Evaluation of Healing Potential of Achyranthes aspera L. (Amaranthaceae) seeds in excision, incision, dead space and burn wound model-An in-vivo Study

Sumanta Mondal1, Debjit Ghosh2, Seru Ganapaty1, Motati Sushrutha Reddy1, Karipeddi Ramakrishna2

1 Department of Pharmaceutical Chemistry, GITAM Institute of Pharmacy, GITAM University, Visakhapatnam, Andhra Pradesh, INDIA.
2 Department of Chemistry, GITAM Institute of Science, GITAM University, Visakhapatnam, Andhra Pradesh, INDIA.

ABSTRACT
Objective: The present study deals with the in vivo investigation of healing properties of Achyranthes aspera L. seeds in excision, incision, dead space and burn wound model. Methods: Preliminary phytochemical tests were performed to find out different phytoconstituents in the test extract. Acute toxicity test was performed to find any abnormalities in mice. Wound healing activities were performed on Wistar rats using excision, incision, dead space and burn wound model. The percentage of wound closure and period of epithelization were recorded for excision and thermal burn wound model, whereas the breaking strength was recorded in incision wound model. In dead space wound model, hydroxyproline content and granulation tissue antioxidant enzymes like SOD and CAT were estimated. Results: The phytochemical investigation showed the presence of alkaloids, carbohydrates, tannins, flavonoids, saponins, lipids and triterpenoids. There were no abnormalities observed in the acute toxicity test. The results of wound healing activity revealed that EEAAS treated group showed an enhanced wound contraction and epithelization period in both excision and burn wound models, whereas in incision wound model the wound breaking strength was significantly increased in extract treated group compared to control. The histology of the skin of healed excised wound showed restoration to almost the normal architecture of the skin in extract treated group compared to the control. In dead space wound model, enhanced antioxidant enzymes and connective tissue markers were observed in the extract treated group. Conclusion: Thus the results conclude that EEAAS possesses potent wound healing properties with a reasonable safety profile. 

Key words: Achyranthes aspera L., Antioxidant enzymes, Hydroxyproline, Povidone iodine, Silver sulfadiazine, Thermal burn.

Correspondence: Dr. Sumanta Mondal, Lecturer GITAM Institute of Pharmacy, GITAM University, Visakhapatnam-530045, Andhra Pradesh, INDIA.
Phone no: +91-9703615761
Email: logonchemistry@yahoo.co.in
DOI: 10.5530/pj.2016.3.20

INTRODUCTION
The use of plants for treating different diseases predates human history and forms the source of current modern medicine. Many conventional drugs have been originated from plant sources.1 In the current scenario, herbal medicines are still considered the primary healthcare system in various parts of the world.2 Wound can be defined as a break in the cellular and anatomical architecture of body tissue that includes skin, mucus membrane, deeply lying tissues or surface of internal organs ranging from incision, laceration, abrasion, puncture, and closed wounds such as contusion, hemATOMA and crush.3 Burns injuries are caused by excessive heat, radioactivity, electricity or corrosive chemicals that denature protein in the exposed cells.4 Healing is a complex process which involves a series of cellular and biochemical reactions that are initiated as a response to an injury. The injured tissues are repaired by a sequence of events such as inflammation, proliferation and migration of various cell types.5 Achyranthes aspera L. (Amaranthaceae) is an erect or procumbent, annual or perennial herb of about 1-2 meter in height, often with a woody base. It is usually found on road sides, field boundaries and waste places as a weed and is distributed throughout India, and other tropical areas of the world.6,7,8 Traditionally, the plant is used in asthma and cough, diuretic, purgative and laxative, useful in oedema, piles, boils and eruptions of skin. Crushed plant is boiled in water and is used in pneumonia. The flowering spikes or seeds are grounded and made into a paste with water which is applied externally for bites of poisonous snakes and reptiles. The seeds and flowering spikes are also used in night blindness, cutaneous diseases and injuries.9 This plant is also useful in liver complaints, rheumatism, scabies and other skin diseases and possesses tranquilizing properties.10,11 Several phyto-constituents were reported from the seeds of the plant such as D-Glucuronic acid, β-D-galactopyranosyl ester of D-Glu- curonic acid, oleanolic acid, amino acids, hentriacontane,12,13 10-tricosanone, 10-octacosanone, 4-tritriacontanone.13,14 Three oleanolic acid glycosides were also identified such as α-L-rhamnopyranosyl-(1→4)-(β-D-glucopyranosyluronic acid)-(1→3)-oleanolic acid, α-L-rhamno- pyranosyl-(1→4)-(β-D-glucopyranosyluronicacidicicd)-(1→3)-oleanolic acid-28-O-β-D-glucopyranoside and α-L-rhamnopyranosyl-(1→4)-(β-D-glucopyranosyluronic acid)-(1→3)-oleanonic acid-28-O-β-D-glucopyranoside and α-L-rhamnopyranosyl-(1→4)-(β-D-glucopyranosyluronic acid)-(1→3)-oleanolic acid-28-O-β-D-glucopyranoside.15 Various pharmacological properties such as antimicrobial, antioxidant, antifertility, larvicidal, immunostimulant, hypolipidemic, hypoglycemic, anti-inflammatory, diuretic, antihypertensive, cardiac stimulant, anti spasmodic, anti-anaacra, antiypretic, analgesic, anthelmintic, prothy- rodic, etc., has been reported from various parts of the plant.2 Literature available from all possible scientific sources revealed that the seeds of Achyranthes aspera L. have not yet been screened for wound healing properties, although according to folkloric information it has been used to cure various cutaneous injuries. Thus, the present study is investigated to explore the wound healing properties of ethanol extract of Achyranthes aspera L. seeds (EEAAS) using excision, incision, dead space and burn wound model.
MATERIALS AND METHODS

Collection and authentication of plant materials
The seeds of *Achyranthes aspera* L., was collected from young and matured plant in the month of August, 2012. The plant materials were then authenticated by Dr. M. Venkaiah, Professor (Retd.), Department of Botany, Andhra University, Visakhapatnam, Andhra Pradesh, India. A voucher specimen has been kept in our research laboratory for further reference.

Preparation of extracts
The collected seeds were gently washed in tap water to remove dirt and then they were shade dried in the laboratory under room temperature (24 ± 2°C) for 3-4 weeks. After complete drying, the seeds were pulverized by using a mechanical grinder followed by sieving (sieve no. 40) to obtain a coarse powder. The powdered plant materials (500 g) were defatted with petroleum ether (60-80°C) in a soxhlet extractor. The marc was then air-dried and extracted with ethanol (90%), excess solvents were removed by rotary evaporator (Evator, Media Instrument Mfg. Co., Mumbai, India) and concentrated to obtain a dark greenish residue. The percentage yield of the ethanolic seed extract was calculated by the following formula.

\[
\text{Percentage yield} = \frac{\text{weight of dry crude extract obtained (g)}}{\text{Weight of plant material before extraction (g)}} \times 100
\]

Preliminary phytochemical tests
Preliminary phytochemical studies of EEAAS were performed for the determination of major phytochemical constituents using standard procedures.16,17

Experimental animals and housing conditions
Swiss albino mice (20-25 g) of either sex were used for acute toxicity studies and Wistar albino rats (150–250 g) both male and female were used to evaluate wound healing potential. The animals were housed for at least one week in the laboratory animal room prior to testing and maintained in clean polypropylene cages with optimum light, temperature and humidity (light/dark cycles (12/12 h), Temp: 25 ± 2°C, and 75% relative humidity) and fed with commercially pelleted rat diet (M/s Hindustan Lever Ltd., Mumbai) and water *ad libitum*.

Ethical approval
All experimental protocols were approved by the Institutional Animal Ethics Committee (IAEC) of GITAM Institute of Pharmacy, Visakhapatnam, Andhra Pradesh, India (Registration No.1287/ac/09/CPCSEA and Protocol No: IAEC/GIP-1287/Bpharm/IP/SM-NS/11/2012-13). Experiments were performed according to the guide for the care and use of laboratory animals.

Acute toxicity study
The acute toxicity studies were conducted over Swiss albino mice as per OECD guidelines 423,18 where the limit test dose of 2000 mg/kg, p.o., was used. Observations were recorded continuously for the first 4 h for any general behavioural changes. They were then kept under observation up to 14 days after drug administration to find out the mortality if any. One-tenth of the maximum tolerated dose of EEAAS (200 mg/kg, body weight, p.o.) was selected for dead space wound model study.

Experimental grouping and dosing for evaluation of wound healing activity
The rats were acclimatised for one week prior to use in the experimental models. The selected animals were subdivided into three groups of six animals each (n=6 per group) for evaluating healing potential on excision, incision, and burn wound models. Group I: treated with simple ointment I.P. (vehicle), Group II: treated with reference standards. Group III: treated with test ointment which was prepared by 10% w/w EEAAS in Simple ointment I.P. Povidone iodine ointment (5% w/w) (Cipladine; mfg by Jeps Pharmaceuticals, Sirmour-173025 (H.P.)), Batch No: JM 92) was used as the reference standard for excision and incision wound model, whereas silver sulfadiazine (1% w/w) was used as the reference standard for thermal burn injury model. For evaluating dead space wound model twelve animals were randomly divided into two groups of six animals each (n=6 per group). Group I serve as control, which received only plain drinking water (3 mL/kg, p.o.) and group II were separately administered with EEAAS orally in water at a dose of 200 mg/kg, p.o., daily for 10 days.

Excision wound model
Investigation of wound healing activities using excision wound model on Wistar rats were performed according to the methods described by Morton and Malone, 1972.19 Prior to creation of the wounds, the animals were anesthetized with 1 mL of intravenous ketamine hydrochloride (10 mg/kg, body weight). An impression was made on the dorsal thoracic region 1 cm away from vertebral column and 5 cm away from ear on the rat. The dorsal fur of the animals was shaved with an electric clipper and the predicted area of the wound to be created was outlined on the back of the animals with methylene blue using a circular stainless steel stencil. Toothed forceps, scalpel and pointed scissors were used to create a full thickness of the excision wound of circular area of 500 mm² and 2 mm depth along the markings. Haemostasis was achieved by blotting the wound with cotton swab soaked in normal saline. The entire wound was left open. All surgical procedures were done under aseptic conditions. The wounds were left untreated for a period of 24 h. The test extracts and reference drugs were applied topically at the wound site twice a day. The wound closure rate was assessed by tracing the wound on 5th, 10th and 15th post wounding days using transparent paper and a permanent marker. Wound area was measured by retracing the wound on a millimeter scale graph paper. The percentage of wound closure was calculated using the formula.20,21

\[
\% \text{ wound closure} = \frac{\text{Wound area on day 0} - \text{Wound area on day n}}{\text{Wound area on day 0}} \times 100
\]

Where n=5th, 10th and 15th post wounding days.

The period of epithelialization was also calculated as the number of days required for falling of the dead tissue remnants without any residual raw wound.

Histomorphological analysis of wound tissue from healed excised wound
The cross-sectional full thickness of wound tissues were cut on the 16th day post wounding from all the groups to study the histopathological changes of the healed excised wound tissue. The tissue samples were fixed in Bouin’s solution and were dehydrated through increasing grades of ethanol and then embedded in paraffin wax. The tissue were then cut to 5 μm sections with a rotary microtome, deparaffinised, mounted on clean glass slides and stained with haematoxylin and eosin (HE). The glass slides were then observed under the microscope for histomorphological changes.22,23
Incision wound model

Incision wound model was performed according to the methods described by Ehrlich and Hunt. The rats were anesthetized with 1 mL of intravenous ketamine hydrochloride (10 mg/kg, body weight) prior to and during creation of the wounds. The dorsal fur of the rats was shaved with an electric clipper and a longitudinal paravertebral incision of 6 cm long was made through the skin and cutaneous tissue on the back. After the incision, the part skin was sutured 1 cm apart using a surgical thread and curved needle. The wounds were left open and undressed. The test extract was topically applied over the wound once a day. The sutures were removed on 8th post wound day and the application of the test extract to the wounds continued till 10th post wounding day. The wound breaking strength was measured on the 10th day evening after the last application of the test sample and standard.

Determination of wound breaking strength

The breaking strength of the wound on each animal was measured by the constant water flow method. The rats were anesthetized and transferred to the operation table and a line was drawn 3 mm away from the edge of the wound on either side. Two forceps were firmly applied on to the line facing each other on opposite side of the incision wound. One of the forceps was fixed on stands, while the other was connected to a freely suspended lightweight polypropylene graduated container through a string run over to a pulley. Water was allowed to flow from the reservoir slowly and steadily into the container. A gradual increase in weight was transmitted to the wound site pulling apart the wound edges. As and when the wound just opened up, the water flow was stopped and the volume of water collected in the container (approximately equal to its weight) was noted. The readings were recorded for a given incision wound and the procedure was repeated on the wound on the contra lateral side. The average reading of the group was taken as an individual value of breaking strength and the mean value gives the breaking strength for a given group.

Dead space wound model

Prior to and during creation of the wounds the rats were anaesthetized with 1 mL of intravenous ketamine hydrochloride (10 mg/kg, body weight). After anaesthetisation, 1 cm incision was made on dorso-lumbar part of the back. Two pre-weighted sterilized polypropylene tube (2.5 length×0.25 cm diameter) were placed in the dead space of lumbar region of rat on each side and wounds were closed with a suture material. On the 10th post wounding day, the granulation tissue formed on the implanted tubes was carefully detached from surfaces of the tubes. The wet weight of the granulation tissue collected was recorded. The sample tissues were dried at 60°C for 12 h and weighed to determine the dry granulation tissue weight. Connective tissue parameter such as hydroxyproline content in granulation tissue was also determined by standard methods. Part of the granulation tissue was collected in phosphate-buffered saline for the estimation of antioxidant enzymes superoxide dismutase (SOD) and catalase (CAT).

Estimation of hydroxyproline

On the 10th day, a piece of skin from the wound area was taken and analysed for hydroxyproline content. Hydroxyproline is a basic constituent of collagen. The tissue samples taken from the wound area was dried in hot air oven at 60°C for 12 h to get dry granulation tissue which were weighed and kept in glass stoppered test tubes. To each tube containing about 40 mg dried granulation tissues, 6 N HCl (1.0 mL for 100 mg sample) was added and then kept on boiling water bath (110°C) for 24 h (12 h each day for 2 days) for hydrolysis. The hydrolysate was then cooled and the excess acid was neutralized by 10 N NaOH using phenolphthalein as indicator. 1.0 mL of acid hydrolysate sample was mixed with 1 mL each of 0.01 M copper sulfate solution, 2.5 N sodium hydroxide, and 6% hydrogen peroxide. The solution were then mixed thoroughly and incubated for 5 min at 80°C. After incubation period, the tubes are chilled in an ice water bath, and 4 mL of 3 N sulfurous acid were added with agitation followed by addition of 2 mL of p-dimethylaminobenzaldehyde (5%) solution and mixed thoroughly. Then the tubes were warmed in a water bath for 15 min at 70°C and then cooled in an ice water bath. Then the absorbance was recorded at 540 nm.

Estimation of granulation tissue antioxidants in wet tissue

Superoxide Dismutase (SOD)

The inhibition of the formation of NADH-phenazine methosulphate-nitro blue tetrazolium formazan was measured. 0.6 mL of 0.052 M sodium pyrophosphate buffer (pH 8.3), 50 μL of 186 μM of PMS, 150 μL of 300 μM NBT, and 0.4 mL of distilled water was added to 0.2 mL of tissue homogenate. Reaction was initiated in the above by the addition of 0.1 mL of 780 μM NADH and incubated for 60 sec at 30°C. The reaction was stopped after the incubation period, by the addition of 0.5 mL of glacial acetic acid. Then the reaction mixture was stirred vigorously and shaken with 2 mL of n-butanol and then the mixture was allowed to stand for 10 min, centrifuged at 3000 rpm for 10 min and butanol layer was taken out. Colour intensity of the chromogen in the butanol was measured at 560 nm in spectrophotometer against n-butanol, a system devoid of enzyme that served as control. The results were expressed as IU/mg protein.

Catalase (CAT)

The granulation tissue is homogenized at 1-4°C in M/150 phosphate buffer and centrifuged at 1000 rpm for 30 mins. The supernatant obtained from centrifugation was diluted to 1:10 with water and 0.04 mL was taken for the assay. Decomposition of hydrogen peroxide in presence of catalase was measured at 240 nm. The results were expressed in nU/mg protein.

Thermal burn wound model

The hairs on the dorsal skin were shaved mechanically 24 h before induction of burn. Then the animals were left for 24 h for observing any inflammation caused by shaving. Thermal burn injuries were induced on dorsal skin of the rat by the pressing of metal rod (2.5 cm diameter) heated to 80-85°C for 20 sec. Then the wound area was dressed with sterile gauge and after the animals recovered from anaesthesia the animals were housed separately. Drugs were applied twice daily upon the burn. The wound closure rate was recorded on 5th, 10th, and 15th post wounding days using transparent paper and a permanent marker. The percentage of wound closure was calculated for final analysis of results according to the following formula.

\[ \text{Wound area on day 0} - \frac{\text{Wound area on day n}}{\text{Wound area on day 0}} \times 100 \]

Where n = 5th, 10th, and 15th post wounding days

Statistical analysis

The data obtained in the studies were subjected to one way of analysis of variance (ANOVA) for determining the significant difference. The inter group significance was analyzed using Dunnet's t-test. A p-value <0.05 was considered to be significant. All the values were expressed as mean ± SEM.
RESULTS

Percentage yield and phytochemical analysis of EEAAS

After complete drying of ethanol extract of Achyranthes aspera L., seeds (EEAAS) yielded about 9.8% of extract. Preliminary phytochemical screening of EEAAS for detection of different phytochemicals revealed that the extract contains alkaloids, carbohydrates, tannins, flavonoids, saponins, lipids and triterpenoids (Table 1).

<table>
<thead>
<tr>
<th>Test groups</th>
<th>Inference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaloids</td>
<td>+</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td>+</td>
</tr>
<tr>
<td>Gums and mucilages</td>
<td>-</td>
</tr>
<tr>
<td>Proteins and amino acids</td>
<td>-</td>
</tr>
<tr>
<td>Tannins and phenolic compounds</td>
<td>+</td>
</tr>
<tr>
<td>Steroids and sterols</td>
<td>+</td>
</tr>
<tr>
<td>Triterpenoids</td>
<td>-</td>
</tr>
<tr>
<td>Saponins</td>
<td>+</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>+</td>
</tr>
<tr>
<td>Lipids</td>
<td>+</td>
</tr>
</tbody>
</table>

(-) Absent, (+) Present

Acute toxicity studies

Through the 14 day period following single oral administration no mortality or morbidity was observed in animals. Morphological characteristics like fur, skin, eyes and nose appeared normal. No salivation, diarrhoea, lethargy or unusual behaviours such as self mutilation, walking backward etc., were observed. Gait and posture, reactivity to handling or sensory stimuli, grip strength was all normal. Daily fluctuations of food and water intake were observed within the range of control animals. This indicates that the ethanol extract of Achyranthes aspera L., seeds was safe to a single dose of 2000 mg/kg, body weight. Hence 200 mg/kg oral doses of EEAAS were selected to evaluate dead space wound model.

Effect of EEAAS on healing of Excision wound

The results of excision wound healing model revealed that the ethanol extract from Achyranthes aspera L., seeds demonstrated significant (P<0.05) healing potency compared to the control. However, the reference drug Povidone iodine ointment (5% w/w) exhibited higher significant (P<0.01) potency than the crude extract (Figure 1). The percentage wound closure of the crude extract was 96.02 ± 1.33, while the reference drug achieved a percent wound closure of 97.23 ± 1.62 on the 15th day of the study. The average number of days that took for the shedding of eschar without leaving any residual raw wound (epithelialization period) was also significantly reduced in the standard drug (P<0.01) and EEAAS treated groups (P<0.05) which was 16.63 ± 0.09 and 17.25 ± 1.88 respectively when compared with control group (29.86 ± 3.81).

Histomorphological studies of excision biopsy of healed skin at the 15th day showed almost healed skin architecture with normal epithelization, restoration of adnexa, and fibrosis within the dermis in both reference standard Povidone iodine (5% w/w) and EEAAS (10% w/w) treated groups when compared to control group (Figure 2).

Effect of EEAAS on healing of Incision wound

The results of incision wound model revealed that reference standard Povidone iodine (5% w/w) and EEAAS (10% w/w) showed significant (P<0.01) increase in breaking strength when compared to the control (Figure 3). The wound breaking strength for reference standard Povidone iodine (5% w/w) and EEAAS (10% w/w) was 385.03 ± 4.91 g and 303.32 ± 4.33 g respectively, whereas the breaking strength for control group was 108.26 ± 8.28 g. Thus the increase in wound breaking strength in extract treated group shows the healing effect of EEAAS in incision wounds.

Effect of EEAAS on healing of dead space wound

In dead space wound model the weight of dry and wet granulation tissue were significantly (P<0.01) increased in EEAAS treated group when compared to the control.
The healing of wound is a dynamic and complex process which takes place by restoring tissue layers and cellular structures in damaged tissue. The wound contraction rate which was enhanced by EEAAS treatment was may be due to enhanced contractile property of myofibroblast.

**DISCUSSIONS**

Acute toxicity is considered an initial study on the safety assessment of the drug and it is usually performed to provide us the initial information about the mode of toxic action of a substance by which we can fix a dose of a new compound and help in dose determination in animal studies. In our acute toxicity study a single administration of EEAAS did not produce any abnormities in animals. The healing of wound is a dynamic and complex process which takes place by restoring tissue layers and cellular structures in damaged tissue. The wound healing process can be divided into three phases like inflammatory phase, proliferation phase, and maturation phase. This healing process is dependent on the general state of the host’s health, type and extent of damage and the ability of the tissue to repair. The inflammatory phase is the body’s natural response to injury. This phase is characterised by inflammation and hemostasis. During the proliferation phase, the wound is rebuilt with new granulation tissue which is comprised of collagen and extracellular matrix and into which a new network of blood vessels develop, a process known as angiogenesis. Epithelial cells finally resurface the wound, a process known as epithelialisation. Maturation is the final phase and occurs once the wound has closed. This phase involves remodelling of collagen. In this study four different models were used to evaluate the effect of EEAAS on various phases of wound healing.

**Effect of EEAAS on healing of thermal burn wounds**

The result shows that in thermal induced burn wound there is a significant increase in percentage wound closure in EEAAS (10% w/w) and Silver Sulfadiazine (1% w/w) treated groups when compared with control group. The maximum percentage wound closure was observed on the 15th day of study where it was 96.4 ± 1.25 and 97.1 ± 1.68 in EEAAS (10% w/w) and silver sulfadiazine (1% w/w) treated group respectively. Thus the percentage wound closure increased with increase in number of days (Figure 4). The period of epithelization was recorded for both silver sulfadiazine (1% w/w) and EEAAS (10% w/w) treated group respectively. The wound contraction rate which was enhanced by EEAAS (10% w/w) treated groups which was 17.23 ± 0.81 and 18.44 ± 0.26 days respectively.

**DISCUSSIONS**

Acute toxicity is considered an initial study on the safety assessment of the drug and it is usually performed to provide us the initial information about the mode of toxic action of a substance by which we can fix a dose of a new compound and help in dose determination in animal studies. In our acute toxicity study a single administration of EEAAS did not produce any abnormities in animals. The healing of wound is a dynamic and complex process which takes place by restoring tissue layers and cellular structures in damaged tissue. The wound healing process can be divided into three phases like inflammatory phase, proliferation phase, and maturation phase. This healing process is dependent on the general state of the host’s health, type and extent of damage and the ability of the tissue to repair. The inflammatory phase is the body’s natural response to injury. This phase is characterised by inflammation and hemostasis. During the proliferation phase, the wound is rebuilt with new granulation tissue which is comprised of collagen and extracellular matrix and into which a new network of blood vessels develop, a process known as angiogenesis. Epithelial cells finally resurface the wound, a process known as epithelialisation. Maturation is the final phase and occurs once the wound has closed. This phase involves remodelling of collagen. In this study four different models were used to evaluate the effect of EEAAS on various phases of wound healing. In excision wound model, EEAAS and reference standard (Povidone iodine) treated groups showed enhanced wound closure rate and period of epithelization when compared with control group. This enhanced epithelization may be due to the effect of EEAAS on enhanced collagen synthesis. Histomorphology of the skin area of excised wound on 15th day post wounding showed normal epithelization, restoration of adnexa and fibrosis within the dermis in both EEAAS and standard treated groups. Wound contraction is the process of mobilizing the healthy skin around the wound to cover the denuded area. It usually takes place by the centripetal movement of the wound margin caused due to the activities of myofibroblast. The wound contraction rate which was enhanced by EEAAS treatment was may be due to enhanced contractile property of myofibroblast or increased number of myofibroblast entering the wound area. In incision wound, EEAAS showed an increase in wound breaking strength compared to control which may be due to the increase in...
REFERENCES


CONCLUSIONS

The present study has demonstrated that an ethanol extract of Achyranthes aspera L., seeds (EEAS) possesses promising wound healing properties which were evidenced by enhanced rate of wound contraction and epithelization period in excision and thermal burn wound model. In incision wound model, wound breaking strength were also increased in EEAS treated groups, whereas enhanced antioxidant enzymes (SOD and CAT) and hydroxyproline content in dead space wound model indicates enhanced wound healing in EEAS treated groups. Thus, the wound-healing properties of EEAS may be attributed to the phytoconstituents they contain, which may be either due to their individual or additive effect that fastens the process of wound healing. However, further phytochemical studies are needed to isolate the active compound(s) responsible for the healing activities.

ACKNOWLEDGEMENTS

We are thankful to GITAM University, Visakhapatnam, Andhra Pradesh, India for providing financial support and facilities to carry out this research.

CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

ABBREVIATIONS USED


PICTORIAL ABSTRACT

SUMMARY

- *Achyranthes aspera* L. seeds have not yet been evaluated for wound healing properties whereas the seeds of the plant are used traditionally to treat skin and other cutaneous injuries and infections.
- The wound healing properties of *Achyranthes aspera* L. seeds were evaluated in four different models thus a detailed report of wound healing properties is given.
- *Achyranthes aspera* L. seeds have a reasonable safety profile.
- As per the results of the excision, incision, dead space and burn wound model, *Achyranthes aspera* L. seeds possess potent wound healing activity.

ABOUT AUTHORS

Dr. Sumanta Mondal: (Lecturer & NSS Programme Officer of GITAM Institute of Pharmacy, GITAM University, Andhra Pradesh, India). His research involves bioactivity and phytochemical studies of various medicinal plant species. He has published more than 48 research articles in various international and national journals. He has guided more than 20 M. Pharm students and presently seven students are pursuing PhD under his guidance.

Mr. Debjit Ghosh: Pursuing PhD from Department of Chemistry, GITAM Institute of Science, GITAM University, Visakhapatnam, Andhra Pradesh, India, under the guidance of Prof. K. Rama Krishna & Dr. S. Mondal. His area of expertise and research interest includes isolation and structural elucidation of phyto-constituents, chromatographic and phytochemical analysis, toxicological studies and pharmacological screening.

Prof. Seru Ganapaty: (Dean & Principal of GITAM Institute of Pharmacy, GITAM University, Andhra Pradesh, India). He had 35 years of teaching and research experience. He supervised 35 doctoral students leading to Ph.D degrees and successfully handled six major research projects on Indian medicinal plants. To his credit he has two patents and more than 150 research publications in peer reviewed national and international journals. He also received a good number of awards in recognition of his research contribution in the area of natural products.