Protease Activity of Floral Extracts of *Jasminum grandiflorum* L., a Wound Healing Herb

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Flowers of *Jasminum grandiflorum*, are used in traditional medicine for rapid wound healing. Protease, a potential candidate in wound healing is not so for studied in *J. grandiflorum* flowers. So an attempt was made to determine the protease activity of floral extracts of *J. grandiflorum*. Buffers of different pH range were used for extraction of the flowers to identify the best buffer for extraction of protease. Total protein content and protease activity were determined in the floral extracts as well as in different organs of the flower. Floral extract showed higher protease activity when the extraction was carried out at pH 4.0 and among the floral organs tested; protease activity was maximum in stamens. The results of the present study indicate that protease activity of the flower may be responsible for wound healing property of the flowers of *J. grandiflorum*.

**Keyword:** *Jasminum grandiflorum*, Wound Healing, Floral Extract, Protease.

1. **Introduction**

*Jasminum grandiflorum* belongs to the family Oleaceae. Although cultivated in subtropical and warm temperate regions for its ornamental qualities, *Jasminum grandiflorum* is valued as a medicinal plant as it is effective against number of ailments The root of the plant is useful in paralysis, mental debility, chronic constipation, flatulence, sterility, ring worm, leprosy and skin diseases[¹]. The leaves of the plant are useful in fixing loose teeth, leprosy, skin diseases, dysmenorrhea, ulcers, wounds and corns[²]. The flowers of *Jasminum grandiflorum* are useful in conditions of leprosy, skin diseases, pruitus, ulcers, dysmenorrhea and vitiated conditions of pitta[³]. The whole flowers of *J. grandiflorum* are used in treatment of wound healing in folk medicine. Protease, one of the key enzyme in wound healing is not so for been reported in flowers of *J. grandiflorum*. Proteases are effective in removing damaged and infected tissues from wounds and thus play an important role in the wound healing process. These enzymes may be used to debride both adherent slough and eschar. Enzymatic debridement is a primary technique in certain cases when surgical debridement is not feasible[⁴]. Proteases from various sources such as plant, microbes, maggots and animals were found to be useful in wound debridement[⁵].

Proteases occur naturally in all living organisms. Growth and development in all organisms occur as a result of an overall balance between protein synthesis and proteolysis. Commercially, proteolytic enzymes from the plant sources have received special attention because of their broad substrate specificity as well as activity in wide range of pH and temperature. As there is no report on the study of protease in *Jasminum*, an attempt was made to study proteases in floral organs of *Jasminum grandiflorum* which may play a role in wound healing property of the plant.
2. Materials and Methods
Flowers of *J. grandiflorum* were collected and stored at -20° C immediately after weighing. Either the whole flowers or different floral organs were used as sample based on the experiment. The total protein content and protease activity were detected in the whole flowers with buffers of different pH range. Citrate phosphate buffer (50 mM) was used for the pH range 4, 5 and 6 and sodium phosphate buffer (50 mM) was used for the pH range 7 and 8. The best extraction buffer was selected to determine the total protein content and protease activity in different parts of the flower such as corolla, stamens and gynoecium along with calyx lobes. The calyx and the gynoecia were taken together for determination of total protein content and protease activity. These two organs usually do not have their contribution in the property of the floral extract as they remain on the plant after plucking out the flower.

2.1 Extraction of Proteins from Flowers
Frozen samples were extracted with buffers of different pH range containing 0.01% (w/v) ascorbic acid. The buffers were used in the ratio 1:5 for extraction. The extracts were filtered through cheese cloth and centrifuged at 20,000 × g for 15 min at 4°C. The clear supernatant was used as the protein and enzyme source.

2.2 Determination of Total Protein Content
The protein content of the floral extracts was determined by the dye binding method [6] with Bovine serum albumin fraction V (Sigma chemical Co., USA) as a standard. To 0.1 ml of enzyme solution, 0.9 ml of distilled water was added and to which 2 ml of Coomassie Brilliant Blue G reagent was added, mixed well and immediately read at 595 nm in a spectrophotometer. Protein content of the unknown sample was calculated from the standard graph. Respective buffer along with the reagent was used as blank.

2.3 Protease Assay
Protease assay was carried out following the procedure of Mc Donald and Chen [7]. The reaction mixture contained a known amount of protein in 0.35 ml of buffer (50 mM, pH 5.0) and 0.35 ml of 0.5% (w/v) casein. The mixture was incubated at 37°C for one hr and the reaction was stopped by adding 0.7 ml of 10% (w/v) ice cold TCA. The undigested substrate was removed by centrifugation in a microfuge for 5 min at 10,000 × g. An aliquot (1 ml) of the supernatant was taken and to which 2.5 ml of the reagent (2.9% Na₂CO₃ and 0.3 N NaOH) and 0.75 ml of Folin Ciocalteu’s phenol reagent (1:3 diluted with distilled water) were added. The samples were incubated at 37°C for 20 min and the liberation of tyrosine was read at 650 nm using a spectrophotometer. One unit of protease activity was defined as the amount of enzyme liberating one micromole of tyrosine equivalent under the assay conditions.

3. Results and Discussion
3.1 Protein Content of Floral Extracts
Total protein content of the flowers of *J. grandiflorum* showed variation in response to different pH of the buffer that was used for extraction. Increase in protein content was observed with increase in pH of the buffer. Maximum protein content was observed with pH 7.0. Further increase in pH showed decline in total protein extraction (Fig 1).
3.2 Protease Activity of Floral Extracts
Maximum protease activity/gfw was obtained when the buffer for extraction was citrate phosphate buffer (50 mM), pH 4.0 which was followed by extraction with citrate phosphate buffer (50 mM), pH 6.0 (Fig 2).

3.3 Protein Content of Floral Organs
As protease activity of flowers of *J. grandiflorum* was more in citrate phosphate buffer (50 mM), pH 4.0, the same buffer was used for extraction of proteins from floral organs separately. Of the floral organs tested, stamens and gynoecium along with the calyx showed higher protein content compared to corolla. Stamens recorded highest protein content of 3.57 mg/gfw (Fig 3).
3.4 Protease Activity of Floral Organs
As far as protease activity in floral organs of *J. grandiflorum* is concerned maximum activity was recorded in stamens and the activity was 286 units/gfw. Protease activity in gynoecia along with the calyx was 163 units/gfw and in corolla it was 120 units/gfw (Fig 4). Thus protease activity of stamens may be responsible for wound healing property of floral extract of *J. grandiflorum*.

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6. References