Benefiting of Some Plant Materials in Improving Liver Functions for Hepatically Toxicated Rats with CCl₄

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Abstract

The present study investigated the effects of carrot (Daucus carota L.), Prickly pear (Opuntia ficus-indica L.) and Chicory (Cichorium intybus L.) on CCl₄ induced hepatotoxicity in rats . Forty five male albino rats were divided into (9) groups (5) rats in each group. Two groups were controls, one fed on basal diet only as a negative control and the other one fed on basal diet after injection with CCI₄ as a positive control. The other groups were injected by CCl₄ then received basal diet containing carrot, chicory and prickly pear at the levels 10, 15% and 15% mixture of the tested plants. Liver damage was assessed by estimation of plasma concentration of enzymes activities of aspartate amino transferase (AST), alanine amino transferase (ALT), lipid fraction (total cholesterol and triglyceride), cholesterol fraction (HDL-c, LDL-c, VLDL-c), Uric acid, Urea nitrogen and glucose. Results showed an improvement in case of prickly pear followed by carrot and chicory for the above parameters. The best level was 15% of these plants followed by 10% of tested plants. So, this study concluded that CCl₄ induced liver damage in rats can be ameliorated by administration of 15% prickly pear, carrot and chicory. **Key words:** Carrot - prickly pear - chicory- liver damage - Cholesterol fraction-glucose.

Introduction

Liver diseases are among the most serious aliment. They may be classified as acute or chronic hepatitis (inflammatory liver diseases), hepatosis (non inflammatory diseases) and cirrhosis (degenerative disorder resulting in fibrosis of the liver) (Kumar *et al.*, 2011).

Carbon tetrachloride (CCl₄) has been widely used in animal models to investigate chemical toxin-induced liver damage. The most remarkable pathological characteristics of CCl₄ induced hepatotoxicity are fatty liver, cirrhosis and necrosis (Recknagel et al., 1989). CCl₄ is a well-known hepatotoxic agent (Ilavarasan et al., 2003). A single dose of CCl₄ (20 micro 1/kg) induced hepatotoxicity, manifested biochemically by significant elevation of serum enzymes activities, such as alanine trasaminase (ALT), aspartate aminotransferase (AST), and lactate dehydrogenase (LDH) (Mansour, 2000).

Phytotherapy is the treatment and prevention of diseases using plants or plants part, such as leaves, flowers, roots, fruits, seeds, and rhizomes. Preparation made from them called medicinal plants, or herbs (Weiss and Fintelmann, 2000). Many plants were suggested to ameliorate or care the liver diseases, among them were the birch, celandine, chicory, dandelion, fennel, gentian, avocado 'rosemary, papaya, onion, carrot and lettuce (Morsi, 1992). Medicinal plants have very important place as they not only maintain the health and vitality of human beings and animals, but also cure several disease, including liver disorders without causing any toxicity (Govind and Madhuri, 2010).

Daucus carota linn. (Family :umberliferae) is an annual or biennial herb, whose roots are eaten raw and also cooked in many parts of the world (Muralidharan et al., 2008). Among common fruits and vegetables, carrots are high in fibers, carotenoids, vitamins C and E and phenolic such as P-coumaric, chlorogenic, and caffeic acids (Potter et al., 2011). Carrot polyacetylense possess allelopathic activity which may explain the historical health benefits of carrots since studies investigating β - carotene doesn't seem to adequately explain the reduced risk of certain types of cancer (Brandon and David, 2012).

Carrot could afford a significant protective action in the alleviation of CCl₄ induced hepatocellular injury (Bishayee *et al.*, 1995). Oral administration of carrot extract produced significant hepatoprotection against lindane induced hepatotoxicity in rats. The increase levels of serum enzymes namely aspartate transaminase, alanine transaminase, alkaline phosphatase and the levels of thiobrabituric acid reactive substances, cholesterol, triglycerides and LDL – cholesterol in ilndane administered rates were observed to be decreased significantly in the lindane (+) carrot extract group. The carrot extract also restored the depressed antioxidants and HDL- cholesterol levels to near normal (Balasubramaniam *et al.*, 1998).

Cactus pear or prickly pear, a member of the cactacea family, originated from arid and semi- arid regions of Mexico. More than 1500

known species of cactus are in the genus opuntia. Cactus pear fruit containing betalain pigments is a good potential for the use as a natural food colorant. This fruit contains the red-violet betacyanins in addition to the yellow betaxanthins (Salim et al., 2009). The prickly pears are considered to be a good source of minerals and other nutrients on the basis of compositional analysis. Availability of K, Ca and Mg in prickly pear was assessed using an in vitro digestion and dialysis method (MC Conn and Nakata, 2004). The juice of prickly pear in nutritionally interesting and its dietary intake could provide protection against oxidative damage (Galati et al., 2003). The cactus pear fruit extract were analyzed for determined constituents: ascorbic acid, flavonoids (quercetin, isorhamnetin, myricetin, kaempferol and luteolin), betalains, taurine, total carotenoids and total phenolics. Opuntia ficus indica fruit extract had strong antioxidant capacity and taurine content (Frenandezlopez et al., 2010).

Opuntia ficus indica fruit juice administration exerts protective and curative effects against the CCl₄- induced degenerative process in rat liver (Galati et al., 2005). The pears of Opuntia have been discovered to contain a plethora of biologically active compounds. Owing to their high nutritional value, in terms of dietary fiber, polyunsaturated fatty acidrich oil, minerals, protein, and an assortment of other phytochemicals, the pears are gaining popularity as exotic, gourmet diet (Patel, 2013).

Chicory (Cichorium intybus L.) is a perennial herbs in the Asteraceae family with many commercial uses. Historically, chicory was grown by the ancient Egyptians as a medicinal plant, vegetable crop, and for animal forage (Judzntiene and Budiene, 2008). Inulin from chicory roots is considered a functional food ingredient as it affects physiological and biochemical processes resulting in better health and reduction of the risk of many diseases (Kaur and Gupta, 2002). The aqueous- alcoholic extracts of aerial parts of C. intybus inhibited xanthine oxidase enzyme dose dependently (Pieroni et al., 2002).

Esculetin, aphenolic compound found in *Cichorium intybus* investigated the possible protective effect against poracetamol and CCl₄-induced hepatic damage (Gilani *et al.*, 1998). Cichorium root extract therapy led to normalization of some morphofunctional liver features (decreases glycogen content and cell of necrosis and increases the number of cells with pronounced protein synthesis activity) in rats with CCl₄ -induced hepatitis (Krylova *et al.*, 2006). The present study was carried out to investigate biological effects of carrot, chicory prickly pear and mixture of the tested plants on serum parameters of liver intoxication in rats.

.Material and Methods Materials Plants

The tested plants were chicory (Cichorium intybus L.), carrot (Daucus Carota L.) and prickly pear (Oputia Ficus indica L.). Chicory

was purchased from herbalist of Cairo, Egypt. Carrot and Prickly pear were purchased from the local markets of Shebin El-kom, Menofia Governorate, Egypt.

Carbon tetra choloride (CCl₄) was used as an inducer for liver cirrhosis. It was purchased from El-Gomhorya Company, Cairo, Egypt as 10% liquid solution.

Chemical reagents

Reagent kits were purchased from Diamond Diagnostics (Egypt).

Experimental animals

Forty five white male albino rats weighing about $200 \pm 5g$ were used as experimental animals in the present investigation. They were obtained from the animal house of Research Institute of Ophthalmology, El-Giza, Egypt. They were kept under observation for one week (as adapted period) before the onset of the experiment. The animals were housed in stainless steel cages at normal atmospheric temperature (25 \pm 5°C) and had a 12 h light-dark cycle. Food and water were consumed ad libitum.

Methods:

Induction of liver intoxication in rats

Forty rats were treated subcutaneous injection of carbon tetra chloride in paraffin oil 50% V/V (2ml/kg b.wt) twice a week for two weeks (Jayaserkar *et al.*, 1997).

Preparation of plant powder

These plants were washed and dried in drying oven at 50°C for 3 days, then crushed and milled as a dried powder.

Animals diet

The basal diet was prepared according to AIN (1993). The vitamin mixture was prepared according to Campbell (1963), while salt mixture was prepared according to Hegsted et al. (1941).

Experimental design

Forty five male albino rats $(200 \pm 5g)$ were randomly divided into 9 equal groups (five rats each). All rats were fed on basal diet for one week before starting the experiment for acclimatization. After the adapted period, the initial weight was $205 \pm 5g$. Groups of rats were as the follows:

Group (1): Rats (n=5) were fed on basal diet only as control negative group.

Group (2):Rats (n=5) were kept without any treatment as positive control group and fed on basal diet after injection with CCl4.

Group (3): Rats (n=5) were injected by CCl4 then fed on basal diet containing 10% chicory.

Group (4): Rats (n=5) were injected by CCl4 then fed on basal diet containing 15% chicory.

Group (5): Rats (n=5) were injected by CCl4 then fed on basal diet containing 10% carrot.

Group (6): Rats (n=5) were injected by CCl4 then fed on basal diet containing 15% carrot.

Group (7): Rats (n=5) were injected by CCl4 then fed on basal diet containing 10% prickly pear.

Group (8): Rats (n=5) were injected by CCl4 then fed on basal diet containing 15% prickly pear.

Group (9): Rats (n=5) were injected by CCl4 then fed on basal diet containing 15 % of mixture chicory, carrot and prickly pear(1:1:1).

By the end of the experimental periods (35 days), rats were scarified using diethyl ether anesthesia at fasting state. Part of the blood was taken to determine the level of serum glucose and other portion of blood samples was collected and allowed to coagulate at room temperature; other portion of blood was added to it, EDTA (ethylene diamine tetracetic acid) and centrifuged at 3000 r.p.m for 15 minutes. Serum was carefully aspirated and transferred into clean covet tubes and stored frozen at -20°C until the time of analysis.

Biochemical analysis:

Serum Alkaline phosphatase (ALP) was determined according to the procedure of (IFCC methods., 1983). Aspartate aminotransferase (AST) or (GOT) glutamic -oxaloacetic transaminase and glutamic pyruvic transaminase (GPT) or Alanine aminotransferase (ALT) were carried out according to the method of Henry (1974) and Yound (1975). Serum uric acid was determined according to the method described by Fossati et al. (1980). Serum urea in plasma was determined according to the enzymatic method of Patton and Crouch (1977). Glucose was determined by enzymatic test according to Tietz (1976) and Yound Enzymatic colorimetric determination of triglycerides was (1975).carried out according to Fassati and Prencipe (1982). Total Cholesterol was determined by colorimetric method according to Allain (1974). The determination of HDL was carried out according to the method of Fnedewaid (1972) and Gordon and Amer (1977). The determination of VLDL (very low density lipoproteins) and LDL (low density lipoproteins) was carried out according to the method of Lee and Nieman (1996).

Statistical analysis

Statistical analysis were done using the Statistical Package for the Social Sciences (SPSS for WINDOWS, version 11.0; SPSS Inc, Chicago). Comparative analyses were conducted using the general linear models procedure (SPSS Inc). Values of P<0.05 were considered statistically significant.

RESULTS

1-Effect of feeding different levels of chicory, carrot and prickly pear on ALP, AST and ALT levels of CCl4-intoxicated rats.

The results in table (1) indicated that mean value of ALP enzyme, rats injected with CCl4 (C +ve group) was 230.7±3.47U\L while in normal rats (C -ve) was 99.3±1.32U\L. These results denote that there

was a significant increase in the mean value of ALP enzyme of rats poisoned by CCl4 as compared to normal rats. The mean values of (ALP) of diets from groups 3, 4, 5, 6, 7, 8 and 9 were significantly higher than control negative group. Also, it could be noticed that there is no significant differences between the values of ALP enzyme of groups 4, 5 and 7. Meanwhile, rats given CCl4 then fed on diet of group 6 (rats fed on basal diet with 15% carrot) showed the lowest mean value in ALP enzyme level in the serum which was 199.4±1.35 U\L as compared to control positive group and recorded the best result of all treatment. It could be observed that due to intoxicated rats the serum levels of AST in table (1) showed a significant increase in control positive group as compared to normal rats represents 37.25±5.82 and 22.06±1.07 U/L, respectively. There is no significant differences between groups 3, 7 and control positive group. Also, there is no significant differences between groups 3 and 9. Meanwhile, group 6 (rats fed on basal diet with 15% carrot) showed the lowest level in serum AST and recorded the best results as compared to all treatments. For ALT, in rats given CCl4 then fed on all treatments groups 3, 4, 5, 6, 7, 8, and 9 were showing a significant differences when compared to control negative group. There is no significant difference between groups 3, 4 and 8. The obtained results showed that there is no significant difference in serum levels of ALT in group 6 as compared to normal rats and the best treatment was recorded for group 6 (rats fed on basal diet with 15% carrot).

Table (1): Effect of feeding different levels of chicory, carrot and prickly pear on ALP, AST and ALT levels of CCl₄-intoxicated rats.

Liver function	ALP Mean SD	AST Mean SD	ALT Mean SD
Animal Groups	Wicali SD	Wicali SD	Mean SD
Group (1) Control – ve	99.31.32 ^E	22.061.07 ^D	9.510.94 ^D
Group (2) Control + ve	230.73.47 ^A	37.255.82 ^A	16.101.10 ^A
Group (3) %10chicory	227.35.06 ^A	28.661.76 ^B	12.290.26 ^C
Group (4) %15chicory	222.43.13 ^B	24.870.42 ^C	11.792.28 ^C
Group (5) % 10carrot	222.37.5 ^B	37.527.22 ^A	15.931.25 ^A
Group (6) % 15carrot	199.41.35 ^D	22.311.80 ^D	10.440.79 ^D
Group (7) %10prickly pear	219.72.05 ^B	35.104.92 ^A	14.431.21 ^B
Group (8) %15prickly pear	206.31.35 ^C	26.120.51 ^C	12.800.48 ^C
Group (9) %15mixture of all plant	207.13.11 ^C	31.120.51 ^B	14.800.48 ^B

^{*}Non significant differences between the values had the same letter. Significant at $p \le 0.05$.

2- Effect of feeding different levels of chicory, carrot and prickly pear on total cholesterol and triglyceride levels (mg/dl) of CCl₄ intoxicated rats.

Data in Table (2)revealed that Injection of CCl₄ led to significantly (P≤0.05) increased serum total cholesterol level in hepatotoxic rats. The $(P \le 0.05)$ increased serum total cholesterol level in hepatotoxic rats. The mean value \pm SD of serum cholesterol in hepatotoxic group control (+ve) was 172.55 ± 12.38 mg/dl compared to 89.78 ± 5.25 mg/dl in the control (-ve) group. The mean values of total cholesterol in rats given CCl₄ then fed on all diets of groups 3 , 4 , 5 , 6, 7 , 8 and 9 were significantly lower than positive control group. There is no significant differences in total cholesterol between groups 3 and 7, as well as between groups 5, 6 and 8. Groups 6 and 8 showed the lowest levels in total cholesterol as compared to all groups. Concerning triglycerides, (Table 2) data revealed that rats injected with CCl₄ (control positive group) had higher value ($P \le 0.05$) of serum levels triglycerides compared to normal rats control negative group. There were nonsignificant differences between groups 3 and 7as well as between groups 5 and 9. Meanwhile, group 8 (rats fed on diet contained 15% prickly pear) showed the lowest level in the mean value of serum triglycerides which showed 54.10 ± 4.72 mg/dl as compared to all treatment and recorded the best result.

Table (2): Effect of feeding different levels of chicory, carrot and prickly pear on total cholesterol and triglyceride levels (mg/dl) of CCl4 intoxicated rats.

Lipid Fraction **Animal Groups** Total cholesterol Triglyceride Mean SD Mean SD Group (1) $39.400.96^{G}$ $89.785.25^{\text{F}}$ Control – ve Group (2) 172.5512.38^A 110.703.11^A Control + ve Group (3) 151.296.92^C 91.301.44^C %10chicory Group (4) $103.402.04^{\rm B}$ $165.781.72^{B}$ %15chicory Group (5) 124.124.71^E $70.003.82^{D}$ % 10carrot Group (6) $119.462.62^{E}$ $60.106.74^{E}$ % 15carrot Group (7) 154.316.15^C 96.002.85^C %10prickly pear Group (8) $117.489.24^{E}$ 54.104.72^F %15prickly pear Group (9) 137.081.04^D $74.100.92^{D}$ %15mixture of all plant

^{*}Non significant differences between the values had the same letter. Significant at p≤0.05.

3- Effect of feeding different levels of chicory, carrot and prickly pear on HDL-c, LDL-c, VLDL-c and the ratio between LDL-c/HDL-c levels (mg/dl) of CCl4-intoxicated rats.

It is obvious that in rats injected with CCl₄ (control+ ve) the mean value of serum levels HDL-c was 28.38±5.33 mg/dl. In normal rats (control-ve) the mean value of serum levels the HDL-c was 60.58±3..62 mg/dl in table (3). These finding denote that there was a significant decrease in HDL-c in the serum of rats poisoned by CCl₄ as compared to normal rats in table 3. There were non significant differences between rats given CCl₄ then fed on diet of groups 3, 5,6, 7 and 9. Finally group 8 (rats fed on diet contained 15% prickly pear) showed the highest increase in serum level of HDL-c and recorded the best treatments. It could be noticed that the data in table 3 evidence that, LDL-c levels was significantly elevated in control positive group to 104.03±8.07 from 20.86±2.74 mg/dl in control negative group. All rats intoxicated with CCl₄ then fed on all tested plant materials showed significant decrease in LDL-c as compared to control positive group. Group 8 (rats fed on diet contained 15% prickly pear) showed the lowest value of serum LDL-c and recorded the best results as compared to all treatments. Data presented in Table(3) indicated the effect of feeding CCl₄ intoxicated rats with chicory, carrot, and prickly pear on the serum levels of VLDLc. There were non-significant differences between normal group 7, 8, 9 and negative control group. Group 8 (rats fed on diet contained 15% prickly pear) showed the lowest decrease in serum level of VLDL-c and recorded the best results as compared to all groups in table 3. As regards to rats injected with CCl₄ without treatment (control positive), the serum LDL-c/HDL-c increase dramatically from 0.34 ± 0.04 for control negative group to 3.67 ± 1.03 for control positive group in Table (3). Rats fed on based diet contained 15% prickly pear showed the lowest level in the serum LDL-c/HDL-c and recorded the best results as compared to all treatments.

Table (3): Effect of feeding different levels of chicory, carrot and prickly pear on HDL-c, LDL-c, VLDL-c and the ratio between LDL-c/ HDL-c levels (mg/dl) of CCl₄-intoxicated rats.

Lipid fraction Animal Groups	HDL-C MeanSD	LDL-C Mean SD	VLDL-C MeanSD	LDL-C/HDL-C MeanSD
Group (1) Control – ve	60.583.62 ^A	20.862.74 H	7.920.19 ^E	0.340.04 ^G
Group (2) Control + ve	28.385.33 ^E	104.038.0 7 ^A	22.200.64 ^A	3.671.03 ^A
Group (3) %10chicory	50.054.08 ^C	62.978.48	11.420.94 ^D	1.260.22 ^c
Group (4) %15chicory	43.384.36 ^D	74.856.59	15.000.76 ^B	1.730.19 ^B
Group (5) % 10carrot	49.551.64 ^C	83.303.71	13.160.76 ^C	1.680.13 ^B
Group (6) % 15carrot	52.817.63 ^C	46.076.61	10.870.58 ^D	0.870.57 ^E
Group (7) %10prickly pear	52.034.16 ^C	57.888.58	8.860.28 ^E	1.110.26 ^D
Group (8) %15prickly pear	56.461.94 ^B	35.676.61	7.880.58 ^E	0.630.52 ^F
Group (9) %15mixture of all plant	52.461.04 ^C	45.670.61	8.080.38 ^E	0.860.12 ^E

*Non significant differences between the values had the same letter. Significant at p≤0.05

4- Effect of feeding different levels of chicory, carrot and prickly

4- Effect of feeding different levels of chicory, carrot and prickly pear on glucose levels (mg/dl) of CCl4 intoxicated rats.

It could be observed that, the mean value ±SD of glucose in Table (4) of control positive group significantly increase, as compared to normal rats, it was being 140.14±0.02 and 80.05±2.11 mg/dl, respectively. In rats given CCl4 then fed on all treatments, there were significant increases in the glucose levels as compared to normal group which were 117.14±0.59, 130.13±0.21, 130.12±0.85, 116.11±0.14, 129.11±0.20, 111, 9±0.12 and 125.79±0.72 mg/dl for groups 3, 4, 5, 6, 7, 8 and 9 respectively. There is no significant difference between groups 3,6 and 8. Also, there is no significant difference between groups 4, 5,7 and 9. Finally, group 8 (rats fed on diet contained 15% prickly pear) showed the lowest increase in glucose level which were 111.9±0.12 and recorded the best treatment.

Table (4): Effect of feeding different levels of chicory, carrot and prickly pear on glucose levels (mg/dl) of CCL intoxicated rats

Animal Groups	Glucose Mean SD	
Animai Groups	Glucose Weam SD	
Group (1)	80.052.11 ^D	
Control – ve	80.032.11	
Group (2)	140.140.02 ^A	
Control + ve	140.140.02	
Group (3)	117 140 50 ^C	
%10chicory	117.140.59 ^C	
Group (4)	120 120 21B	
%15chicory	130.130.21 ^B	
Group (5)	120 120 05B	
% 10carrot	130.120.85 ^B	
Group (6)	116 110 140	
% 15carrot	116.110.14 ^C	
Group (7)	120 110 20B	
%10prickly pear	129.110.20 ^B	
Group (8)	111.00.120	
%15prickly pear	111.90.12 ^C	
Group (9)	125 700 72B	
%15mixture of all plant	125.790.72 ^B	

^{**}Non significant differences between the values had the same letter. Significant at p≤0.05

5- Effect of feeding different levels of chicory, carrot and prickly Uric acid and Urea nitrogen levels (mg/dl) CCl4intoxicated rats.

Results revealed that, treated rats with CCl₄ -intoxicated diet control positive group led to a significant increase (P\le 0.05) in serum uric acid when compared with control negative group. The mean values of uric acid of groups 4, 5, 6, 7, 8 and 9 were significantly lower than positive control group (Table 5). Non-significant differences were observed between groups 3 and control positive group. Meanwhile, group 9 (rats fed on diet contained 15% mixture of all plant materials) showed the

lowest level in serum uric acid among all treatment and recorded the best results compared to normal group. For urea nitrogen, there is nonsignificant difference between group 3 and control positive group. Also, group 4 was similar to normal group. Groups 5, 6,7,8 and 9 showed lower (P<0.05) in urea nitrogen than both control groups. Finally, group 9 (rats fed on basal diet with 15% mixture of all plant materials) showed the lowest level of urea nitrogen among all treatment groups.

Table (5): Effect of feeding different levels of chicory, carrot and prickly pear on Uric acid and Urea nitrogen levels (mg/dl) of CCl₄ intoxicated rats.

Kidney function Animal Groups	Uric acid Mean SD	Urea Nitrogen Mean SD
Group (1) Control – ve	1.050.11 ^C	14.70.9 ^B
Group (2) Control + ve	1.140.12 ^A	15.62.2 ^A
Group (3) %10chicory	1.130.11 ^A	15.61.4 ^A
Group (4) %15chicory	1.140.09 ^A	14.90.6 ^B
Group (5) % 10carrot	1.120.05 ^B	13.61.9 ^C
Group (6) % 15carrot	1.110.15 ^B	14.10.8 ^C
Group (7) %10prickly pear	1.110.10 ^B	13.80.9 ^C
Group (8) %15prickly pear	0.90.72 ^D	13.11.2 ^D
Group (9) %15mixture of all plant	$0.790.72^{E}$	10.11.2 ^E

^{**}Non significant differences between the values had the same letter. Significant at p≤0.05

Discussion:

The reactive electrons species from CCl₄ induces rat liver cirrhosis that resembles the human disease, and it can serve as a suitable animal model for studying human liver cirrhosis (An et al., 2006). Toxicity experienced by the liver during CCl₄ poisoning results from the production of a metabolite, CCl₄ which is a direct hepatotoxin responsible for change in cell permeability and it inhibits mitochondrial activity followed by cell death (Ambrose et al., 2009). It has also been reported that chronic CCl₄ exposure produced cirrhosis in rats (Chieli and Malvadi, 2008). An obvious sign of hepatic injury is the leakage of cellular enzyme into plasma (Schmidt et al., 1975). When the liver cell plasma membrane is damaged, a variety of enzymes normally located in the cytosol are released into blood stream. Their estimation in the serum is a useful quantitative marker for the extent and type of hepatocellular damage (Ansari et al., 1991). ALT and AST are the most often used and

most specific indicators of hepatic injury and represent markers of hepatocellular necrosis. These liver enzymes catalyze transfer of alphaamino group aspartate and alanine to the alpha-ketoglutaric acid. Whereas ALT is primarily localized to the liver, AST is present in a wide variety of tissue, including heart, skeletal, kidney, brain, and liver. AST is present in both the mitochondria and cytosol of hepatocytes, but ALT is found only in the cytosol. In an asymptomatic person with isolated elevation of AST or ALT level, diagnostic clues can be garnered from the degree of elevation (Posen and Kaeffe, 1998). Possible of the from the degree of elevation (Rosen and Keeffe, 1998). Results of the current study revealed that administration of CCl₄ caused significant increases in the levels of aspartate aminotransferase, alanine aminotransferase, glucose levels, lipid profile and kidney enzymes and these are in agreement with **Túnez** et al. (2005). On the other hand, the current study demonstrated that the treatment with Cactus pear extract caused marked ameliorations of transaminase analysis. caused marked ameliorations of transaminase enzymes activity (ALT and AST). The results are in accordance with **Tapiero** et al. (2002) who showed the effect of carrot extracts on carbon tetrachloride—induced hepatotoxicity in rats. The mechanism by which the cactus pear fruit hepatotoxicity in rats. The mechanism by which the cactus pear fruit induces its hepatoprotective activity is not certain. However, it is possible that β-sitosterol, a constituent of cactus pear, which is at least partly responsible for the protective activity against CCl₄ hepatotoxicity (Tesoriere *et al.*, 2004). An additional and important factor in the hepatoprotective activity of any drug is the ability of its constituents to inhibit the aromatase activity of cytochrome P-450, thereby favoring liver regeneration. On that basis, it is suggested that flavonoids in cactus pear could be a factor contributing to its hepatoprotective ability through inhibition of cytochrome P-450 aromatase (Kowalska *et al.*, 1990). In addition, the recorded content of vitamin C in the chicory (35 -38 mg per 100 g) may also play a role in hepatoprotection. Previous in vivo studies indicate that hepatic microsomal drug metabolism decreases in ascorbic acid deficiency and is augmented when high supplements of the vitamin are given to guinea pigs (Burtis and Ashwood, 2001). Blood glucose concentration is known to depend on the ability of the liver to absorb or produce glucose. The liver performs its glucostatic function owing to its produce glucose. The liver performs its glucostatic function owing to its ability to synthesize or degrade glycogen according to the needs of the organism, as well as via gluconeogenesis (Ahmed et al., 2006). The blood sugar level after overnight fasting in cirrhotic patients is believed to decrease only in severe hepatic failure (Kruszynska and McIntyre, 1991). This is confirmed by our data that indicate that glucose levels in cirrhósis decreased.

Conclusions

The study clearly demonstrates that 15% of tested plants and its mixture have potential for treatment and prevention of CCl_4 -induced hepatic cytotoxicity. This study, along with other research, targets cactus pear as a potentially safe and effective fruit that has important medicinal

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الاستفادة من بعض المواد النباتية في تحسين وظائف الكبد لدى الفئران المصابة بالتسمم الكبدى برابع كلوريد الكربون أ.د/ يحيي عبدالمنعم عبدالهادي د/ نهاد طاحون د/ حنان سعيد حجران

الملخص العربي

تهدف هذه الدراسة إلى معرفة تأثير الجزر والتين الشوكى والهندباء على الفئران المصابة بالتسمم الكبدى. قسمت الفئران (45 فار ذكور بيضاء) إلى 9 مجموعات بكل مجموعة 5 فئران. تم استخدام مجموعتان ضابطتان غذيت المجموعة الأولى على الوجبة الأساسية فقط كمجموعة ضابطة سالبة والمجموعة الثانية فتغنيت على الوجبة الأساسية بعد حقنها برابع كلوريد الكربون كمجموعة ضابطة موجبة ، أما المجموعات الأخرى فتم حقنها برابع كلوريد الكربون ثم تغذيت على الوجبة الأساسية محتوية على نسبة 10% ، 15% من الجزر والتين الشوكى والهندباء و 15% خليط من النباتات المستخدمة . تم تقدير وجود تلف الكبد عن طريق تقدير نشاط وتركيز الإنزيمات في البلازما وهي انزيم اسبرتات ترانس امينيز والانين امينو ترانسفيريز واكوليستيرول الكلى والتراى جلسريد والليبوبروتينات منها الليبوبروتين عالى الكثافة والليبوبروتين منخفض الكثافة جدا وحمض البوريك واليوريا والجلوكوز . أظهرت النتائج وجود تحسن لهذه القياسات باستخدام الجزر والتين الشوكي والهندباء وكانت أفضل النتائج باستخدام نسبة 15% يليها نسبة 10% من هذه النباتات. لذلك نستخلص من هذه الدراسة ان تلف الكبد بالفئران الناجم عن رابع كلوريد الكربون يمكن تحسنه باستخدام من هذه الدراسة ان تلف الكبد بالفئران الناجم عن رابع كلوريد الكربون يمكن تحسنه باستخدام من هذه الدراسة ان تلف الكبد بالفئران الناجم عن رابع كلوريد الكربون يمكن تحسنه باستخدام من هذه الدراسة ان تلف الكبد بالفئران الناجم عن رابع كلوريد الكربون يمكن تحسنه باستخدام