Ameliorative Potential of *Allium cepa* Lam. Leaves on Diabetes Induced and Chronic Constriction Injury Induced Neuropathic Pain in Experimental Rats

Dureshahwar Khan¹, Mubashir Mohammed¹, Aman Upaganlawar², Chandrashekhar D. Upasani², Hemant D. Une¹,*

¹Department of Pharmacology, Y. B. Chavan College of Pharmacy, Dr. Rafiq Zakaria Campus, Dr. Rafiq Zakaria Marg, Rauza Bagh, Aurangabad, Maharashtra, INDIA.
²Department of Pharmacology, SNJB’s SSDJ College of Pharmacy, Neminagar, Chandwad, Maharashtra, INDIA.

**ABSTRACT**

**Background:** Neuropathy can be induced in rats by peripheral injuries, depending on compression of complete or a section of sciatic nerve and chemically by Streptozotocin (STZ). **Materials and Methods:** In the present study neuropathic pain were induced in rats by two methods, chronic constriction injury (surgical model) and STZ (40mg/kg/i.p.) induced diabetes (drug induced model). In both the models, behavioural as well as markers of oxidative stress were studied. Behavioural parameters were tested using vonfrey hair and Randall Selitto analgesiometer whereas biochemical parameter includes glycosylated haemoglobin and markers of oxidative stress. The study was further supported by histopathology of sciatic nerve. Flavonoid rich extract of *Allium cepa* Lam. leaves was administered at three different doses viz. 25, 50 and 100mg/kg/p.o to the rats with neuropathic pain. Both the models of neuropathic pain showed significant alteration in behavioural as well as oxidative stress parameters. **Results:** Treatment of *Allium cepa* leaves extract showed dose dependent improvement in behavioural and biochemical parameters towards normal (*p* value <0.001, <0.05 and <0.01). The altered histopathological changes in sciatic nerve were also significantly improved as compared to rats with neuropathic pain. **Conclusion:** The neuroprotective effects of the *Allium cepa* leaves extract is a virtue of its strong antioxidant activity.

**Key words:** *Allium cepa*, Neuropathy, CCI, Oxidative stress, Sciatic nerve.

**INTRODUCTION**

Different animal models of neuropathy showed a strong correlation of sciatic nerve with neuropathic pain. Peripheral nerve injury models involve surgical procedures at sciatic nerve to induce neuropathy, whereas drug induced neuropathy and disease induced neuropathy are models developed due to oxidative stress, degeneration or neurotoxicity of peripheral nerves i.e sciatic nerve.¹ Surgical models targeting nerve result in a chronic mechanical allodynia on injured paw. These models include chronic constriction injury, sciatic nerve cuffing, partial sciatic nerve ligation, spinal nerve ligation or common peroneal nerve ligation.² While the disease induced neuropathy commonly include diabetic neuropathy model in research groups,¹ chronic hyperglycemia induces oxidative stress through multiple pathways like redox imbalances, altered protein kinase C activity and mitochondrial overproduction of superoxide³ and this plays a significant role in development of diabetic complications including diabetic neuropathy.⁴ While, sciatic nerve injury is associated with excessive production of reactive oxygen species, calcium ions entry and apoptosis; calcium overload through Transient Receptor Potential (TRP) cation channel and pain...
intensity are associated with resulting oxidative stress following CCI.\(^5\)

Being a reliable model of inducing diabetic neuropathy, Streptozotocin (STZ) in multiple sub-diabetogenic doses results in chronic hyperglycemia from partial damage to islets followed by an inflammatory process. In previous research work, it was found to develop diabetic neuropathy in rats.\(^6\) Chronic hyperglycemia and diabetic neuropathy leads to dysfunction and degeneration of the sciatic nerve.\(^7\)

*Allium cepa* Lam. is a common and medicinally valuable plant. Leaves are reported to be aphrodisiac, anti-spasmodic, antihelminthic, alterative, carminative, digestive, diuretic, emollient, expectorant, mild laxative, stimulant and tonic along with presence of flavonoids,\(^8\) vitamin A, thiamin and ascorbic acid.\(^9\) Outer layers of scales of its bulbs in different varieties were studied by researchers for their effects on chronic constriction injury induced neuropathic pain and resulted into significant antioxidant effect and improvement in neuropathic pain.\(^10\)

In the present study flavonoid rich extract of *A. cepa* leaves was used to evaluate its efficiency for protecting sciatic nerve against neuropathy and oxidative stress by examining the extract effects on biomarkers of neuropathy in both surgical and diabetic neuropathy models followed by evaluation of oxidative stress after 5 weeks of diabetic neuropathy and 3 weeks of treatment and later animals were sacrificed to study histology of sciatic nerve.

**MATERIALS AND METHODS**

**Drugs and chemicals**

Streptozotocin (STZ, Spectrochem Pvt. Ltd.) was mixed in cold 0.01 M citrate buffer, pH 4.5, prepared freshly before administration,\(^6,11\) and all other analytical grade chemicals were consumed in investigation.

**Plant material**

The leaves of *A. cepa* were collected from the local area of Aurangabad, Maharashtra, India. They were authenticated by Department of Botany, Dr. Babasaheb Ambedkar Marathwada University, Aurangabad, Maharashtra and a voucher specimen 0587 has been deposited in the herbarium of Dr. Babasaheb Ambedkar Marathwada University, Aurangabad (MS), India. Flavonoid rich extract was obtained from methanolic extract of *A. cepa* leaves followed by fractionation using ethyl acetate (ACEA- *A. cepa*: Ethyl acetate fraction) from the technique reported in earlier research work.\(^6\)

**Experimental animals**

Sprague Dawley rats of either sex weighing between 180-250g were used. Prior to the experimental work, acclimatization of rats to the conditions of animal house for one week was carried out. Normal pellet diet and water *ad libitum* was followed for them. They were housed at standard conditions of temperature (23±12°C), humidity (45±5%) and 12 hr. light and dark cycle. The experimental protocol for animal experiments was approved by the Institutional Animal Ethics Committee (IAEC) of Y. B. Chavan College of Pharmacy, Aurangabad (Approval no. CPCSEA/IAEC/P’col-52/115).

**Induction of Neuropathy**

**Induction of experimental diabetic neuropathy**

Following overnight fasting experimental rats were injected with a multiple dose of 40 mg/kg/i.p. of STZ for 3 consecutive days. Soon within 5 min after mixing in citrate buffer pH 4.5 the STZ solution was administered. The rats in group NC were injected with distilled water as a vehicle control. The animals were allowed to drink 5% glucose solution *ad libitum* overnight to overcome hypoglycemia. On inducing diabetes random blood glucose was estimated followed by regular analysis until attaining stable hyperglycemia. Rats may develop hyperglycemia and additional clinical diabetic indications within 3 days of STZ injection. After 1 week rats showing blood glucose above 250mg/dl were preferred to study, 24 rats were selected for further studies. After 3 weeks of stable diabetes drug treatment was performed and pretreatment (pre T/T) studies of parameters of neuropathy were performed. On completion of dosing period rats were again tested for different parameters of Neuropathy.\(^6,11,12\)

**Experimental Design**

Animals were distributed in various groups composed of six animals each. Group NC: Normal control, vehicle treated, Group DC: Diabetic control, vehicle treated, Group ACEA\(_{25}\): Diabetic, ACEA treated (25 mg/kg, p. o.), Group ACEA\(_{50}\): Diabetic, ACEA treated (50 mg/kg, p. o.), Group ACEA\(_{100}\): Diabetic, ACEA treated (100 mg/kg, p. o.)

**Surgical Model of Neuropathy**

In order to obtain the surgical control group, partial sciatic nerve ligation technique was followed. Rats were allowed to fast overnight. Skin incision was made dorsal to pelvis followed by anaesthesia. The common sciatic nerve was first exposed by separating the muscles then loosely ligated with a chromic gut suture. Fascia,
muscle and skin incision were closed using silk suture. After the completion of surgery and recovery of animals, the group was used for comparative study to assess tactile alldynia.\textsuperscript{2,13,14}

\textbf{Experimental design}

Animals were allotted various groups with six animals in each group. Group NC: Normal control, vehicle treated, Group DC: Diabetic control, vehicle treated, Group ACEA\textsubscript{25}: Nerve ligation, ACEA treated (25 mg/kg, p. o.), Group ACEA\textsubscript{50}: Nerve ligation, ACEA treated (50 mg/kg, p. o.), Group ACEA\textsubscript{100}: Nerve ligation, ACEA treated (100 mg/kg, p. o.)

\textbf{Blood Collection and blood glucose estimation}

Blood samples from the experimental groups were collected from tail vein. The samples so collected were analyzed for glucose estimation using FAD-GDH method by Contour TS glucometer.\textsuperscript{6,11}

\textbf{Assessment of Behavioural Parameters}

\textbf{Tactile Allodynia}

\textbf{Von Frey Filament model}

Stimulus presentation and testing paradigms as described by S.R. Chaplan \textit{et al.} was followed to obtain 50\% threshold calculations and estimate tactile allodynia in rats. Animals were placed in acrylic chamber with wire mesh at bottom which gives full access to the paws. Animals were conceded to get acclimatized in the cage for about 15 min. The paw was touched with vonfrey filaments following up-down method in order to calculate 50\% withdrawal threshold. Vonfrey hairs were subjected to the paw perpendicularly with sufficient force to cause buckling response against paw and held for about 6-8 sec. Sharp withdrawal of paw was considered to be positive response.\textsuperscript{11,15,16}

50\% g threshold was calculated by formula:

\[(10^{\text{f(\text{kg})}})/10,000\]

Where \(f\) – Value (in log units) of final vonfrey hair used.

k - Tabular value for the pattern of positive / negative responses (as provided in table)\textsuperscript{15}

δ- Mean difference (in log units) between stimuli (here, 0.224)

\textbf{Randall Selitto Analgesiometer}

Nocteceptive withdrawal threshold was estimated on Randall selitto test apparatus on the paws of experimental animals. Hold animals with soft cotton cloth in order to immobilize it. Place gently the paw on the apparatus and allow the tip of device to apply on paw with application of increasing mechanical force and withdrawal latency to the pressure supported was noted down.\textsuperscript{3,11}

\textbf{Assessment of Biochemical Parameters}

\textbf{Glycosylated Haemoglobin}

Glycosylated Haemoglobin (HbA\textsubscript{1c}) estimation was carried out by calorimetric method. Before sacrificing animals for biochemical and histopathological studies blood was collected using fine glass capillary puncturing of retro-orbital plexus in epindorff tubes. Serum and packed blood cells were separated by centrifugation for collecting RBCs. The packed cells were washed 3-4 times in normal saline. After final wash distilled water and CCl\textsubscript{4} 0.5 ml each were added and mixed vigorously followed by 20 min centrifugation at 3000 rpm. The supernatant haemolysate were separated and its Hb concentration was adjusted to 10 gms % with distill water. To 2 ml of haemolysate, 1ml of 0.3 N oxalic acid was added and kept in boiling water bath for 60 min after covering the test tube using cotton plug. After cooling 1 ml of 40\% TCA was added, shaken vigorously and centrifuged at 3000 rpm. To 2 ml of this supernatant 0.5 ml of 0.05 M thiobarbituric acid was added and kept at 37\°C for 40 min. The resultant yellow color was read on colorimeter at 443 nm, reading was taken as blank 2ml distilled water and 0.5ml of thiobarbituric acid (HbA\textsubscript{1c} was calculated on assumption that 1\% HbA\textsubscript{1c} corresponds to an absorbance of 0.029 at 443 nm).\textsuperscript{3,18}

\textbf{Estimation of markers of oxidative stress}

At end of the treatment and behavioral assessments, animals were sacrificed under deep anesthesia preceded by overnight fasting. By using Remi homogenizer, sciatic nerves on rapid removal were homogenized in a cold 50 mM phosphate buffered saline (pH 7.4). The 10\% w/v tissue homogenate was centrifuged at 1000 rpm for 20 min at 4\°C for estimation of catalase activity while for SOD and GSH homogenate was centrifuged at 12000 rpm as long as 60 min at 4\°C. Catalase activity was estimated applying H\textsubscript{2}O\textsubscript{2} just as substrate. In brief, H\textsubscript{2}O\textsubscript{2} decomposition by catalase was monitored following the decrease in absorbance at 240 nm.\textsuperscript{19} SOD activity in sciatic nerve was estimated by its ability to inhibit auto-oxidation of Pyrogallol at 420 nm.\textsuperscript{20} Concentration of GSH was estimated spectrophotometrically using DTNB reagent at 412 nm.\textsuperscript{21,22}

\textbf{Histopathological evaluation}

Animals were sacrificed and the sciatic nerve was carefully removed from each animal. The nerve tissue was washed with saline and fixed in 10\% formalin for histopathological study. Slides were prepared by embedding
the nerve in paraffin and staining with hematoxylin and
eosin. The histology of sciatic nerves was studied for
axonal degeneration, dilatation, edema and vacuoliza-
tion using a bright field microscope and images were
taken for reference.

**Statistical Analysis**
All the data expressed by Mean ± SEM (n=6), two-
way ANOVA, followed by Benferroni test using Prism
Graph Pad version 5 with \( p < 0.001, <0.05 \) and
<0.01 statistical significance.

**RESULTS**
In the previous research work it was reported that
ACEA fraction is flavonoids rich having presence
of quercetin with a potential ant diabetic activity and
protective in thermal hyperalgesia, cold allodynia and
motor inco-ordination behavioral biomarkers of diabetic
neuropathy.

**Effects of ACEA on mechanical alldynia (Vonfrey
filaments) in neuropathic pain.**
Nociceptive withdrawal threshold expressed in terms
of 50% threshold of diabetic control and surgical
control was significantly \( p<0.001 \) decreased compar-
able with that of normal control. The ACEA treated groups
shows dose dependant significant \( p<0.05, 0.001 \)
increase in paw withdrawal threshold comparable with
that of diabetic and surgical control (Table 1).

**Effects of ACEA on Mechanical or Static Hyperalgesia
(Randall Selitto test) in neuropathic pain.**
Nociceptive withdrawal threshold expressed in terms
of supported pressure resulting from application of
randall-selitto probe of diabetic and surgical control has
found to decrease significantly \( p<0.001 \) comparable with
that of normal control while ACEA has shown
dose dependant increase in supported pressure compa-
rable with that of diabetic and surgical control signifi-
cantly \( p<0.001 \), (Table 1).

**Effects of ACEA on Glycosylated Haemoglobin level in
neuropathic pain.**
The HbA1c (%) value of diabetic control has found to
increase significantly \( p<0.001 \) comparable with that of
normal control while ACEA has shown dose dependant
decrease in value comparable with that of diabetic con-
trol significantly \( p<0.05, p<0.001 \), (Figure 1).

**Effects of ACEA on oxidative stress markers in
neuropathic pain.**
The biomarkers of oxidative stress were assessed,
ACEA dose dependently increased the Catalase, SOD
and GSH levels significantly \( p<0.01, p<0.001 \) in sci-
atic nerve comparable with that of diabetic and surgical
control group (Figure 2) thus, resulted in normalization
of STZ as well as CCI induced biochemical abnormalities
in a significant manner.

| Table 1: Effect of ACEA on mechanical allodynia in neuropathic pain. |
|---------------|-----------------|--------------|-------------|--------------|
| Tactile Allodynia | Surgical induced neuropathy |
|                | NC       | SC        | ACEA25     | ACEA50      | ACEA100     |
| Vonfrey Filaments (50% threshold (gm)) | 8.81±0.7 | 1.72±0.1** | 3.41±0.2** | 4.35±0.1**  | 5.7±0.2**   |
| Randall-Sellito apparatus (Supported Pressure (gm)) | 318.06±18.9 | 82.54±3.2** | 123.13±4 | 140.60±1.5  | 234.80±28.2** |
| STZ induced neuropathy |
|                | NC       | DC        | ACEA25     | ACEA50      | ACEA100     |
| Vonfrey Filaments (50% threshold (gm)) | 8.81±0.7 | 1.37±0.09** | 3.36±0.2*  | 4.20±0.1**  | 5.61±0.2**  |
| Randall-Sellito apparatus (Supported Pressure (gm)) | 318.06±18.9 | 81.16±3.4** | 122.84±4.1 | 140.23±1.3  | 234.49±28.6** |

Data is represented in the form of Mean ±SEM (n=6), Two way ANOVA followed by Benferroni test using Prism Graph Pad version 5 with *p<0.05 and **p<0.01 significance for groups are compared with normal group while #p<0.05 and ###p<0.001 for groups compared with diseased group. (NC-Normal Control, DC-Diabetes Control, ACEA25-Extract treated, 25mg/kg p.o., ACEA50-Extract treated, 50mg/kg p.o. and ACEA100-Extract treated, 100mg/kg p.o.)
DISCUSSION

Different animal models for neuropathic pain circles around sciatic nerve dysfunction, from which the surgical model and diabetes induced neuropathy were successfully followed in the present work. The surgical nerve injury model mimics symptoms of chronic nerve compression in humans. Previous research work has proved STZ induced diabetic neuropathy model to be reliable and widely accepted model and has also proven ACEA to be a flavonoid rich extract also it was found that free radical generation and oxidative stress has long been involved in neuropathic pain pathogenesis. On the basis of previous reports of signs of STZ induced diabetic model after one week from induction of diabetes, animals were tested for stable hyperglycemia and hyperalgesia and were used for further studies. In tactile allodynia quantitative assessment is estimated by Vonfrey filaments and Randall selitto test, 50% threshold for nociceptive withdrawal latency provides quantitative assessment of tactile allodynia expressed by Vonfrey filaments. Randall selitto is another adequate and sensitive method to quantify tactile allodynia in neuropathy, threshold for withdrawal latency is measured in the form of supported pressure in randall selitto test. Thus, in the present study tactile allodynia has been successfully quantified and proved to be useful and versatile techniques for assessment of appearance of neuropathic pain.

Glycosylated Haemoglobin (HbA\textsubscript{1c}) is a result of haemoglobin molecule undergoing post translational changes, their levels correlate with glycemic levels over the previous six to ten weeks. Under physiological condition, a reaction between glucose and N terminal valine of Beta chain of Hb molecules proceeds to glycosylation of haemoglobin. In the existing study this parameter was used to support STZ induced neuropathic pain model follow up of animals.

All of the histopathological parameters of sciatic nerves are shown in Figure 3. In diabetic control and surgical control groups, it was determined severe axonal degenerations; mild dilatations and edema were observed in nerve epineuriums. While ACEA treated group depicted repaired sciatic nerve with no axonal degenerations, dilatation, edema and vacuolization.

Effects of ACEA on Histopathological changes in sciatic nerve.

Figure 3: Histopathology study of Sciatic nerve with H and E stain, where ad: axonal degeneration, d: dilatation and e: edema. (ACEA\textsubscript{100} - Extract treated, 100mg/kg p.o.)
neuropathy and it was found ACEA has attenuating action over it too (Figure 1).

It has been reported that sciatic nerve ligation induces imbalance in reactive oxygen species and antioxidant enzymes also oxidative stress has been noted to induced by hyperglycemia.\(^{25}\) In our study, ACEA has shown restoration of biochemical parameters such as catalase, SOD and GSH level in nerve tissues.

It is well established by previous reports that chronic hyperglycema induces oxidative stress and proves to be neurotoxic that leads to neuronal apoptosis.\(^{26}\) These neuronal damages can be studied with the help of morphological study of sciatic nerve. Histopathology study of sciatic nerve highlighting axonal degeneration, dilatation, edema and vacuolization acts as degree of nerve damage\(^ {24}\) due to surgical and diabetes induced neuropathy were obtained in results. Present work determines histopathology of sciatic nerve of diabetic control and surgical control groups showed significant derangement of nerve cells with signs of axonal degeneration, dilatation and edema while that of ACEA treated group shows signs of repair of nerve cells from structural alteration as there are no signs of degeneration seen as compared to surgical and diabetic control groups (Figure 3). Improved nerve structures are considered as a marker for protective action of extract on damaged nerve.\(^ {23}\)

**CONCLUSION**

Surgical as well as STZ induced neuropathy model were found successful in inducing neuropathy and oxidative stress in present work. ACEA has depicted protective action in biomarkers of tactile allodynia, oxidative stress; glycosylated haemoglobin and conformation of protective effects were obtained through histopathological study of sciatic nerve. ACEA has significantly proven to have ameliorative potential over STZ and CCI induced neuropathic pain addressing it as a better approach to drug development in treatment of neuropathy.

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**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

**ABBREVIATIONS**

**ACEA:** Ethyl acetate fraction from methanolic extract of leaves of *A. cepa* Lam;  **FAD-GDH:** Flavin Adenine Dinucleotide- Glucose Dehydrogenase.

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SUMMARY

The current research work shows surgical as well as STZ induced neuropathy model to be successful in inducing neuropathy and oxidative stress. Leaves of *Allium cepa* L. provides protective action in quantitative assessment of tactile allodynia, oxidative stress; glycosylated haemoglobin and histopathological study of the sciatic nerve. This plant sample has proven ameliorative potential in STZ and CCI induced neuropathic pain.

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