ANTI-INFLAMMATORY ACTIVITY OF ROOT OF DECALEPSIS HAMILTONII

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Received 01 Dec 2012; Review Completed 04 Jan 2013; Accepted 12 Jan 2013, Available online 15 Jan 2013

ABSTRACT:

Powdered root of Decalepsis hamiltonii was used to evaluate the acute & chronic anti-inflammatory activities in rodents. Albino rats of either sex weighing between 150-200 grams were used for this experiment. The rats were kept in pollysulfone rat cages and maintained at a 12 h light / dark cycle. The temperature was maintained at 22 - 26°C while the relative humidity was 50-60%. The experiment was conducted in the Department of Pharmacology, Medical College Kolkata in collaboration with West Bengal University of Animal and Fishery Sciences, Mohanpur Campus. Rats were fed on standard pellet diet and provided filtered autoclaved ad libitum drinking water. The experiments were performed following approval by the Institutional Animal Ethics Committee of WBUAFS.

INTRODUCTION:

Decalepsis Hamiltonii (Wight& Arn), popularly known as sarsaparilla, is a climber with stout, smooth branches shrub and a native of the forests of Deccan peninsula and Western Ghats of India. The leaves are curvaceous, orbicular or elliptical, with rounded tip. Its tubers are consumed as pickles and the juice for its alleged health promoting properties The Root has a sweet sarsaparilla-like taste; contains 9.2% fleshy matter and 8% woody core. It contains quercetin, kaempferol, coumarin and rutin and considered as “Sariva Bheda” in Ayurveda where these find use as an alternative to the roots of Hemidesmus indicus in the preparation of several herbal drugs like Amrutasalaka taila (hair tonic), Drakshadi churna (general vitalize), Shatavari rasayana (adapagenic) and Yeshtimadhu taila (mild analgesic, anti rheumatism). The root can be stored for longer period and remains unaffected by microorganisms and insects, apparently due to the presence of a volatile principle with bacteriostatic and toxic properties which can be isolated by steam distillation. It has been extensively used as preservative and also as a source of bio insecticide in stored food grains due to the presence of strong aroma and 4-0-methyl resorcyldaldehyde (0.9-12%).

Although the roots of D. hamiltonii have been used for their alleged health benefits, scientific investigation in this regard need to be done. Though many researchers establish the anti oxidant, anti bacterial and anti ulcerogenic role of the root extract of D. hamiltonii, very few have explored the anti inflammatory potential. The anti-inflammatory drugs (NSAIDs) presently available are not free of side effects like gastric ulceration, kidney damage etc so an alternative drug obtained from natural sources devoid of side effects would be useful in treatment of acute and chronic inflammatory diseases. The aim of the present study was therefore to evaluate the acute & chronic anti-inflammatory activities of the extract of root of Decalepsis Hamiltonii in rodents.

MATERIALS & METHOD:

Animals

Albino rats of either sex weighing between 150-200 grams were used for this experiment. The rats were kept in polysulfone rat cages and maintained at a 12 h light / dark cycle. The temperature was maintained at 22 - 26°C while the relative humidity was 50-60%. The experiment was conducted in the Department of Pharmacology, Medical College Kolkata in collaboration with West Bengal University of Animal and Fishery Sciences, Mohanpur Campus. Rats were fed on standard pellet diet and provided filtered autoclaved ad libitum drinking water. The experiments were performed following approval by the Institutional Animal Ethics Committee of WBUAFS.

Plant Extract

Powdered root of Decalepsis Hamiltonii (5 Kg) was obtained from the Chemical Research and Extraction. Supply Unit of CCRAS and extracted with petrol ether (60 – 80 °C) in sox let apparatus for 15 hours and solvent was removed by distillation. The residual gummy material obtained was then suspended in 5% gum acacia and used for oral administration to the experimental animals at a dose of 250 mg/ kg body wt after standardizing according to their LD 50 & ED 50 determined before selecting the dose.
**Drugs and Chemicals:**
Carrageena (σigma), 5HT (Roche Pharmaceuticals), Bradykinin (BRA 640, San Doz), Formaldehyde, Phenylbutazone (SG Pharmaceuticals), Dexamethasone (MSD), Gum acacia.

**Procedure:**
As the indigenous drug was insoluble in water it was suspended in 5% gum acacia solution. The rats were divided in five groups (n=6) and the drugs were given orally with the help of rat feeding canula fitted to 2 ml glass syringe keeping the volume of medicament constant (1ml) one hour prior to Carrageenan injection (for inducing inflammation).

GR I served as control (receiving gum acacia only)  
GR II received root extract of D. hamiltonii, 250mg/kg  
GR III received root extract of D. hamiltonii, 500mg/kg  
GR IV received Phenylbutazone 50 mg / Kg  
GRV received Dexamethasone 0.5 mg / Kg

The average volume of hind paw of all the rats was measured with the help of a plethysmograph by the method of Buttle et al after some modification and noted before inducing acute inflammation with Carrageenan.

**Paw edema induced to produce acute inflammation:**
Carrageenan, 5HT and bradykinin induced edema was adopted as introduced by Winter, Risley and Nuss with slight modification by Ghosh & Singh. Carrageenan 1% in 0.9% w/v Sodium Chloride was taken and 0.1 ml of such solution was injected in sub planter region of hind paw of all the rats one hour after the administration of the drugs. The paw volume was again measured twice after 1 hour and at the end of 3 hours and noted. Increase in volume of paw edema was recorded by subtracting the initial paw volume from final paw volume measured with the help of Plethysmograph after 1 hour and 3 hours of Carrageenan injection.

Percentage of inhibition of inflammation rate was calculated as

\[ \frac{V_c - V_t}{V_c} \times 100 \]

Where

Vc- mean increase in volume of paw edema in control group.  
Vt- means increase in paw volume of treated group of animals.

Same procedure is followed after inducing acute inflammation with sub planter injection of 0.1 ml of 1% sol of 5HT and 0.1 ml 1 micro gram of bradykinin respectively.

**Paw edema induced to produce chronic / immunologically induced inflammation (arthritis model)**
A new set of rats were taken and grouped in a similar manner and the established method was followed to produce chronic inflammation. A volume of 0.1 ml of 2% formaldehyde (arthritis model) was injected into the sub planter region of the left hind paw of each rat. The degree of inflammation was assessed by measuring the paw volume daily for 13 days according to the method of Buttle et al. As in acute inflammation, Groups I, II, III, IV received the drugs once daily orally in similar dosage for 13 days. The degree of inflammation was tabulated on alternate days till the 13th day and compared as shown in Table IV. Percentage of inhibition of inflammation rate was calculated in a similar manner.

**Statistical analysis:**
The results were obtained in mean± SE and comparison between the control and experiment groups was done. Statistical analysis was carried out by one-way analysis of variance (ANOVA) and P<0.05 was considered as significant.

**RESULTS:**
The effects of Decalepsis hamiltonii, Phenylbutazone and Dexamethasone on Carrageenan, 5HT and bradykinin induced edema of rat hind paw is shown in Table no I, II & III respectively and Graph I& II. There was inflammatory inhibition of 35.55% in carrageenan induced, 42.62% in 5HT induced and 36.62% in bradykinin induced acute inflammatory models with study compound. In the formaldehyde induced chronic inflammatory model, a progressive inhibition of 36.84% (3rd day), 36.45% (5th day), 51.45% (7th day) and 66.39% (13th day) was observed with study compound. The efficacy was comparable to the standard drugs phenylbutazone and Dexamethasone. (Table IV)

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean increase in paw volume (in ml ± SE)</th>
<th>% inhibition</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>GR I</td>
<td>0.9 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GR II</td>
<td>0.58 ±0.06</td>
<td>35.55</td>
<td>P&lt;.05</td>
</tr>
<tr>
<td>GR III</td>
<td>0.43±0.11</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GR IV</td>
<td>0.23 ± 0.02</td>
<td>74.44</td>
<td>P&lt;.001</td>
</tr>
<tr>
<td>GR V</td>
<td>0.3 ± 0.03</td>
<td>66.66</td>
<td>P&lt;.01</td>
</tr>
</tbody>
</table>

Table-I: Effects of Decalepsis hamiltonii, Phenylbutazone and Dexamethasone on carrageenan induced edema of rat hind paw (n=6)
TABLE - IV: Effects of Decalepsis hamiltonii, Phenylbutazone and Dexamethasone on Formaldehyde induced paw edema in rats (n=6)

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean increase in Paw Volume (ml ± SE)</th>
<th>3rd Day</th>
<th>5th Day</th>
<th>7th Day</th>
<th>9th Day</th>
<th>11th Day</th>
<th>13th Day</th>
</tr>
</thead>
<tbody>
<tr>
<td>GR I</td>
<td>0.95 ± 0.02</td>
<td>0.96 ± 0.02</td>
<td>1.03 ± 0.05</td>
<td>1.16 ± 0.12</td>
<td>1.23 ± 0.13</td>
<td>1.13 ± 0.09</td>
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<tr>
<td>GR II</td>
<td>0.71 ± 0.03</td>
<td>0.66 ± 0.04</td>
<td>0.63 ± 0.04</td>
<td>0.6 ± 0.03</td>
<td>0.55 ± 0.03</td>
<td>0.5 ± 0.05</td>
<td>55.75%</td>
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<tr>
<td></td>
<td>25.26%</td>
<td>31.25%</td>
<td>38.83%</td>
<td>48.27%</td>
<td>P &lt; .01</td>
<td>P &lt; .01</td>
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<td></td>
<td>P &lt; .001</td>
<td>P &lt; .001</td>
<td>P &lt; .001</td>
<td>P &lt; .001</td>
<td>P &lt; .001</td>
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<tr>
<td>GR III</td>
<td>0.7 ± 0.03</td>
<td>0.65 ± 0.02</td>
<td>0.62 ± 0.04</td>
<td>0.58 ± 0.03</td>
<td>0.5 ± 0.04</td>
<td>0.4 ± 0.04</td>
<td>64.60%</td>
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<td></td>
<td>26.32%</td>
<td>32.29%</td>
<td>39.80%</td>
<td>59%</td>
<td>P &lt; .01</td>
<td>P &lt; .01</td>
<td>P &lt; .001</td>
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<td></td>
<td>P &lt; .001</td>
<td>P &lt; .001</td>
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<td>P &lt; .001</td>
<td>P &lt; .001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GR IV</td>
<td>0.60 ± 0.06</td>
<td>0.61 ± 0.06</td>
<td>0.50 ± 0.06</td>
<td>0.46 ± 0.04</td>
<td>0.43 ± 0.04</td>
<td>0.38 ± 0.03</td>
<td>66.39%</td>
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<tr>
<td></td>
<td>36.84%</td>
<td>36.45%</td>
<td>51.45%</td>
<td>60.34%</td>
<td>P &lt; .01</td>
<td>P &lt; .01</td>
<td>P &lt; .001</td>
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<td>P &lt; .001</td>
<td>P &lt; .001</td>
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</table>

DISCUSSION:

Medicinal plants are nature’s hidden and to a large extent unexplored treasure. They have been used as a source of safe and effective medicine since time immemorial, as they are less toxic, cheap and suitable for use over a prolonged period. This potential resource has hardly been commercially tapped. India is endowed with about 8000 species of medicinal plants. According to a recent estimate of the Planning Commission, Government of India, the potential for plant-based crude drugs is about Rs. 400 billion. Globally, the demand for medicinal plants and their derivatives is growing at a rate of 7-15%. Only a few medicinal plants have been selected for scientific testing and experimentation as inflammatory agents. Moreover, the active principles responsible for the alleged health promoting activity need to be isolated to elucidate their exact mode of action.

Chopra6 pioneered the usefulness of indigenous drugs various diseases including arthritic conditions. Earlier works have shown that the D. hamiltonii roots contain aldehydes, amyrins, lupelos and volatile flavour compounds such as 2-hydroxy-4methoxybenzaldehyde, vanillin etc and essential oil like 4-methylresorcyaldehyde, atlantone, terpinene, geraniol etc. A combinational molecule containing pectic polysaccharide with bound phenolics identified in the root of D. hamiltonii and their break down products have been known to have health beneficial properties.7 This highly aromatic root has been subjected to over exploitation by destructive harvesting that has endangered the survival of this plant. A method for rooting of D. hamiltonii for field transfer is reported8.

Flavonoids and occasionally polysaccharides present in various plant products have frequently been implicated as antioxidant and antiulcer agents. Various workers have proved the antibacterial activity supercritical extract of swallow root10. Some claim that it is gastro protective in nature and possibly inhibits the H’K’ ATPase enzyme responsible for HCl secretion in stomach.11 while some state that it has multitope ulcer protective activity,12 and antioxidantive properties13. The inhibition of carrageenan-induced inflammation in rats is an established model to screen compounds for potential anti-inflammatory activity. According to Vinegar et al14 the development of carrageen induced edema is biphasic; the first phase occurs within one hour of carrageenan administration and attributed to the release of cytoplasmic enzymes, histamine and serotonin, from the mast cells. The second phase (>1.0 h) is mediated by an increased release of prostaglandins in inflammatory area and continuity between the two phases is maintained by kinins. Taken together, it suggests root of D hamiltonii possesses potent anti-inflammatory activity (Table I-III) possibly due to inhibition and/or release of inflammatory mediators, principally the prostaglandins. Anti inflammatory effect is also seen in inflammation produced by an i...
by 5HT and Bradykinin, that proves it may have some action by inhibiting the pain pathway that caused by 5HT and Bradykinin.

Formaldehyde induced arthritis is generally used to study the efficacy of a drug against the proliferative phase of inflammation i.e. chronic inflammation. There was significant inhibition of formaldehyde induced paw edema by Decalepis hamiltonii with the increase of number of days which suggests that it would be beneficial in treatment of arthritis and related disorders. Results of the present study corroborate with Lakshman et al15 who examined the effectiveness of the methanol extract of roots of Decalepsis hamiltonii at 250 and 500 mg/kg doses orally in the carrageenan-induced rat paw edema and cotton pellet-induced chronic inflammatory models. The extract showed significant dose-dependent anti-inflammatory activities in both models.

Though efficacy of phenylbutazone was the most in acute inflammatory model and dexamethasone was most efficacious in chronic inflammation, study compound exhibited a comparable result in both the models.

ACKNOWLEDGEMENT:
I want to thank all the members of department of Pharmacology, Medical College, Kolkata for their constant support to carry out my experiment.

Conflict of Interest: Nil.