RESEARCH ARTICLE

NEPHROPROTECTIVE EFFECT OF METHANOLIC EXTRACT OF PETROSLINUM CRISPUMS LEAVES AGAINST RIFAMPIN INDUCED NEPHROTOXICITY IN RATS.

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Abstract

The study was conducted for the renal protective activity of methanolic extract of, Petroslinum crispum leaves against renal toxicity induced by rifampin injection in rats. This plant is used in traditional medicine for the treatment of renal failure, inflammation of kidney and other disease. The renal toxicity was produced by administration of rifampin at dose rate 70 mg/kg I/p for 28 days. The plant extract was given by simultaneous oral administration of methanolic extract of Petroslinum crispum leaves at a dose rate of 300 and 600mg/kg for 28 days. It was found that the induced toxicity was inhibited by decreased level of serum urea, creatinin, sodium (Na) and potassium (K). Histopathological examination revealed protection of the kidney from the marked necrosis of renal tubules that induced by rifampin.

Introduction:

The kidneys are a pair of organs located in the back of the abdomen. Each kidney is about 4 or 5 inches long -- about the size of fist. The kidneys' functions are to filter the blood. All the blood in our bodies passes through the kidneys several times a day. The kidneys remove wastes, control the body's fluid balance, and regulate the balance of electrolytes. As the kidneys filter blood, they create urine, which collects in the kidneys' pelvis -- funnel-shaped structures that drain down tubes called ureters to the bladder. Produce rennin, an enzyme that helps regulate blood pressure, Produce erythropoietin, which stimulates red blood cell production and Produce an active form of vitamin D, needed for bone health. Chronic kidney disease (CKD), also known as chronic renal disease, is progressive loss in kidney function over a period of months or years. The symptoms of worsening kidney function are not specific, and might include feeling generally unwell and experiencing a reduced appetite. Often, chronic kidney disease is diagnosed as a result of screening of people known to be at risk of kidney problems, such as those with high blood pressure or diabetes and those with a blood relative with CKD. This disease may also be identified when it leads to one of its recognized complications, such as cardiovascular disease, anemia, pericarditis or renal osteodystrophy (the latter included in the novel term CKD-MBD).¹² CKD is a long-term form of kidney disease; thus, it is differentiated from acute kidney disease (acute kidney injury) in that the reduction in kidney function must be present for over 3 months. CKD is an internationally recognized public health problem affecting 5–10% of the world population.

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Nephrotoxicity is poisonous effect of some substances, both toxic chemicals and medication on the kidneys. Many of the drugs causes toxicity to the kidney, these drugs used in chemotherapy such as Cisplatin, Carboplatin, Carmustine and high dose methotrexate biologic therapy such as Interleukin-2 or Interferon Alfa, antibiotics and antimicrobial such as Amphotericin B, Gentamicin, Vancomycin and Rifampin (Rifadin). Currently, medicinal plants are widely used in the treatment of many diseases owing their safety, cheapness and nontoxicity when compared with the synthetic drugs. It has been estimated that about 80% of the world population relies on natural and botanical preparations as medicine to meet their health needs. Also, the dietary intake of carotenoids, vitamins and flavonoids, which are widely distributed in plants, could be useful in protecting against drug-induced nephrotoxicity.

Tuberculosis (TB), a serious disease caused by Mycobacterium tuberculosis, is a growing international health concern. It primarily affects lungs and secondarily smooths course of antibiotics including isoniazid (INH), rifampicin (RIF), pyrazinamide (PZA) and ethambutol or streptomycin. The use of isoniazid and its combination with rifampicin is associated with liver damage that could result in acute liver failure with a high mortality rate. The rate of their hepatotoxicity is much higher in developing countries compared to that in developed countries, even with using the same dose schedule. Although the hepatotoxic effects of the anti-tuberculosis drugs have been extensively studied, their induced nephrotoxicity have been poorly documented. The study of Mahmouds et al demonstrated that oxidative stress is implicated in the INH-induced toxicity in rats. In addition, Lopez-Novoaetal reported that reactive oxygen species (ROS) and oxidative stress represent the central key in the pathogenesis of drug-induced renal damage. Therefore, the use of antioxidants could offer protection against drug-induced renal damage.

Petroselinum crispum seed was used traditionally as a carminative to decrease flatulence and colic pain. The root was used as a diuretic and the juice to treat kidney ailments. Petroselinum crispum oil also has been used to regulate menstrual flow in the treatment of amenorrhea and dysmenorrhea, and is purported to be an abortive. Bruised leaves have been used to treat tumors, insect bites, lice, skin parasites, and contusions. At one time, parsley tea was used to treat dysentery and gallstones. Other traditional uses include treatment of diseases of the prostate, liver, and spleen. Historically, Petroselinum crispum also has been used in the treatment of anemia, arthritis, and cancers, as an expectorant, antimicrobial, aphrodisiac, hypotensive, diuretic, and laxative. It also has been used as a scalp lotion to stimulate hair growth. However, there have been no clinical trials to confirm these uses.

Material and method:-
Animals:-
Twenty rats Wister white (albino) weighting 100-150gm, were obtained from the atomic energy Research institute. They were housed in laboratory cages, maintained in a room under standard environmental condition, controlled temperature (22±2c), relative humidity (60%) with free access to water and formula rat feed (corn powder, protein). Animals were apparently healthy and there were identified by color tail marks. 10 days was allowed as a preliminary adaptive period.

Plant material:-
Petroselinum crispum leaves belong to the family of Apiaceae were collected in May 2014 from Alkhartoum city. The plant was authenticated by the botanists in medicinal and aromatic plants research institute.

Preparation of extract:-
500 g of Petroselinum crispum was successively extracted by soaking in 80% methanol for about seventy two hours of solvent with daily filtration and evaporation. Solvents were evaporated under reduced pressure to dryness using rotary evaporator apparatus.

Experimental design:
Group A: serving as control for 28 days. Group B: was injected by rifampin I/p-0.8 ml/Kg induced nephrotoxicity for 28 days. Group C: was injected by rifampin I/p-0.8 ml/Kg and treated with the low dose (300mg/Kg) of the
methanolic extract of *Petroselinum crispum* leaves. **Group D:** was injected by rifampin i/p-s 0.8 ml/Kg and treated with high dose (600mg/Kg) of methanolic extract of *Petroselinum crispum* leaves. Serum concentration of urea, creatinine, and K were chemically estimated.

**Result and discussion:**

Kidney injury is a severe complication that can disturb treatment and cause permanent kidney damage. It has been reported that nephropathy is an important microvascular complication of anti-tuberculosis therapy multiple studies have reported that oxidative stress and inflammation are implicated in drug-induced organ toxicity. However, renal injury induced by anti-tuberculosis therapy is usually reversible and could be treated appropriately if detected early. Therefore, the use of antioxidants could protect against RIF-induced oxidative stress and nephrotoxicity. Oral administration of rifampin in dose of 70mg/kg i/p for 28 days to rats caused nephrotoxicity manifested by significant (P < 0.05) increased in serum level of urea, creatinine, Na and K when compared with the (negative) control group. Oral administration of methanolic extract of *Petroselinum crispum* leaves for 28 days induced significant (P< 0.05) decreases in high serum levels of urea, creatinine, Na and k when compared with rifampin intoxicated rats Table (1to 4) and Fig(1 to 4). We designed the current study to test the hypothesis that *petroselinum crispum* leaves extract could protect against Rifampin(RIF)-induced renal damage in rats through attenuation of inflammation and prevention of oxidative stress.

The present findings showed that the administration of Rifampin (RIF) has induced renal injury and glomerular dysfunction as evident by the elevated serum urea, creatinine, sodium and potassium levels. These parameters are often regarded as reliable markers of renal damage [15]. In addition, serum creatinine has been used to estimate glomerular function and its elevation is an indicator of renal failure [16]. The elevated serum markers of renal toxicity is agreement with the study of Hashmi et al [17] who reported increased serum levels of urea and creatinine in albino rabbits following administration of anti-tuberculosis drugs Isoniazid (INH) and Rifampin (RIF) induced nephrotoxicity. In the present study, results suggested that methanolic extract of *petroselinum crispum* 600 mg/kg significantly reduced Rifampin-induced elevated serum levels of creatinine, urea, sodium, and potassium. The histological findings also supported the nephroprotective action of methanolic extract of *petroselinum crispum*. It can be concluded from the present study that nephroprotective effect of *petroselinum crispum* was showed due to the presence of one or more of flavanoids.

**Fig (1)**
Fig(2)

Methanolic extract of Petroselinum Crispum

Fig(3)

Methanolic extract of Petroselinum Crispum
Table (1):

| Groups     | Urea in zero time | Urea in 7 days | Urea in 14 days | Urea in 21 days | Urea in 28 days |
|------------|-------------------|----------------|-----------------|-----------------|-----------------|
| Group A    | 27.00± 2.08       | 29.01± 6.11    | 29.33± 6.36     | 28.67± 5.81     | 29.00± 3.46     |
| Drug       | 26.00± 2.08       | 42.00± 6.11    | 75.33± 6.36     | 85.67± 5.81     | 95.00± 3.46     |
| Low dose   | 19.67± 1.20       | 38.67± 2.33    | 60.33± 3.18     | 57.67± 1.45     | 57.00± 1.53     |
| High dose  | 22.00± 3.06       | 44.00± 3.06    | 63.00± 2.08     | 58.33± .88      | 52.33± .88      |

Table (2):

| Groups     | Creatinine zero time | Creatinine 7 days | Creatinine 14 days | Creatinine 21 days | Creatinine 28 days |
|------------|----------------------|-------------------|--------------------|-------------------|-------------------|
| Group A    | .67± .07             | .63± .42          | .67± .38           | .83± .17          | .50± .32          |
| Drug       | .57± .07             | 2.53± .42         | 3.57± .38          | 4.73± .17         | 5.40± .32         |
| Low dose   | .53± .03             | .57± .03          | 1.83± .15          | 1.73± .07         | 1.73± .07         |
| High dose  | .60± .06             | 1.13± .12         | 1.90± .17*         | 1.80± .10         | 1.43± .03         |

Table (3):

| Groups     | Na zero time | Na 7 days | Na 14 days | Na 21 days | Na 28 days |
|------------|--------------|-----------|------------|------------|------------|
| Group A    | 137.00± .58  | 138.67± 1.20 | 138.67± 1.76 | 138.67± 2.85 | 138.00± 1.15 |
| Drug       | 135.33± .33  | 137.67± 1.20 | 140.67± 1.76 | 146.67± 2.85 | 151.00± 1.15 *|
| Low dose   | 136.00± 1.53 | 143.67± .33  | 151.33± 2.03  | 149.67± .88  | 148.00± 1.53 *|
| High dose  | 136.67± .88  | 144.67± 1.45  | 149.00± 2.08  | 150.67± .88 * | 145.00± .00   |
Table (4):

| Groups      | K zero time | K 7 days | K 14 days | K 21 days | K 28 days |
|-------------|-------------|----------|-----------|-----------|-----------|
| Group A     | 3.60±.12    | 3.83±.03 | 4.23±.09  | 4.07±.09  | 4.03±.09  |
| Drug        | 3.53±.09    | 3.73±.03 | 4.13±.09  | 4.97±.09  | 5.83±.09  |
| Low dose    | 3.67±.12    | 4.30±.15 | 5.10±.21  | 5.23±.13  | 5.17±.12  |
| High dose   | 3.53±.09    | 4.30±.21 | 5.43±.23  | 5.20±.12  | 4.90±.06  |

Histopathological examination:

Histological examination of kidneys of normal rats showed normal histological structure of renal glomeruli and tubules (Fig.5A). Kidney of rats intoxicated with rifampin (70 mg/kg) for 28 days revealed marked tubular necrosis (Fig.5B) Kidneys of GM - nephrotoxic rats received the low dose (300 mg/kg) of parsley leaves extract showed large vaculations and peritubular leukocytes infiltration (Fig.5 C). Kidneys of nephrotoxic rats pre-treated orally with the large dose (600mg/kg) of parsley leaves extract showed almost normal histological architecture of renal glomeruli and tubules (Fig.5D).
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