PHYTOCHEMICAL AND BIOLOGICAL STUDY OF CENTAUREA CINERARIA L. CULTIVATED IN SYRIA

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In the current study, C. cineraria plant was subjected to chemical and biological study. The plant extract was screened for its phytochemical constituents, its quantitative content of phenolics, and its activity in scavenging free radical (DPPH). The plant was subjected to a biological study, which includes investigating the potential cytotoxic effect of the plant towards human erythrocytes membranes, and the ability of the plant to protect erythrocytes against oxidative hemolysis. The results indicated that the extract did not contain saponins. Our study also showed that the extract contains a good content of phenolics (41.12±0.6 mg GAE/g, and has good DPPH radical scavenging activity (79.96%±1.14). The results showed that the extract does not have a toxic effect on erythrocytes membranes, and it is highly effective in protecting the membranes from oxidative damage (IC50 value was 408.86 μg/ml). We hope that future studies will focus on the pharmaceutical aspects of this plant, and will not be considered as ornamental plant only.

INTRODUCTION

Plants have been a great resource of medicinal products which are used for the treatment of several ailments. The Asteraceae is one of the largest families of flowering plants, with 1600–1700 genera and 24,000–30,000 species. Plants of this family are widely spread in the tropics and warm temperate regions of South, South-East and East-Asia, Africa and central South America. Centaurea L. is a large genus in the Asteraceae family, with over 500 species widespread across the world. Some species are cultivated for spice, as ornamental plants, and in gardens. Centaurea species produce a vast array of secondary metabolites belonging to different classes of bioactive compounds. The plant Centaurea cineraria is perennial herb. Stems up to 80 cm, erect, rarely procumbent, with few branches above. Leaves more or less tomentose, rarely glabrescent; lower lyrate to 2 pinnatisect. Capitula solitary. Bracts broadly ovate; appendages usually dark brown, the apex acuminate, not spinose; fimbriae 0.5–2 mm. Pappus as long as achene, rarely absent.

- Taxonomical Position of Centaurea species
  - Kingdom: Plantae
  - Phylum: Magnoliophyta (Flowering Plants)
  - Class: Magnoliopsida (Dicotyledons)
  - Subclass: Asteridae
  - Superorder: Asterales
  - Order: Asterales
  - Family: Asteraceae
  - Subfamily: Carduoideae
  - Tribe: Cynareae
  - Subtribe: Centaureinae
  - Genus: Centaurea
  - Species: C. cineraria

In general, C. cineraria is known as ornamental plant, and only very few studies have gone into investigating its chemical content and biological activities. The current study includes a chemical investigation of C. cineraria plant to screen its phytochemical compounds, determine its quantitative content of phenolic compounds, and test its activity in scavenging free radical (DPPH), in addition to a biological study, which includes investigating...
the potential cytotoxic effect of the plant towards human erythrocytes membranes, and the ability of the plant to protect healthy human erythrocytes against $H_2O_2$ induced hemolysis.

**METHODOLOGY**

**Plant material**

*C. cineraria* seeds were purchased, and then cultivated within strictly defined environmental conditions. For this study, the aerial parts of the cultivated plant were collected, shade dried and powdered using mechanical grinder.

**Preparation of plant extract**

Aqueous methanolic extract (AqME) of *C. cineraria* dried aerial parts was prepared according to Falleh et al. with few modifications.

The powdered plant material (50 gm) was mixed with 250 ml of aqueous methanol 70% in a bottle. The extraction procedure was carried out using ultrasonic bath with frequency of 15 kilohertz (kHz). The bottle was put in a sonic bath for 15 min. After that, the extract was vacuum filtered, the filtrate was collected and the residual was extracted three times using the same method. The total filtrate was evaporated by using rotary evaporator under reduced pressure at 45°C, and stored in a desiccator. Finally, the extraction yield was calculated by the formula:

$$
\text{Yield\%} = \left( \frac{\text{weight of dry extract}}{\text{weight of dry powdered plant material}} \right) \times 100
$$

**Qualitative analysis of phytochemical components**

*C. cineraria* aerial parts subjected to qualitative tests in order to characterize several chemical groups using standard procedures (Phenols, Flavonoids, Tannins, Carbohydrates, Saponins). The symbols + and - denote present, and absent.

**Total phenolic content**

Total phenolic content of the AqME extract of *C. cineraria* was spectrophotometrically determined using Folin-Ciocalteu method, with Gallic acid as standard (published in our previous study). The total phenolic content was reported as mg of gallic acid equivalent (GAE) per gram of dry extract.

**Free radical scavenging activity (DPPH method)**

The free radical scavenging activity of the AqME extract was determined according to Rashed, with modifications. First, DPPH solution was prepared in ethanol with final concentration of 45 µg/ml. Then, 1 ml of each plant extract (0.005 g/ml) was added to freshly prepared 3 ml of DPPH solution. After 30 min, the absorbance was measured at 517 nm. Control sample was prepared using distilled water instead of the plant extract. The percentage of inhibition was calculated by comparing the absorbance values of the control and test samples using the equation:

$$
\text{Free radical scavenging activity (\%)} = \left( \frac{(A_t - A_c)}{A_c} \right) \times 100
$$

$A_t$ is the absorbance of the control. $A_c$ is the absorbance of the test sample.

**Hemolysis assay**

*C. cineraria* AqME extract was tested for its cytotoxic effects on normal human erythrocytes. Five milliliters of blood from fifteen healthy donors - after their approval - was collected in EDTA vials. Normal human erythrocytes suspension (10%), and the concentrations of plant extract were prepared in phosphate buffered saline (PBS) solution, (pH 7.4). In sterile Eppendorf tubes, (1 ml) of each concentration (100-500-1000-2000-4000) µg/ml of the prepared extract was added to (1 ml) of erythrocytes suspension (isolated from the blood samples of ten healthy donors). The mixture was then incubated in a water bath at 37°C for 30 min, then samples were centrifuged (2500 rpm/min) for 10 mins, and the amount of free hemoglobin was estimated by measuring the absorption of the supernatant using a Spectrophotometer at the wavelength 540 nm.

In order to compare the results, a negative control sample containing (1 ml) of PBS and (1 ml) of erythrocytes suspension, and a positive control sample containing (1 ml) of distilled
water and (1 ml) of erythrocytes suspension were prepared17.

The percentage of hemolysis was calculated according to the following formula:

\[
\text{Hemolysis} \% = \frac{(\text{OD}_{\text{test}} - \text{OD}_{\text{con}})}{(\text{OD}_{\text{con}+} - \text{OD}_{\text{con}})} \times 100
\]

\( \text{OD}_{\text{test}} \): optical density of test sample.
\( \text{OD}_{\text{con}} \): optical density of negative control (PBS).
\( \text{OD}_{\text{con}+} \): optical density of positive control (distilled water).

Assessment of anti-hemolytic activity (hemolysis induced by oxidative factor)

The anti-hemolytic activity of \( C. \) cineraria AqME extract was determined by the spectrophotometric procedure. Five milliliters of blood from fifteen healthy donors - after their approval - was collected in EDTA vials. (1 ml) of each concentration of the extract (100-500-1000-2000-4000) µg/ml was mixed with (0.5 ml) of the erythrocytes suspension (10%) in sterile Eppendorf tubes. The mixture was incubated at laboratory temperature (5 min), then (0.5 ml) of hydrogen peroxide solution (prepared using PBS) was added to induce oxidative degradation of the membrane lipid (haemolysis). The mixture was incubated for 4 hrs at a temperature of 37°C. Then samples were centrifuged at a speed of 2500 rpm for 10 min.

In order to compare the results, a negative control sample containing (1 ml) of a PBS solution and (0.5 ml) of the healthy erythrocyte suspension, and (0.5 ml) of hydrogen peroxide solution was prepared.

Note: The hydrogen peroxide concentration was chosen to be able to cause hemolysis at approximately 95% in the negative control sample.

The percentage of hemolysis was calculated by estimating the amount of liberated hemoglobin using the spectrophotometer at the wavelength of 540 nm18. Based on this, the ability of plant extracts to protect erythrocyte from oxidative hemolysis was calculated according to the following equations:

\[
\text{Hemolysis} \% = \frac{\text{OD of sample}}{\text{OD of control}} \times 100 
\]

\[
\text{Protection} \% = 100 - \text{[hemolysis]} 
\]

Statistical analysis

Results were presented as average ± standard deviation using Microsoft Excel 2010. Statistical Package for Social Science (SPSS) was used to perform the statistical tests.

RESULTS AND DISCUSSION

Characteristics and yield of the plant extract

\( C. \) cineraria AqME extract was characterized by a brown color, bitter taste and a pleasant smell. The extraction yield was 12.64%.

Phytochemical screening

The results of the qualitative chemical analysis showed that the extract contain bioactive compounds as carbohydrates, phenols, tannins, flavonoids, while it was found that it does not contain saponins.

The phytochemical analysis of plants is important and have commercial attention in pharmaceuticals companies and research institutions for the industrialization of the new drugs19. The chemical compounds that have been found in \( C. \) cineraria AqME extract have important biological activities, which gives this species medical importance, in addition to its use as an ornamental plant in different countries of the world. No reference studies were found on the phytochemical analysis for this plant to discuss our results. Our study may be the first chemical and biological study of this plant.

Total phenolic content

Our results showed good phenolic content in \( C. \) cineraria AqME extract with (41.12±0.6) mg GAE/g.

It should be noted here that the phenolic content in the plant species in general is affected by the different environmental conditions for each geographical area, in addition to the different time of collecting the plant samples, and the difference of the solvent used in preparing the plant extracts20.

Free radical scavenging activity

Hydrogen donating ability of \( C. \) cineraria extract was measured by using DPPH assay. The results were expressed in percentage of inhibition of DPPH. The extract showed good
DPPH radical scavenging activity with (79.96%±1.14).

When testing the free radical scavenging activity of *C. cineraria* extract. It showed good DPPH radical scavenging activity. The ability to scavenge free radicals is determined by a 2,2-Diphenyl-1-picrylhydrazyl (DPPH) test, which is a free radical with a dark purple color in methanol and ethanol solutions. This test is based on an evaluation of the ability of plant extracts to reduce DPPH, which is inferred by the conversion of purple color to yellow.

**Hemolytic effect**

Hemolytic assay experiment was designed to evaluate the effect of five different concentrations of *C. cineraria* extract on red blood cells. The results showed that there were no toxic effects of the extract towards the erythrocyte membranes, although there was an increased direct relationship between the concentrations used from the plant extracts and the measured absorbance values. The tested concentrations caused a slight hemolysis (less than 5% at the concentration 4000 µg/ml) and this percentage is very weak when compared with the corresponding high concentration (table 1).

**Table 1:** The percentages of hemolytic effect by *C. cineraria* extract.

| Concentration (µg/ml) | Hemolysis %  |
|----------------------|-------------|
| 100                  | 0.26 ± 0.4  |
| 500                  | 0.49 ± 0.4  |
| 1000                 | 2.36 ± 0.6  |
| 2000                 | 3.72 ± 1.2  |
| 4000                 | 4.84 ± 0.9  |

All values are represented as mean ± SD. (*) Significant differences at (*p* < 0.05).

Before developing pharmaceutical agents from natural sources, it is important to screen them for cytotoxicity, and hemolytic activity represents a good indicator of cytotoxicity toward healthy erythrocytes. Triterpene and steroid saponins, which are abundant in plant extracts, have hemolytic properties.²¹ When conducting a qualitative phytochemical analysis of *C. cineraria*, it was found that the studied extract did not contain saponins, and the extract did not actually cause noticeable cytotoxic effects on erythrocytes’ membranes.

**Anti-hemolytic activity against oxidative hemolysis**

In this test, the extract’s ability to reduce the damage caused by oxidative stress in the erythrocyte membranes was evaluated. In general, the results indicated that the addition of the gradient concentrations of the *C. cineraria* extract resulted in the inhibition of oxidative hemolysis, which was stimulated *in vitro*, and in a relationship proportional to the gradual increase in the concentrations used, as shown in (table 2). A significant decrease in the percentages of erythrocyte hemolysis was demonstrated at high concentrations (1000-4000) µg/ml, which indicates the effectiveness of the extract in inhibiting the action of the oxidizing agent (H₂O₂). The IC₅₀ value was 408.86 µg/ml (Fig. 1). This result is consistent with the result of the free radical scavenging test, which confirms the plant’s activity in scavenging DPPH.

When comparing the results of the current study of the plant *C. cineraria* cultivated in Syria with a previous study¹⁴, that we conducted on the plant *C. iberica* wild spread in Syria, it became clear to us the great similarity in the chemical and biological effectiveness of the two species, and this may support the classification of these two species within the same genus.

**Table 2:** The percentages of hemolysis and protection by *C. cineraria* extract against oxidative hemolysis.

| Concentration (µg/ml) | Hemolysis % | Protection % | IC₅₀ (µg/ml) |
|----------------------|-------------|--------------|-------------|
| 100                  | 83.64         | 16.36±2.6    | 408.86      |
| 500                  | 49.91         | 50.09±1.2    |             |
| 1000                 | 18.44         | 81.56±3.1    |             |
| 2000                 | 12.96         | 87.04±1.4    |             |
| 4000                 | 5.54          | 94.46±1.5    |             |

Protection values are represented as mean± SD.
Conclusion
In the light of the results obtained, we concluded that the aerial parts of *C. cineraria*, are very rich in phytochemical compounds, and has very low toxicity against human erythrocytes. The plant also has an antioxidant activity that is demonstrated by the ability of the extract to scavenge the free radical (DPPH), and protect normal human erythrocytes from oxidative damage induced by H$_2$O$_2$.

Conflict of interest statement
No conflict of interest.

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Centaurea cineraria L. دراسة فيتوكيميائية وحيوية لنبات
المزروع في سوريا

ربى جوجه
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تم في هذا البحث إجراء دراسة كيميائية وحيوية لنبات Centaurea cineraria لإجراء الكشف الكيميائي للتحري عن المكونات الكيميائية الرئيسي في المستخلص النباتي، كما تم تحديد محتواه النوعي من المركبات الفينولية، واختبار فعاليته في كبح الجذور الحر. DPPH أجريت أيضًا دراسة حيوية تضمنت التحري عن السمية الخلية المحتملة لنبات تجاج أشبة الكريات الحمر البشرية، وكذلك اختبار قدرة النبات على حماية الكريات الحمر البشرية من الانحلال الدموي التأكسدي. أشارت نتائجنا إلى أن المستخلص المختبر لا يحتوي على مركبات الصابونيات، وتملك محتوى جيد من DPPH المركبات الفينولية، كما أبدى المستخلص النباتي فعالية جيدة في كبح الجذور الحر. DPPH أيضا أن المستخلص لا يبدى تأثيرا سمنا تجاج أشبة الكريات الحمر البشرية، بل يبدى فعالية عالية في حماية الأشبة من الأذى التأكسدي. نأمل أن توجه الدراسات المستقبلية للتركيز على أهمية هذا النبات في النواحي الصيدلانية، وعدم التعامل معه كنبات تزييني فقط.