A RELATIVE ASSESS ON WOUND HEALING AND ANTI SCAR ACTIVITY OF CRUDE *Echinops heterophyllus* EXTRACT AND SOME OF ITS BIOACTIVE FRACTIONS

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INTRODUCTION

A large number of plants has been used by folk medicine at diverse parts of Iraq to treat different human ailments. *Echinops heterophyllus* P.H. Davis (Compositae), is naturally inhabitant in the Kurdistan region, it has been traditionally used for various skin problems including cuts, abrasion, burns and snake bites. 

Echinops, is a genus includes many plants which are independently referred to as globe thistle. Echinops has more than 120 species of perennials, annuals, and biennials[1,2]. The genus belongs to the daisy family Asteraceae, and its species are found in eastern and Southern Europe, North Africa and Asia[3]. Prof. Dr. Ali Al-Raw an an Iraqi plant classification specialist mention 11 species of Echinops in his book[4], one of them was *Echinops heterophyllus* P.H. Davis. This plant is indigenous in Iraq mainly in the Kurdistan region, especially in Erbil and Suhimani cities. In Hanara village and surrounding area in Wadi Bastora and shaklawa in Erbil governorate, the plant is called (Shakroka). The term [Shakroka] is come from that the circle-like part of the plant, before getting harder in the late spring, is eaten and the taste is sweet, therefore, it is called shakroka. Shakr means sugar , Shakroka------ sweet like sugar (in local Iraqi language). The *Echinops heterophyllus* (figure-1) is a perennial, 40-100 cm high. Stems are simple or branching from the base, sparsely cobwebby-cane scent. Leaves are lanceolate or oblong-lance late, the lower ones are 10-15 cm long, 4-6cm wide, with triangular-lance late, prickly lobes, greenish, shiny, sub glabrous above, densely whitish-tomentose below[3] Echinops plant was reported to possess variety of compounds belonging to various classes like: alkaloids, flavonoids, terpenoids, lipids, steroids and polycylenes [6]. And many literatures survey revealed different pharmacological activities of *Echinops* plant like, antibacterial activity[7], Antifungal [8], Antioxidant activity [9], Protective effects on testosterone-induced prostatic hyperplasia [10], Hepato-protective[11] and anti-ulcerogenic activity[12]. So this study was emphasized on the identification of active constituents in the Iraqi species of *Echinops* plant, and investigated the pharmacological activity of this plants crude extract and some of its bioactive fractions in promoting wound healing and as anti-scar agent.

Fig. 1: Iraqi *Echinops heterophyllus*

Plant materials

The whole plant of *Echinops heterophyllus* of the Family (Compositae) was collected from Nazali, 7Km north of Erbil. The plant was authenticated by Dr. Abdul-hussein alkhait (an Iraqi plant classification specialist). The plant were cleaned and dried at room temperature in the shade and then pulverized by mechanical mills and weighed.

Preliminary qualitative phytochemical analysis

Chemical tests were carried out using the ethanolic extracts from plants and or the powdered specimens, using standard procedures to identify the active constituents[13-15].

Test for alkaloids

Ten (10ml) of alcoholic extract was stirred with 5 ml of 1% HCL on a stream bath. Mayer’s (1.53gm mercuric chloride in 60ml water + 5gm potassium iodide in 10ml water) and Wagner’s reagents (1.27g of iodine and 2g of potassium iodide in 100ml of water) were added, white and reddish brown color precipitate respectively, were taken as evidence for the presence of alkaloids.
Test for flavonoids
Lead acetate test: 5ml of alcoholic extract, 3ml of 10% lead acetate solution was added. The formation of a yellowish-white precipitate was taken as a positive test for flavonoids.

Tests for steroids
Liebermann-Burchard test extract (3ml) was treated with chloroform, acetic anhydride and drops of sulphuric acid was added. The formation of dark pink or red color indicates the presence of steroids.

Test for tannins
About 10mg plant material in 10ml distilled water was filtered, then 3ml of the filtrate + 3ml of FeCl₃ solution (5% w/v) were mixed. The formation of a dark green or blue black precipitate was considered an indication for the presence of tannins.

Test of saponins
Five ml of aqueous extract was shaken vigorously with 5ml of distilled water in a test tube and warmed. The formation of stable foam, honey comb in shapes, was taken as an indication for presence of saponins.

Tests for anthraquinoids
Borntreger's test: 3ml of alcoholic extract was shaken with 3ml of benzene, filtered and 5ml of 10% ammonia solution was added to the filtrate. The mixture was shaken and the development of a pink, red or violet color in the ammoniac (lower) phase indicates the presence of free anthraquinoid.

Test for terpenoids
Two ml of the organic extract was dissolved in 2ml of chloroform and evaporated to dryness. 2ml of concentrated sulphuric acid was then added and heated for about 2 min. A grayish color was considered an indication for the presence of terpenoids.

Extraction and fractionation of different active constituents
A (200gm) of shade-dried coarsely powdered plant materials separately were defatted with hexane for 24 hours then allowed to dry at room temperature. The defatted plant materials was extracted with (2 L) of 80% ethanol in soxhlet apparatus until complete exhaustion. The alcoholic extract was evaporated to dryness, under reduced pressure at a temperature not exceeding 40°C. To give a dark greenish-yellow residue designated as a crude fraction. Crude fraction was acidified with 300ml of 15% hydrochloric acid to pH 2 and partitioned (three times) with equal volume of ethyl acetate to get two layers (aqueous acidic and ethyl acetate layer). The aqueous acidic layer was then separated and basify with 5% sodium hydroxide to pH 10 and extracted with chloroform in the separatory funnel to get two layers, the aqueous basic layer was then separated and evaporated under reduced pressure at a temperature not exceeding 40°C. To give blackish residue designated as fraction 1 (F-1 which contains alkaloids) and aqueous basic layer. The ethyl acetate layer of the original alcoholic extract (crude fraction) was evaporated to dryness under reduced pressure and basify with 300ml of 5% sodium hydroxide to pH 10 and extracted with ethyl acetate in the separatory funnel to get two layers, the aqueous basic layer which was separated, evaporated to dryness and acidify with 5% HCL to pH 2 then extracted with ethyl acetate to get fraction designated as fraction 2 (F-2 which contained flavonoids)[14].

A relative assess on wound healing activity of crude *Echinops heterophyllus* extract and some of its bioactive fractions (F-1 and 2) was done as follow

In vivo experiment
Plant material
Crude plant extract, bioactive fractions (Alkaloids and flavonoids).

**Experiment Animals**
Twenty four adult male rabbits were used. Aged between six months to one year, obtained from the local market and placed in sterilized cages subjected to constant environmental conditions.

**Induction of wounds**
Surgical preparations were made at the upper back region after clipping, shaving and washing the area with tap water and drying. Then, standard longitudinal incisions (1x2 cm²) were implemented using a surgical scalpel[16,17].

**Fig. 2:** (1x2 cm²) Wound incision at the back region.

**Blood collecting**
Blood samples were obtained from each animal at day one, four, six and twelve. Measuring the blood glucose, protein, albumin, GOT and GPT, to ensure that animals are in a healthy state and the wound healing process was not affected by other factors. Two ml of blood was obtained through percutaneous cardiocentesis in anesthetized rabbits approaching the heart from the lateral left side and the midline under the sternum side aiming the needle toward the heart. Using 19 to 25G needle with 3 to 5 ml syringe and blood sample collection tubes with an anticoagulant agent[18-20].

**Animal groups**
Twenty four adult male rabbits were used and divided into four equal groups. The effect of crude *Echinops* extract and its bioactive fractions (Alkaloids and flavonoids) was evaluated visually and through histopathological changes. Treatment was applied three times daily using a cotton swab.

First group: was considered as a control group, each rabbit was wounded and left without treatment.

Second group: was considered as a treatment group, rabbits were wounded and treated with crude *Echinops* extract.

Third group: was also considered as a treatment group, wounded rabbits were treated with alkaloid fraction obtained from *Echinops* extract.

Fourth group: was considered as a treatment group, wounded rabbits were treated with flavonoids fraction obtained from *Echinops* extract.

**Histological Evaluation**
Specimens were taken from day one, four, sixth and twelfth. Animals were anesthetized using (Xylazine and Ketamine) in a dose of 5mg/Kg and 15 mg/Kg respectively[21]. Later, the specimens were kept in buffered formalin (10%) solution and examined.

**RESULTS**

**Preliminary qualitative phytochemical analysis**
The results of phytochemical screening are given in table-1.
Abdulrasool et al.


Table 1: Phytochemical Screening of Echinops heterophyllus

<table>
<thead>
<tr>
<th></th>
<th>Alkaloids</th>
<th>Flavonoids</th>
<th>Steroids</th>
<th>Tannins</th>
<th>Saponins</th>
<th>Anthraquinoin</th>
<th>Terpenoids</th>
</tr>
</thead>
<tbody>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
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<td>-</td>
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<td>+</td>
</tr>
</tbody>
</table>

+, - represent presence and absence of phytoconstituents respectively.

Phytochemical screening of Iraqi Echinops plant indicated that the plant contains different chemical classes of active constituents, alkaloids (basic compounds), flavonoids (acidic compounds) and steroids (neutral compounds) so the fractionation based on the conversion of basic compound to its salt by aqueous mineral acids, and when the salt of an alkaloid is treated with hydroxide ion, nitrogen gives up a hydrogen ion and the free amine is liberated which is taken or extracted by specific organic solvent like (chloroform) to get free alkaloids (F-1) leaving quaternary alkaloids and water soluble compounds in the aqueous layer. The same principle is applied to the acidic compounds to get flavonoids as a free aglycon in fraction-2 leaving neutral components in the organic layer.

**Blood test**

Results demonstrated normal levels for glucose, protein, albumin, GOT and GPT as in tables 2, 3, 4 and 5.

Table 2: Blood Samples from Day One.

<table>
<thead>
<tr>
<th></th>
<th>Glucose mg/dl</th>
<th>Protein gm.</th>
<th>Albumin g/dL</th>
<th>GOT U/liter</th>
<th>GPT U/liter</th>
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</thead>
<tbody>
<tr>
<td>Group 1</td>
<td>100</td>
<td>8.0</td>
<td>35</td>
<td>20</td>
<td>40</td>
</tr>
<tr>
<td>Group 2</td>
<td>185</td>
<td>7.5</td>
<td>4.5</td>
<td>30</td>
<td>45</td>
</tr>
<tr>
<td>Group 3</td>
<td>120</td>
<td>8.0</td>
<td>4.0</td>
<td>32</td>
<td>53</td>
</tr>
<tr>
<td>Group 4</td>
<td>275</td>
<td>8.0</td>
<td>4.0</td>
<td>44</td>
<td>62</td>
</tr>
</tbody>
</table>

Table 3: Blood Samples from Day Four

<table>
<thead>
<tr>
<th></th>
<th>Glucose mg/dl</th>
<th>Protein gm.</th>
<th>Albumin g/dL</th>
<th>GOT U/liter</th>
<th>GPT U/liter</th>
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</thead>
<tbody>
<tr>
<td>Group 1</td>
<td>95</td>
<td>8.0</td>
<td>3.5</td>
<td>28</td>
<td>39</td>
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<tr>
<td>Group 2</td>
<td>156</td>
<td>7.2</td>
<td>4.6</td>
<td>34</td>
<td>46</td>
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<tr>
<td>Group 3</td>
<td>170</td>
<td>6.3</td>
<td>4.2</td>
<td>43</td>
<td>67</td>
</tr>
<tr>
<td>Group 4</td>
<td>185</td>
<td>8.2</td>
<td>3.8</td>
<td>22</td>
<td>64</td>
</tr>
</tbody>
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Table 4: Blood Samples from Day Six

<table>
<thead>
<tr>
<th></th>
<th>Glucose mg/dl</th>
<th>Protein gm.</th>
<th>Albumin g/dL</th>
<th>GOT U/liter</th>
<th>GPT U/liter</th>
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<tr>
<td>Group 1</td>
<td>97</td>
<td>5.6</td>
<td>4.4</td>
<td>31</td>
<td>39</td>
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<tr>
<td>Group 2</td>
<td>160</td>
<td>7.4</td>
<td>5.0</td>
<td>26</td>
<td>33</td>
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<tr>
<td>Group 3</td>
<td>146</td>
<td>7.2</td>
<td>7.0</td>
<td>37</td>
<td>45</td>
</tr>
<tr>
<td>Group 4</td>
<td>158</td>
<td>8.3</td>
<td>4.5</td>
<td>29</td>
<td>58</td>
</tr>
</tbody>
</table>

Table 5: Blood Samples from Day Twelve

<table>
<thead>
<tr>
<th></th>
<th>Glucose mg/dl</th>
<th>Protein gm.</th>
<th>Albumin g/dL</th>
<th>GOT U/liter</th>
<th>GPT U/liter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1</td>
<td>206</td>
<td>5.5</td>
<td>3.6</td>
<td>24</td>
<td>47</td>
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<tr>
<td>Group 2</td>
<td>189</td>
<td>5.4</td>
<td>4.4</td>
<td>36</td>
<td>44</td>
</tr>
<tr>
<td>Group 3</td>
<td>123</td>
<td>6.3</td>
<td>5.9</td>
<td>33</td>
<td>46</td>
</tr>
<tr>
<td>Group 4</td>
<td>99</td>
<td>5.8</td>
<td>4.8</td>
<td>32</td>
<td>45</td>
</tr>
</tbody>
</table>

**Visual remarks**

**Group one (wounding without treatment)**

Normal healing took place which involves continuous cell–cell and cell–matrix interactions that allow the process to proceed in three overlapping phases: inflammation (0–3 days), cellular proliferation (3–12 days) and remodeling (3–6 months), so in this group there is some inflammatory signs seen from the first day with partial wound closure starting from the 4th day and some scar tissue at the 12th day. Figures (3-5).

Fig. 3: Group-1 day-1 Fig. 4: Group-1 day-6 Fig. 5: Group-1 day-12
Group two (wounding with treatment with crude Echinops extract): healing signs were very clear starting from day one. The complete fading of any inflammatory signs at day four and six. Finally, a complete wound closure at day twelve. Figures (6-8).

Fig. 6: Group-2 day-1  Fig. 7: Group-2 day-6

Fig. 8: Group-2 day-12

Group three (wounding with treatment with alkaloid fraction): remarkable wound healing signs from day one as there was no inflammatory signs. At day six wound edges started to convene. Lastly, an absolute wound closing at day twelve. Figures (9-11).

Fig. 9: Group-3 day-1  Fig. 10: Group-3 day-6  Fig. 11: Group-3 day-12

Group four (wounding and treatment with flavonoids fraction): mild inflammatory signs occurred at day one. A gradual healing appeared from day four till day twelve and few scar tissue at day 12th. Figures (12-14).

Fig. 12: Group-4 day-1  Fig. 13: Group-4 day-6  Fig. 14: Group-4 day-12

Histology
The control group was used as a model to be compared with the other three groups treated with the crude Echinops extract, alkaloid and flavonoids bioactive fractions obtained from Echinops heterophyllus extract. Hematoxylin and Eosin (H&E) Staining were used. Results demonstrated that the treatment group with the crude
Echinops extract gave the best results, while the alkaloid bioactive fraction was more potent than flavonoids bioactive fraction. The following figures demonstrates these results.

**Group one (wounding without treatment):** normal histological signs was observed.

![Image]

**Fig. 15:** Group one day one: a very clear appearance of inflammatory signs at the wound area during the first 24hrs, Fibrous threads took place and new epidermal layer formed the marginal ends started to thicken.

![Image]

**Fig. 16:** Group one day 12: There was no signs of inflammation. There was also an increase in collagen proliferation. Wound is not healed yet.

**Group two (wounding with treatment with crude Echinops extract)**

![Image]

**Fig. 17:** Group two-day one: (X 200) A selection of skin showing few amounts of inflammatory cells at the upper dermis layer and oedema, a clear migration of epidermal cells.
Fig. 18: Group two- day six: normal healing of dermis and epidermis layer.

Fig. 19: Group two- day 12: there was a full thickness epidermal regeneration which covered completely the wound area.

Group three (wounding with treatment with alkaloids fraction)

Fig. 20: Group three day one: some inflammatory cells at the upper dermis layer and oedema

Fig. 21: Group three day six: marked infiltration of the inflammatory cells, increased blood vessel formation and enhanced proliferation of cells as a result of treatment with alkaloids fraction
Fig. 22: Group 3 day 12: No inflammation, accumulation of granulation tissue, increase in the tensile strength. No scar formation.

Group four (wounding and treatment with flavonoids fraction)

Fig. 23: Group 4 day 1: a very clear appearance of inflammatory signs with oedema

Fig. 24: Group 4 day 6: almost complete healing

Fig. 25: Group 4 day 12: There was no signs of inflammation, accumulation of granulation tissue, increase in the tensile strength.
DISCUSSION

Wounds are referred to as disruption of normal anatomic structure and function. Skin wounds could happen through several causes like physical injuries resulting in opening and breaking of the skin. The most common symptoms of wounds are bleeding, loss of feeling or function below the wound site, heat and redness around the wound, painful or throbbing sensation, swelling of tissue in the area and pus like drainage. Wound healing is a very complex, multifactor sequence of events involving several cellular and biochemical processes. The aim in these processes is to regenerate and reconstruct the disrupted anatomical continuity and functional status of the skin. Healing process, a natural body reaction to injury, initiates immediately after wounding and occurs in four stages. The first phase is coagulation which controls excessive blood loss from the damaged vessels. The next stage of the healing process is inflammation and debridement of wound followed by re-epithelisation which includes proliferation, migration and differentiation of squamous epithelial cells of the epidermis. In the final stage of the healing process collagen deposition and remodeling occurs within the dermis. The results showed wound healing and repair, accelerated by applying crude extract of Echinops plant which was highlighted by the full thickness coverage of the wound area by an organized epidermis. The enhanced capacity of wound healing with the plant could be explained on the basis of anti-inflammatory effects of the active constituents of the plant (quinoline alkaloids, flavonoids, steroids and terpenoids).

Study on animal models showed enhanced rate of wound contraction and drastic reduction in healing time than control, which might be due to enhanced epithelisation. The animals treated with plant extract showed significant results when compared with untreated groups. The treated wound after six days itself exhibit marked dryness of wound margins with tissue regeneration.

REFERENCES