., 2013)[23]. Although it is classified as a mild analgesic, paracetamol does not have significant anti-inflammatory activity. (McKay et al., 2013)[24].

Paracetamol apparently might modulate the endogenous cannabinoid system in the brain through its metabolite, AM404, which appears to inhibit the reuptake of the endogenous cannabinoid/vanilloid anandamide by neurons, making it more available to reduce pain. AM404 also appears to be able to directly activate the TRPV1 (older name: vanilloid receptor), which also inhibits pain signals in the brain (Ghanem et al., 2016)[34].

Pharmacokinetics
After being taken orally, paracetamol is rapidly absorbed by the gastrointestinal (GI) tract. (Prescott et al., 1980)[29] and its volume of distribution is roughly 50 L. (Graham et al., 2013)[29]. The concentration in serum after a typical dose of paracetamol usually peaks below 30 µg/ml (200 µmol/L). After 4 hours, the concentration is usually less than 10 µg/ml (66 µmol/L). (Marx et al., 2013)[30]. Paracetamol is metabolized primarily in the liver, into toxic and nontoxic products. Mehta et al., (2012)[33] have noted three metabolic pathways namely Glucuronidation (45-55%), Sulfation (Sulfate conjugation) (20-30%) (Fortuny et al., 2006)[35] N-hydroxylation and dehydration, then glutathione conjugation, (less than 15%). (Ronald et al., 1995)[31].

All three pathways yield final products that are inactive, nontoxic, and eventually excreted by the kidneys. In the third pathway, however, the intermediate product NAPQI is toxic. NAPQI is primarily responsible for the toxic effects of paracetamol; this constitutes an example of toxicity (Brayfield et al. 1995)[32].

Methods
Animal Management and Grouping
30 healthy adult male wister rats (weighing about 120-210g). The animals were housed in Plastic cages with iron nettings and maintained under standard laboratory conditions (Temperature 24 °C ± 2 °C, with relative humidity of 60-70% under 12hours light and dark cycles) and were provided easy access to food (growers mesh) and water ad libitum. The animals were allowed to acclimatize for two weeks prior to the experiment. At the end of the acclimatization, twenty-four (24) were selected randomly and divided into (6) groups of four (4) animals each. They were labeled group 1-6 which group 1 served as the control group. The body weight were obtained using an electronic weighing scale before and after acclimatization, and also continued at weekly intervals during the experiment.

Animal Treatment
Group A received water and feed (positive control group), group B received 550mg/kg1 of acetaminophen and 100mg/kg1 of Solanum macrocarpon (low dose), group C received 550mg/kg1 of acetaminophen and 200mg/kg1 of Solanum macrocarpon (middle dose), group D received 550mg/kg1 of acetaminophen and 400mg/kg1 of Solanum macrocarpon(high dose), group E received 550mg/kg1 of acetaminophen only (negative control group) while group F received 550mg/kg1 of acetaminophen and 50mg/kg1 of vitamin E. Acetaminophen induced nephrotoxicity period lasted for ten (10) days followed by another ten (10) days of oral administration of Solanum macrocarpon extract.

<table>
<thead>
<tr>
<th>Group</th>
<th>No of rat</th>
<th>Total duration</th>
<th>Induced substance (pcm)</th>
<th>Dosage</th>
<th>Treatment</th>
<th>Duration of administration of drug</th>
<th>Duration of administration of extract</th>
</tr>
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<tbody>
<tr>
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<td>20 days</td>
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<td>Nil</td>
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<td>10 days</td>
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<td>100mg</td>
<td>Extract low dose</td>
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<td>Group c</td>
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<td>20 days</td>
<td>550mg</td>
<td>200mg</td>
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<td>10 days</td>
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<tr>
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<td>Nil</td>
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<tr>
<td>Group f</td>
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<td>20 days</td>
<td>550mg</td>
<td>50mg</td>
<td>Vitamin E</td>
<td>10 days</td>
<td>10 days</td>
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</table>

Collection of brain tissue (Cerebral cortex)
Under diethyl ether anesthesia, animals of each group were sacrificed on the 20th day, and the brain was harvested and fixed in 10% formalin (Dougnon et al., 2009) in a sample bottle with tight fitting lids for 72 hours as a prerule to improve staining quality and to aid optical differentiation of cell contents.

Tissue preparation for microscopy: The tissue were subsequently trimmed, dehydrated in ascending grades of alcohol (70%,80%,90% and absolute alcohol), cleared in three (3) changes of xylene and embedded in molten paraffin wax. Sections of 5um thickness with a rotary microtome were made, floated in water bath (45°C) and incubated at 60 ºc to dry. The 5um thick sections were subsequently stained in using hematoxylin and Eosin stains. The photomicrographs were taken using a motic®5.0 megapixels microscope camera.

Results
Behavioral Tests: Y- Maze test (YMT) and elevated plus maze test (EPM) were carried out in this study. The tests were performed before, during and after administration of acetaminophen in days 0, 15 and 22. Behavioral tests were performed 24 hours after a previous acetaminophen administration so as to allow time for drug distribution that may be used to access possible behavioral changes that may be caused by acetaminophen absorption in the blood stream. (Yoon et al. 2016, Ferri, Fred F. (2016),)[1, 2] All behavioral tests were recorded with a camcorder and manually timed.

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Y – Maze Test
This test is used to access spatial working memory. Arm entries were counted. An arm entry is when all four legs of the rat are completely in the arm. Spontaneous alternation was also counted. Spontaneous alternation is when an animal enters three different arms consecutively in triple sets. Percentage of alternation was calculated as \(\frac{\text{spontaneous alternation}}{\text{total number of arm entries} - 2}\) * 100 (Reza, et al., 2009) \(^3\) Each rat was placed in the Y maze apparatus for 5 minutes after which the apparatus was cleaned with 5% ethanol to clean up any possible odour left by the last animal before introducing another.

Elevated Plus Maze Test
Elevated Plus Maze (EPM) is used to access the rate of anxiety in laboratory rodents (Sun, et al., 2008) \(^4\). The EPM apparatus look like a plus sign with two open arms (50cm X 10cm) and two closed arms in opposite directions. The walls of the closed arms are 30cm high and the maze is elevated 60 cm above the floor. The open arms contain a slight ledge (4 mm high) to prevent the rats from slipping and falling off the edge. The central square is 5cm X 5cm. Each rat was placed in the centre of the elevated plus maze facing an open arm, and allowed 5 minutes to freely explore the apparatus. The following parameters were measured: open arm entries, time spent in open arms, closed arm entries, time spent in close arms and total arm entries. Arm entry is defined as when the hind paws of the rats are completely within the arm. The closed arms provide a sense of safety because they are enclosed. Behaviors such as open arm activity and head dipping are considered exploratory, and a greater frequency of these activities shows a greater level of exploration (Brown, et al., 1999) \(^5\).

Statistical analysis
All behavioral tests were carried out using one way ANOVA and Bonferroni post test. Data for each experimental group were analyzed and compared with the control group. Graphpad prism version 5.03, Graph pad.

Results
Behavioural test
Results for elevated plus maze test

<table>
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<th>GROUPS</th>
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<th>HIGH DOSE</th>
<th>SD</th>
<th>ACT</th>
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<td>DEFECATION EPM</td>
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<td>CONTROL</td>
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Fig 1: Result showing animals response to fear which shows in the amount of fecal boil deposited during test. Results did not show any significant difference in comparison of fecal boil deposits. \(P = 0.0467\) with \(P \leq 0.05\). Not statistically significant. (Kruskal Wallis Test)

Fig 2: Result showing duration of animals in the closed arm (DCA) of the elevated plus maze. Medium dose group showed the least time spent in the closed arm. \(P = 0.4159\). Test result was not statistically significant with 0.05 as level of significance. (Kruskal Wallis Test)

Fig 3: Result showing duration of animals in the open arm (DOA) of the elevated plus maze test. Only low dose and act groups showed activity in the open arm with \(P = 0.4239\). Test result was not statistically significant with 0.05 as level of significance. (Kruskal Wallis Test)

Fig 4: Result showing rate at which animals entered the close arm of the elevated plus maze across the groups. Number of Closed Arm Entries (NCAE). Animals in the low dose group made more entries into the closed arm depicting high levels of anxiety. \(P = 0.3123\). However, test result was not statistically significant with 0.05 as level of significance. (Kruskal Wallis Test)
Results for y maze test

Fig 5: Showing frequency of rearing displayed by the animals across the groups with the control and medium dose groups displaying more exploratory behavior. P = 0.4159 with P ≤ 0.05. (Kruskal Wallis Test)

Fig 6: Showing grooming activity of animals across groups with control and high dose groups displaying more response to novelty and possible stress. P = 0.4159 with P ≤ 0.05. Not statistically significant. (Kruskal Wallis Test)

Fig 7: Y maze result showing frequency of deposition of fecal boil by the animals across the groups with high level of anxiety recorded by animals in the sd group. Animals in the low dose group showed the lowest level of anxiety with low fecal boil deposits. P = 0.4123. Test result showed no significant difference in anxiety levels across the groups with 0.05 as level of significance. (Kruskal Wallis Test)

Fig 8: Y maze result showing the number of arm entries made by animals across the groups. More arm entries indicate high loco motor activity. High dose group indicated high loco motor activity followed by the low dose group. This is an indication of intense psychomotor activity with p value at 0.2124. Results showed no significant difference in the number of entries made with level of significant at 0.05. (Kruskal Wallis Test)

Fig 9: Y maze result showing level of spontaneous alternation showed by animals across the groups. It shows the willingness of the animals to explore new environments. The low dose group of animals showed marked willingness to explore with p value at 0.2134. Results showed no significant difference in willingness to explore new environments with 0.05 as level of significance. (Kruskal Wallis Test)

Fig 10: Showing rearing activity amongst animals across the groups. Animals in high dose groups displayed more exploratory activity. P=0.4159 with P ≤ 0.05. Not statistically significant. (Kruskal Wallis Test)
Fig 11: Showing animals response to novelty and possible stress with the high dose group displaying more grooming activity. P=0.4257 with P ≤ 0.05, statistically significant (Kruskal Wallis Test)

Histological findings
Group A (CONTROL GROUP)

Plate 1: Photomicrograph of the cerebral cortex showing normal neural cell bodies in the gray matter (NC) (H & E. X400)

Group B

Plate 2: Photomicrograph of cerebral cortex induced with 550mg/kg of acetaminophen and 100mg/kg Solanum macrocarpon extract showing mild moderately condensed neurons (CN) and moderate microcytic spaces (MS). (H & E. X400)

Group C

Plate 3: Photomicrograph of cerebral cortex induced with 550mg/kg of acetaminophen and 200mg/kg of Solanum macrocarpon showing moderate healing with mildly condensed neurons (CN) and mild microcytic space (MS) (H & E. X400)

Group D (High dose)

Plate 4: Sections of cerebral cortex induced with 550mg/kg of acetaminophen and 400mg/kg solanum macrocarpon shows moderate healing with mildly condensed neurons (CN), mild microcytic space (MS). H & E. X400

Group E

Plate 5: Photomicrograph of cerebral cortex induced with 550mg/kg of acetaminophen and vitamin E showing moderate healing with mild microcytic spaces (MS). H & E. X400
Group F

Plate 6: Photomicrograph of the cerebral cortex induced with 550mg/kg of acetaminophen showing moderate degeneration with microcystic space (MS), shrinking of the neurons (SN) moderate coagulative necrosis (ES) and eosinophilic substance within the vacuolated space (ES). H & E. X400.

Discussion
This work was aimed at investigating the curative effect of *Solanum macrocarpon* leaves extract on acetaminophen induced brain injury in adult wistar rats. All animals except control group were given 550mg/kg of acetaminophen. Result On the present study indicates that there was no significant change on anxiety levels amongst wistar rats used for this study. There was minimal possibility of causing motor deficits after administering 550mg/kg of Acetaminophen. Behavioral test findings revealed slight increase in anxiety levels with low dose group with slight exploratory tendencies especially in animals in the high dose extract group. Acute exposure to solanum Macrocarpon leaf extracts indicated potential anxiolytic properties (Melina Giorgetti & Giuseppina Negri, 2011) [37]. The animals in the low dose group displayed less exploratory tendencies. This is possibly via decrease activity of midbrain dopaminergic neurons of nigrostriatal system (Seppa T et al. 2000) [38]. Acetaminophen group displayed high anxiety traits. Results show that acetaminophen alone increases brain serotonin as well as norepinephrine levels with a concomitant inhibition of liver TDO activity. (Maharaj et al, 2004) [38]. Grooming and rearing activity showed no significant difference in anxiety levels as animals’ response to novelty was higher in the high dose group as this in a possible indicator of stress. There were observable histological observations attributed to the effects of solanum macrocarpon extracts. These effects observably increased with increases in the doses of the extract; this implies that the effects were dose dependent. Results reveal moderately condensed neurons in animals administered with 100mg/kg, 200mg/kg and 400mg/kg of solanum macrocarpon leaf extract. Plate 6 given 550mg/kg of ACT which did not receive any therapeutic intervention, showed neuronal degeneration and coagulative necrosis. There were also evidences of alterations of the cerebral histoarchitecture and the neuropil. Reports showed that the extract at all dosages had deleterious effects on the brain tissue and the gravity of effects increased with the dosage (Owolabi et al, 2016) [40]. Plate 4 revealed that sections of cerebral cortex induced with 550mg/kg of acetaminophen and 400mg/ kg *Solanum macrocarpon* shows moderate healing with mild condensed neurons. This is an indication of the curative ability of solanum macrocarpon extracts. Olusola et al, proposed that solanum macrocarpon extracts have protective properties in the cells of the brain using wistar rats (Olusola et al, 2014) [12, 20].

Acetaminophen overdose resulted in locomotor retardation, excessive self-grooming, working-memory impairment, anxiety, derangement of liver/kidney biochemistry, antioxidant imbalance, and histological changes in the liver, kidney and cerebral cortex (Olatunji et al, 2017) [39].

Conclusion
This present study proved that aqueous extract of *Solanum macrocarpon* leaf exhibited a dose dependent mild anxiolytic activity on the brain against acetaminophen induced inflammation on adult wistar rats and showed mild curative properties in the cerebral cortex. It is however recommended that further clinical and phytochemical investigations should be carried on the efficacy and safety of use this extract on various brain centers and also in astrocyte and microglial population in the limbic system as these cells perform inflammatory reactions towards drug interactions.

List of abbreviations
ACT: Acetaminophen
EPM: Elevated Plus Maze
DCA: Duration in close Arm
DOA: Duration in Open Arm
NCAE: Number of Close Arm entries

References

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38. Maharaj H, Maharaj DS, Saravanan KS, Mohanakumar KP, Daya S. Aspirin curtails the acetaminophen-induced

