EFFECT OF CELERY FLAVONOIDS ON LIVER ENZYME GOT AND GPT IN MICE

Aseel Shakir Mahmood and *Ibtisam Qahtan Abdul Kreem

Institute of Genetic Engineering and Biotechnology for Post Graduate Studies, University of Baghdad, Iraq

*Graduate Center \ Baghdad University

ABSTRACT

Celery is good source of flavonoid. They have also other positive effects on human health. The aim of current research was to study and compare an antioxidant activity. The hexanolic extracted and purification flavonoids from celery plant by using Thin layer chromatography (TLC). The study showed the effects of different three concentrations (10,20,30) of celery flavonoid extract on the effectiveness of liver enzymes (GOT,GPT) in blood and liver mice through injection Subcutaneous of laboratory mice during period of tow weeks in same age mice. The results of experiments demonstrate that the indicated ther were a significant differences (P<0.05) to concentration of the hexanolic extract on the effectiveness of tow types specific liver enzymes (GOT,GPT). The study showed of their inhibitory effects on certain enzymes and antioxidative activity. In mice treated with carbon tetrachloride CcL4, Through experiments show that the percentage of 30% is the best ratio of antioxidant.

INTRODUCTION

Celery (Apium graveolens L.) was introduced from Caucasia into China in about fifteen century. Ancient medicinal herbs records and modern research have proved that Celery has a series of medicinal properties, such as reducing blood press, sedation, promoting digestion, diuresis and moistening lung. Celery is, hepaxanthic herb grown as abieni and under certain conditions, as an annual. Celery is a native of Eurasia and is grown mainly in coastal regions. Celery is widely cultivated in the temperate Zones as an important garden crop and the bleached leaf stalks are relished as a popular vegetable. A. graveolens is one of ingredients in 8 of the 33 indian polyherbal formations with reuted life-protecting activity. Celery is also used as an effective remedy for various ailments such as bronchitis, liver and spleen disease, arthritic pain and this natural holistic approach to health is becoming mor and mor popular now adays (Kolarovic et al., 2010). (Jung 2011). Flavonoids (or bioflavonoids), collectively known as Vitamin p and citrin, are a class of plant secondary metabolites which are ubiquitous in photosynthesizing cell and are commonly found in fruits, vegetables, nuts, seeds, stems, flowers, tea, Wine, propolis and honey. For centuries, preparations containing these compounds as the principal physiologically active constituents have been used to treat human disease. (Mink et al., 2007). The function of flavonoids in flowers is to provide colors attractive to plant pollinators. In leaves, these compounds are increasingly believed to promote physiological survival of the plant, protecting it from, for example, fungal pathogens and UV-radiation, flavonoids are in photosensitization, energy transfer, the actions of plant growth hormones and growth regulators, control of respiration and photosynthesis, morphogenesis and determination. (Das et al., 2009). The basic structural feature of flavonoid compounds is the 2-phenyl-benzopyrene or flavane nucleus, which consiste of tow benzene rings (A and B) linked through a heterocyclic ring (C). Increasingly, this class of natural products is becoming the subjects of anti infective, and many groups have isolated and identified the structures of possessing antifungal, antiviral and antibacterial activity. (Shohaib et al., 2011).

The therapeutical effect of many traditional drugs are attributed to this group of compounds because of their inhibitory effects on certain enzymes and antioxidative activity. They have been shown to posses antibacterial, antifungal, antiviral and antinflammatory activities. Their anti allergic, antioxidative and antimutagen activities have been proven. Reduced risk of breast, prostate and colon cancers is related to isoflavonoid activity. Flavonoids have been studied in the prevention of menopausal symtoms and oseporosis. It was shown that their biological activity depended on the location of the free hydroxyl groups on ring A more so than that on ring B. (Dragan et al., 2007). Celery flavonoids will play very important role in theoretical and practical significance for Celery. The distribution of flavonoids from different Celery resources in Hunan was studied, and characteristics fingerprints of Apigenin, Apin from Celery with chromatographic technique. Biological activity and pharmacological functions of Celery flavonoids compounds on resistance liver cancer, leukemia, lowering lipid, anti-inflammatory were studied.

*Corresponding author: Ibtisam Qahtan Abdul Kreem, Institute of Genetic Engineering and Biotechnology for Post Graduate Studies, University of Baghdad, Iraq
(Amaowicz et al., 2004., Scott, 2012). The aim of search to studied flavonoid effecting on GPT, GOT blood and liver enzymes.

MATERIALS AND METHODS

The A. graveolens leaves were procured from local market in 2010 Baghdad Iraq. The leaves were washed thoroughly in tap water to remove adhering mud particules, rinsed in distilled water, drained, dried in a hot air oven at 50 tempereture. The dried leaves were finely powdered. The dried powder (50) gm was extracted with 500 ml hexanol for 10 h in soxhlate. The obtian extracts were filtered through filter paper. The crude extracted concentrated in rotary in tempereture 40 C°, So green material that give, and suspended in 250 ml Ethanol for 8 h. Solvents were removed in Royary and extracts were obtained, respectively. The residues were dissolved in 25 ml NaoH 5% in separation funal, then added 25 ml chloroform for 15 min and show tow layer the upper layer (water layer), lowring layer (chloroform layer). The upper layer had taken and Measured in PH 7 by used HCL, Then added 25 ml Chloroform. Total flavonoid content of celery leaves were determined by using way (Bajis et al., 1999), By mixed 10 ml from extracted with 50 ml Ethyl alcohol 95% in 1:1 vol:vol. TLC sorbents and three mobile phase were used for the analysis of the flavonoid exudates. Toluin : Diethyl ethir : acitic acid 10% (50:50:50) V \ V \ V, was used for the development of the exudates on silica gel plates (20x20 cm (0.2) mm layer. Detection with U.V light at (280) nm. Flavonoid reaction products were identified by RF values and chromatography with authentic substances on TLC in different solvent systems.

Lab animal

20 male mice were tested, with two months age and 21 gm weight, for 5 groups evry one contain 4 mice in Cages. So these mice that were nutritions on concentrated food and water, then several concentration from flavonoid extracted (10,20,30) mg/Kg (0.1) ml had injected dialy subcotenuos s/c for 14 days, these which laboratory animals were autopsied To take the blood and liver to hold the rest of the experiments.

Prepare of blood solution

Subjected mice under anesthesia, then later took heart blood and put in test tubs that contain EDTA. These solution was mixed with phosphate buffer, centerfigation 2000 cycle/min for 10 min. Supernated was took and 1 ml distal water added for it. GPT, GOT in blood extract were analaysed with least significant difference in p< 0.05.

Liver preparation

Liver mice had tooked, then cutting for small pices and mixed with 3 ml phosphate buffer, then centerfigation, supernated liver result have determination been the GOT,GPT activity in spectrophotometer.

RESULTS AND DISCUSSION

The accumulation prossecing consider first important steps, Dring plante the second step to lowring moisting of plante, to prevent enzymatic and microbial activity may be happen in plant tissue (Jung et al., 2011). The isolation and purification of flavonoids from the celery crud extract was succefully established by using Hexanol to remove volitical oils and fatties were found in celery plant without effecting on flavonoid, so another step using ethyl alcohol as phase solvent, following separation prossec NaoH sodium hydroxide 5% due to it solve in water layer, and added equal volium from chloroform to remove larg amount from chlorofil material (kavaratskhella 2004). (Das et al., 2009). Whene show yellowesh precepetae after added 10 ml extracted crude with 50 ml ethyle alcohol 95%. These lead to flavonoides found, these results successful with (Al Zubaidy, 2008).

The isolation and purification of flavonoids from the celery crud extract was succefully established by using toluin, diethyl ether, acetic acid 10% (50:50:50) as the tow- phase solvent. Celery flavonoids glycosides monomer was tracked in 280 nm wavelength by TLC thin layer chromatography, Celery flavonoids can very good separation. Flavonoid reaction products were Identified by RF values (0.68) and cochromotography with authentic substances on TLC in different solvent systems (Figure 1). Researchers are under way to study the effect of flavonoids in GOT and GPT enzymes in blood and liver in mouse. Affecting GPT, GOT enzyme level GPT enzyme found alarge ratio in liver more than another enzymes so that indicator for liver cell dameg, Celery flavonoid play a major role in decreased these enzymes ratio whene high resonance. (Robert et al., 2001) (Jenifer et al., 2009).

Figure 1. Purification flavonoid on TLC interaction with other enzyme system. Compared with research on the antioxident capacities of flavonoid, there has been relatively little research on other beneficial effects of flavonoids. The major effects of flavonoid may be the result of radical scavenging. Another possible mechanism by which flavonoids act is through interaction with various enzyme systems. Another interesting effect of flavonoid on enzymes system is the inhibition of arachidonic acid (Hanneken et al., 2006). Show in Table 1 enzymes GPT, GOT for two weeks report decreasing (P < 0.05) after added the material,
comparative with positive controlling during injection Subcutaneous period of different extract concentration. In the experiments periods, lead to that these materials have been effective in liver enzymes these results Conformable with (Jung et al., 2011), compounds because of their inhibitory effects on certain enzymes and antioxidative activity. (Dragant 2007) (Alkubasy et al., 2002).

Table 1. Affected different extracted concentration in GPT, GOT enzymes ratio in blood lab.animal for two weeks (P < 0.05)

<table>
<thead>
<tr>
<th>Effecting concentration</th>
<th>Quality activation (unit/L) GPT Blood</th>
<th>Quality activation (unit/L) GOT Blood</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positve controlling</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distal water</td>
<td>79.7±0.24</td>
<td>54.25±5.50</td>
</tr>
<tr>
<td>Negative controlling (Ccl4)</td>
<td>128.76±4.57</td>
<td>167.67±1.73</td>
</tr>
<tr>
<td>10% flavonoid</td>
<td>104.05±0.12</td>
<td>146.74±2.04</td>
</tr>
<tr>
<td>20% flavonoid</td>
<td>97.68±4.25</td>
<td>113.023±0.14</td>
</tr>
<tr>
<td>flavonoid 30%</td>
<td>90.35±0.76</td>
<td>79.11±0.40</td>
</tr>
</tbody>
</table>

Also when GPT, GOT enzyme measure in lab. Animal liver flavonoid was inhibitory effects on certain enzymes after give lab. Animal Ccl4, So table (2) show 30% the best ratio near from normal, these equal with (Robert et al., 2001), (Pansiri, 2004).

REFERENCES

Al subaide Ali Hafed 2007. Studied of Mutagenic aptility of water extractions and alcoholic for Maramia plant Salvia officinalis by used bacterial system. Baghdad University /instituted Genetic Engineering


**********