IN VITRO ANTI-INFLAMMATORY ACTIVITY OF PROTEINS ISOLATED FROM MUNTINGIA CALABURA PLANT ROOT

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ABSTRACT

Objective: To investigate the in vitro, Antioxidant and Anti-inflammatory activity of the proteins isolated from Muntingia Calabura plant root. Methods: Proteins of the extract was isolated by Ammonium sulphate protein precipitate method. In vitro antioxidant studies were carried out for the Muntingia Calabura root proteins using superoxide radical scavenging method and anti-inflammatory studies was done using membrane stabilization assay were performed for Muntingia Calabura root proteins. Results: The superoxide radical scavenging ability of the proteins was compared with standard antioxidant like BHA and Ascorbic acid at a maximum dosage of 8µg, where Muntingia Calabura root protein showed a maximum inhibition of 71% and BHA and Ascorbic showed 65 and 60% respectively. The anti-inflammatory activity of Muntingia Calabura root proteins were compared with the standard drug Diclofenac sodium, where at a
dosage of 1000µg/ml, plant root protein showed maximum inhibition where as the drug Diclofenac Sodium showed at a dosage of 200µg/ml. **Conclusion:** The results of the present study concluded that the *Muntingia Calabura* root proteins possess significant antioxidant and anti-inflammatory activity and it may due to the presence of proteins.

**KEYWORDS:** *Muntingia Calabura*, proteins Antioxidant, Antiinflammant.

**INTRODUCTION**

Inflammation is a sign of the biological response of body tissues to harmful environmental stimuli, such as pathogens, damaged cells, or irritants resulting in pain, redness, swelling, loss of function etc.[1][2] Anti-inflammant means, it is the property of a substance or that reduces inflammation or swelling. Non-steroidal anti-inflammatory drugs (NSAID) available in Pharmaceutical market alleviate pain by counteracting the cyclooxygenase (COX) enzyme, which synthesizes prostaglandins and creating inflammation.[3] The use of NSAID drug medication is still used by most of the clinicians for joint and spine related inflammatory pain, despite their commonly known side effects, that, in the long usage of these anti-inflammants, can cause gastric erosions, which can become stomach ulcers and in extreme cases can cause severe haemorrhage, resulting in death.[4] Hence, there is a need of searching of nontoxic, inexpensive, easily available herbal sources or medicinal plant sources or dietary sources, which one can use easily as a part of dietary component as ant antioxidant and as well as anti-inflammant.

In Ayurveda, Chinese medicine it is quoted that, my herbs like Turmeric, Ginger, and Garlic had antioxidant and antiinflammant properties, but these types of preparations are not scientifically evaluated. *Muntingia calabura* plant, the sole species of genus *Muntingia*, is a flowering plant native to southern Mexico, the Caribbean, Central America, and western South America south to Peru and Bolivia. The leaves and flowers of the plant are used in the preparation of Tea and jam. It is reported that, the leaves and fruits are having antinociceptive, anti-inflammatory, antipyretic, antimicrobial and antipruritic properties.[5][6][7][8] These results encouraged us to study the anti-inflammatory activity of the plant root.

**MATERIALS AND METHODS**

**Extraction of proteins from *Muntingia Calabura* plant root**

10g of cleaned *Muntingia Calabura* root collected from authentic source, cleaned with 0.1% KMnO₄ solution, followed with double distilled water, crushed, shade dried and powdered (British Pharmacopoeia 100 mesh) and stored in glass bottle. The root powder mixed with
200 ml of boiling double distilled water, boiled for five minutes and vortexed for 4 hours at 20°C using magnetic stirrer. The vortexed mixture is centrifuged at 6000 rpm for 20 minutes, the supernatant was separated. The supernatant was subjected to 65% ammonium sulphate precipitation and vortexed over night. The mixture was centrifuged at 10000 rpm. The precipitated protein was collected and subjected to dialysis using 2.5kDa molecular cutoff biomembrane against water for 76 hours with an interval of 6 hours. The dialyzed precipitated was separated and stored at -10°C for further analysis.

**Stability to proteases**

The *Muntingia Calabura* root proteins were tested for its ability to withstand with proteases like trypsin, and pepsin. 500μg of *Muntingia Calabura* root proteins was incubated at 37°C for 1 hour with 20μg of trypsin in 20mM phosphate buffer pH 8.0 or 20μg of pepsin in 20mM sodium acetate buffer, pH 2.0. The reaction was arrested by keeping the tubes in ice. The incubation mixture contained *Muntingia Calabura* root proteins in the presence or absence of proteolytic enzymes in a ratio of 25:1 w/w was used. Aliquots of samples were then subjected to check their antioxidant capacity.

**Antioxidant activity**

**Superoxide scavenging activity**

The Superoxide radical (O2•-) scavenging activity of was measured according to the method of Lee et al.\(^9\) with minor modifications. The reaction mixture containing 100μl of 30mM EDTA (pH 7.4), 10μl of 30mM hypoxanthine in 50mM NaOH, and 200μl of 1.42mM nitro blue tetrazolium with or without *Muntingia Calabura* root proteins and SOD serving as positive control at various concentrations ranging from 50-300μg. After the solution was pre-incubated at ambient temperature for 3min, 100μl of xanthine oxidase solution (0.5U/ml) was added to the mixture and incubated for one hour at 37°C, and the volume was made up to 3ml with 20mM phosphate buffer (pH 7.4). The solution was incubated at room temperature for 20 min, absorbance was measured at 560 nm. Appropriate controls were included to rule out the artifacts induced reaction. The control was without any inhibitor. Inhibitory effect of *Muntingia Calabura* root proteins on superoxide radicals was calculated as

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\text{% Superoxide radical scavenging activity} = \frac{\text{Absorbance of control} - \text{Absorbance of test sample}}{\text{Absorbance of control}} \times 100
\]
**In vitro anti-inflammatory activity**

**Membrane stabilization assay**

The Human Red Blood Cells (HRBC) membrane stabilization method has been used to study the anti-inflammatory activity.\[10\][11] Blood was collected from the healthy volunteers and mixed with equal volume of sterilized Alsever solution (2% dextrose, 0.8% sodium citrate, 0.05% citric acid and 0.42% sodium chloride in water). The blood was centrifuged at 3000 rpm and packed cells were washed with isosaline (0.85%, pH 7.2) and a suspension was made with isosaline (10%v/v). The assay mixture contained 1 ml of Phosphate buffer (0.15M, pH 7.4), 2 ml of hyposaline (0.36%), 0.5ml of HRBC suspension and 1 ml of various concentration of the extract. Diclofenac sodium was used as standard drug. In the control solution, instead of hyposaline, 2ml of distilled water was added. The mixtures were incubated at 37°C for 30 min and centrifuged. The absorbance of the supernatant solution was read at 560nm spectrophotometrically. The % haemolysis was calculated by assuming the haemolysis produced in the presence of distilled water as 100%. The percentage of HRBC membrane stabilization was calculated using the formula.

\[
\text{% membrane stabilization} = \frac{100 - \text{O.D. of drug treated sample}}{\text{OD of Control}} \times 100
\]

**Statistical analysis**

The data were expressed as means ± standard deviations (SD). All the experiments were repeated at least three times and the values are expressed as Mean ± SD. The significance of the experimental observation was checked by student’s test and the value of p value.

**RESULTS AND DISCUSSION**

The boiling water extraction of *Muntingia Calabura* root was done as explained in materials and methods. Further the supernatant was subjected to Ammonium Sulphate precipitation to precipitate proteins of the root, followed by dialysis against water to removed unwanted salts and confirmed the precipitate rich with proteins by proximate analysis as explained by Mohamed Azmathulla Khan et al., 2015.\[12\] To confirm the antiprotease activity or protease inhibitory activity of the root proteins, the *Muntingia Calabura* root proteins of 500μg was treated with 20μg of pepsin/trypsin. Hydroxyl radical scavenging activity was done by deoxyribose assay as described in methods. The appropriate controls were included in all the experiments. The control was without any The *Muntingia Calabura* root proteins or enzyme
and the % hydroxyl radical scavenging activity was calculated accordingly (Results not shown). The antioxidant activity of the proteins of Muntingia Calabura root, was analyzed by the superoxide radicals scavenging activity studies. Here the superoxide radicals are the free radicals and are generated in a variety of biological systems by auto-oxidation processes or by enzymatic activities.\[^{12}\] Moreover, superoxide anions produce other kinds of cell damaging free radicals and oxidizing agents.\[^{13}\] Herein, we used the NBT assay system to check Muntingia Calabura root proteins scavenge superoxide radicals. As shown in Figure-1, a dose dependent study was done towards scavenging superoxide radicals by Muntingia Calabura root proteins along with other antioxidants like BHA and Ascorbic acid. Here Muntingia Calabura root proteins inhibited NBT reduction by a maximum dose of 8µg. BHA shows a maximum inhibition at a dosage of 7 µg and Ascorbic shows maximum inhibition of reduction of NBT at a dosage of 8 µg. This observation indicates that Muntingia Calabura root proteins are superoxide scavengers. It also confirms that, the root proteins are also heat stable.

Further, the anti-inflammatory activity of the root proteins was studied by Membrane stabilization assay. The prevention of hyptonicity induced HRBC membrane lysis is taken as a measure of anti-inflammatory activity as HRBC membrane are similar to lysosomal membrane components.\[^{14}\] As shown in figure 2, the Muntingia Calabura root crude proteins shows maximum anti-inflammatory activity at the concentration of 1000µg/ml which is comparable to that of standard drug Diclofenac sodium (200µg/ml). The anti-inflammatory activity of the crude protein was concentration dependent. The proteins exhibited membrane stabilization effect by inhibiting hyptonicity induced lysis of erythrocyte membrane. The erythrocyte membrane is comparable to the lysosomal membrane and hence, its stabilization shows that, the Muntingia Calabura root proteins may stabilize lysosomal membrane. The above results shows that, the proteins of Muntingia Calabura root boiling water extract are good antioxidants when compared to standard antioxidants and also having anti-inflammatory studies. These results need to be confirmed by in vivo anti-inflammatory studies.
Fig. -1 Superoxide radical scavenging activity of Muntingia Calabura root proteins:
Dose dependent.

Dose-dependent Super oxide anions scavenging activity of *Muntingia Calabura* root protein. The control was without protein or BHA or Ascorbic acid. The Super oxide radical scavenging activity was calculated accordingly as described in methods.

Results are shown as mean ± SD (n = 3).

Fig. -2: *In vitro* anti-inflammatory effect of *Muntingia Calabura* root crude proteins:
Dose dependent.

Dose-dependent anti-inflammatory studies of *Muntingia Calabura* root protein (10 to 1000µg/ml) and standard drug Diclofenac sodium (100 and 200µg/ml).

Results are shown as mean ± SD (n = 3).
REFERENCE
