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2 **Biochemical and molecular studies on the role of**
3 **rosemary extract in reducing liver and kidney**
4 **toxicity due to Vepesid in male rats**
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8
9 **ABSTRACT**
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Aims: Vepesid is chemotherapeutic drugs that inhibit topoisomerase II activity and long been used for treatment of human malignancies, where it is a semi-synthetic compound derived from the plant podophyllum pelltatum. The current study was designed to investigate the possible protective effect of rosemary extract against Vepesid -induced changes in liver and kidney functions, and DNA damage in rats.

Materials and Methods: A total of 50 male Wister albino rats were divided randomly into four groups (1st group was control; 2nd group was treated with rosemary, 3rd group was received Vepesid, and 4th & 5th groups was co- and post treated groups respectively).

Results: The administration of Vepesid revealed a significant increase in serum ALT, AST, ALP, creatinine, urea, potassium ions, chloride ions, and DNA damage. In contrast; a significant decrease in albumen, total proteins, sodium ions, and calcium ions were when compared with control group. This increased in ALT, AST, ALP, creatinine, urea, potassium ions, chloride ions, and DNA damage was reduced after administration of rosemary when co-treated with Vepesid (G4), or post-treated after Vepesid (G5) for four weeks with lowest damage in G4. Also; this decreased in albumen, total proteins, sodium ions, and calcium ions was increased after administration of rosemary when co-treated with Vepesid (G4), or post-treated after Vepesid (G5) for four weeks with lowest damage in G4.

Conclusion: it could be concluded that rosemary has a promising role and it worth to be considered as a natural substance for protective the liver and kidney toxicity induced by Vepesid chemotherapy.

11
12 *Key words:* Chemotherapy; Vepesid; Rosemary; Liver; Kidney; Rat.
13

14 **1. INTRODUCTION**
15

16 Today, there are many different kinds of chemotherapy that used for cancer treatments. It is therefore
17 important to search for therapies which can reduce the side effects of anticancer treatments without
18 altering their efficacy or increasing toxicity or damage in target organs [1-8]. Vepesid or VP-16 is the
19 trade name for etoposide. Etopophos and toposar or etoposide phosphate are other names for Vepesid.
20 In some cases, health care professionals may use the trade name VP-16 or other names Vepesid or
21 etopophos or toposar or etoposide phosphate when referring to the generic drug name etoposide.
22 Vepesid is chemotherapeutic drugs that inhibit topoisomerase II activity and long been used for
23 treatment of human malignancies, where it is a semi-synthetic compound derived from the plant
24 podophyllum pelltatum [9,10].

25 Vepesid is commonly used alone or with another anticancer agent for the treatment of Hodgkin's
26 lymphoma and AID's and sexual organ cancers as testicular, ovarian, uterine, bladder and prostate or
27 for the treatment of other organs as lung and stomach cancer [10]. Although Vepesid is effective in

28 the treatment of different types of cancers, it causes the death of normal proliferating cells, including
29 male germ cells [9].

30 Many plant extracts and their products have been shown to have significant antioxidant activity which
31 may be an important property of medicinal plants associated with the treatment of several ill-fated
32 diseases including liver toxicity [11-19].

33 Rosemary (*Rosemarinus officinalis*) is one of household herbs that contains a number of
34 phytochemicals, including rosmarinic acid, camphor, caffeic acid, ursolic acid, betulinic acid, and the
35 antioxidants carnosic acid and it used in traditional medicine to treat a variety of disorders [20-22].
36 Extracts of rosemary leaves contains flavonoids and phenols and it possess a variety of bioactivities in
37 vitro including anti-tumor, antioxidant, antibacterial, antinociceptive, antidiabetic, antithrombotic,
38 antiulcerogenic, antidiuretic and anti-inflammatory agents [10,22]. Therefore; the present study was
39 conducted to examine the possible modifying effects of rosemary aqueous extract against the changes
40 in the liver and kidney function and DNA damage, induced by Vepesid in male rats.

41

42 **2. MATERIAL AND METHODS**

43

44 **2.1. Animals**

45 The experiments were performed on 50 male rats weighing 130 ± 10 g and of 10 week's age. The rats
46 were kept in our Faculty animal house for one week before the experimental work and maintained on
47 a standard rodent diet and water available *ad libitum*. After one weeks of acclimation, rats were
48 equally divided into four groups.

49

50 **2.2. Experimental groups**

51 Rats were equally divided into four groups.

52 1st group: Control group included rats received no treatment.

53 2nd group: Rosemary group included rats received by oral gavages rosemary extract at a dose of (220
54 mg/kg b.w. /twice weekly) for six weeks [21].

55 3rd group: Vepesid group included rats that injected intraperitoneally with Vepesid (1 mg /kg
56 B.W/day) for six weeks [10].

57 4th group: Co-treated group included rats that injected by Vepesid (1 mg/kg B.W. /day) for six weeks
58 and received rosemary (220 mg /kg b.w. /twice weekly) orally for the same six weeks.

59 4th group: Post treated group included rats that injected by Vepesid (1 mg/kg B.W. /day) for six weeks
60 and then received rosemary (220 mg /kg b.w. /twice weekly) orally for another six weeks.

61 At the end of the experimental period, rats were fasted overnight; euthanized with intraperitoneal
62 injection with sodium pentobarbital and subjected to a complete necropsy. Blood samples were
63 individually collected from the inferior vena cava of each rat in non-heparinized glass tubes for
64 estimation of liver and kidney functions biomarkers [23]. Blood samples were incubated at room
65 temperature for 10 minutes and left to clot then centrifuged at 3000 r.p.m for 15 min and the serum
66 were collected, serum was separated and kept in clean stopper plastic vial at -80°C until the analysis
67 of serum parameters.

68

69 **2.3 Liver function Biomarker:**

70 Alanine transaminase (ALT) and aspartate transaminase (AST) activities in serum were assayed by
71 using commercial kit that was supplied by Humann (Germany) according to the method of Tousson et
72 al. [24] and Bolkinny et al. [25] respectively while alkaline phosphatase (ALP) was estimated in the rat
73 serum according to El-Moghazy et al. [11]. Serum albumin was estimated according to Basuony et al.
74 [6] while serum total proteins level was estimated according to Tousson et al. [26].

75

76 **2.4 Electrolytes and kidney functions Biomarker:**

77 Serum urea and creatinine were determined in the rat sera according to Salama et al. [27] and Eldaim
78 et al. [28] respectively. To measure the levels of serum electrolytes (Potassium, sodium, calcium and
79 chloride ion) by using commercial kits (Sensa core electrolyte, India) according to El Atrash et al.
80 [29].

81

82 **2.5 Comet assay**

83 One gram of crushed kidney tissue was transferred to 1 mL of ice-cold phosphate-buffered saline
84 (PBS) and the assay was performed according to Eldaim et al. [28], for visualization of DNA damage,
85 observations were carried out on GelRed-stained DNA using a 40× objective on a fluorescent
86 microscope. Komet 5 image analysis software developed by Kinetic Imaging, Ltd. (Liverpool, UK)
87 linked to a CCD camera was used to assess the quantitative and qualitative extent of DNA damage in
88 the cells by measuring the length of DNA migration and the percentage of migrated DNA. Finally, the
89 program calculates tail moment. Generally, 50–100 randomly selected cells are analysed per sample.
90

91 **2.6. Statistical Analysis**

92 Data were expressed as mean values ± SE and statistical analysis was performed using one way
93 ANOVA to assess significant differences among treatment groups. The criterion for statistical
94 significance was set at $P < 0.01$ for the biochemical data. All statistical analyses were performed using
95 SPSS statistical version 16 software package (SPSS® Inc., USA).
96

97 **3. RESULTS**

98

99 **3.1. Serum markers of liver damage**

100 Data presented in Figure (1) showed that serum ALT, AST, ALP and total bilirubin levels were
101 significantly ($P < 0.05$) increase in treated rats with Vepesid as compared to control group. In contrast;
102 a significant ($P < 0.05$) decrease in total protein and albumin levels in treated rats with Vepesid as
103 compared to control group (Figure 1). Treatment of rats with Vepesid and rosemary (as in G4&G5)
104 revealed a significant ($P < 0.05$) decrease in ALT, AST, ALP and total bilirubin levels and a significant
105 ($P < 0.05$) increase in total protein and albumin levels when compared with treated rats with Vepesid
106 (Figure 1). Also; with lowest damage in G4

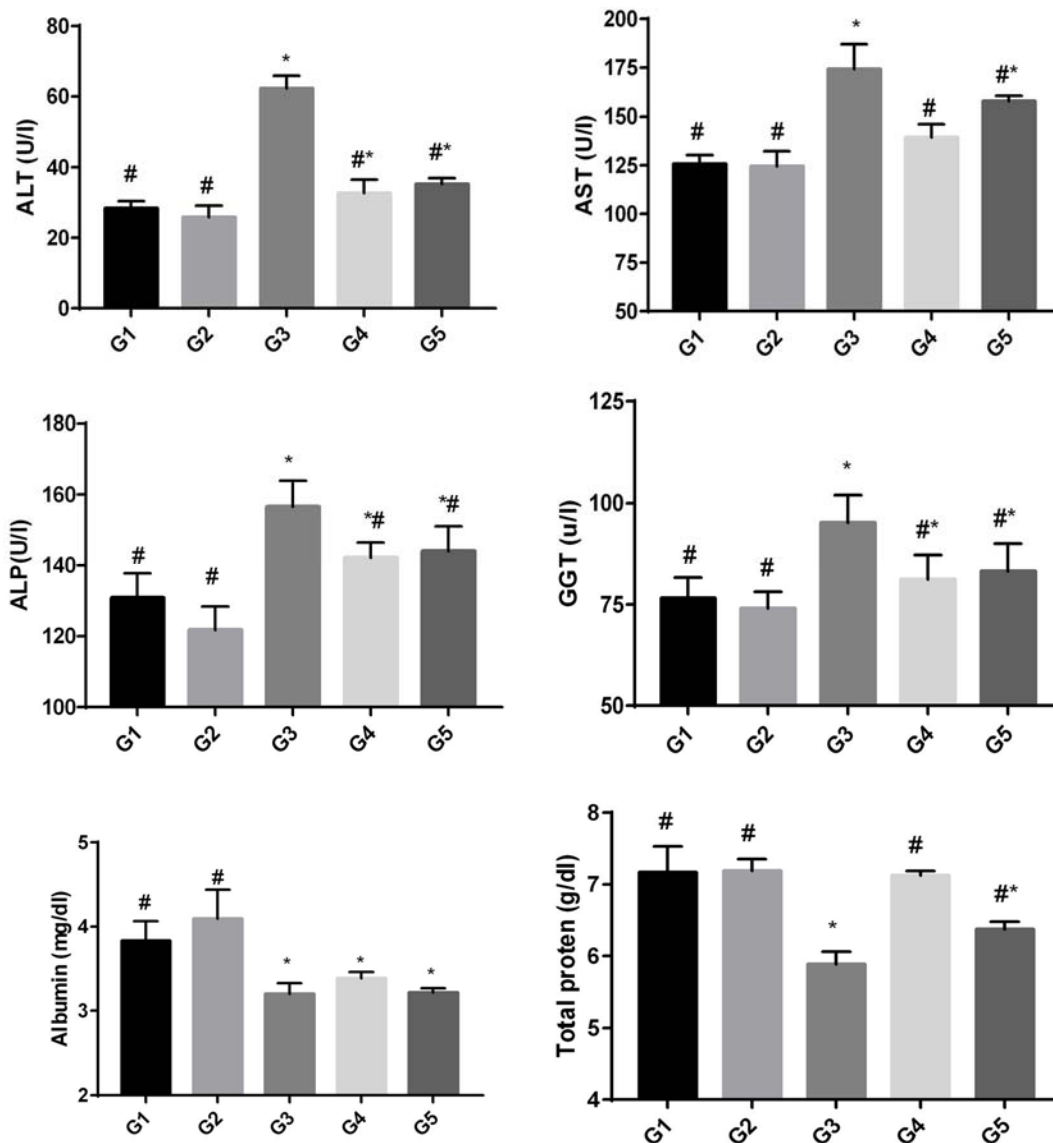
107 **3.2. Serum markers of kidney damage**

108 Data presented in Figure (2) showed that serum creatinine, urea, potassium and chloride ions levels
109 were significantly ($P < 0.05$) increase in treated rats with Vepesid as compared to control group. In
110 contrast; a significant ($P < 0.05$) decrease in serum sodium and calcium ions levels in treated rats with
111 Vepesid as compared to control group (Figure 2). Treatment of rats with Vepesid and rosemary (as in
112 G4&G5) revealed a significant ($P < 0.05$) decrease in creatinine, urea, potassium and chloride ions
113 levels and a significant ($P < 0.05$) increase in sodium and calcium ions levels when compared with
114 treated rats with Vepesid (Figure 1).

115 **3.3. DNA damage in liver tissues**

116 A comet assay was performed to assess DNA damage in liver of rats after treatment by rosemary
117 and/or Vepesid as compared to normal control. The results of comet assay were shown in Figures (3)
118 and Tables (1). Administration of Vepesid (G3) led to significant increase in liver DNA damage ($P <$
119 0.05) that was indicated by increase in tail length, tail DNA% and tail moment as compared to normal
120 control (G1) and rosemary (G2) groups (Table 1 & Figure 3). This increased liver DNA damage was
121 reduced after administration of rosemary when co-treated with Vepesid (G4), or post-treated after
122 Vepesid (G5) for four weeks with lowest damage in G4. On the other hand, no significant difference
123 in liver DNA damage (tail length) was observed between normal control (G1) and rosemary treated
124 groups (G2).

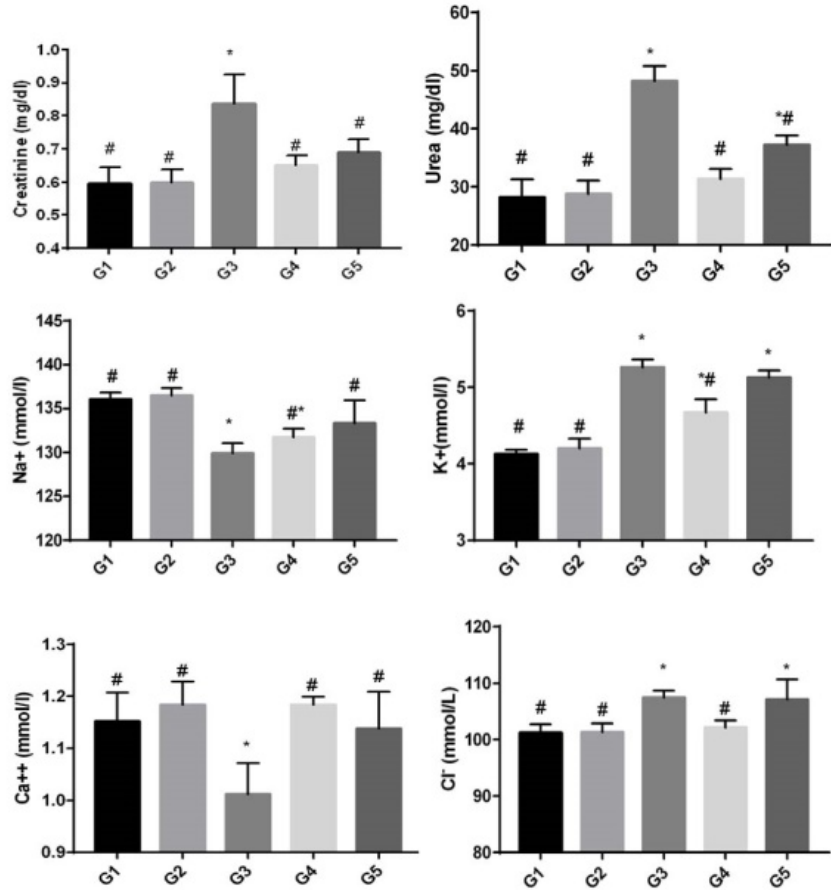
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126

127 **Figure 1:** Changes in serum ALT, AST, ALP, GGT, total protein and albumin levels in different
 128 groups under study. Data are expressed as mean \pm SE of 10 observations. *Significant difference from
 129 control group at $P < 0.05$. #Significant difference from Vepsid group at $P < 0.05$. Where G1, control
 130 group; G2, rosemary group; G3, Vepsid group; G4, co-treated Vepsid with rosemary group; G5, post-
 131 treated Vepsid with rosemary group.

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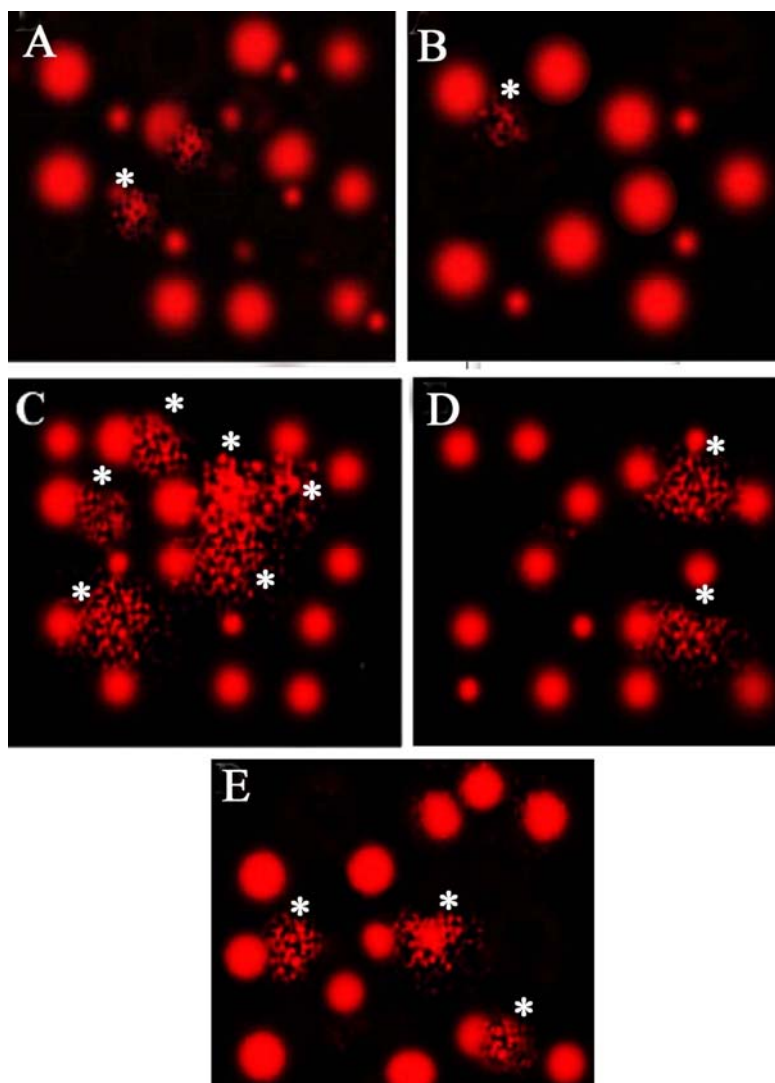
134 **Figure 2:** Changes in serum kidney functions (creatinine and urea) and electrolytes (sodium,
 135 potassium, calcium and chloride ions) levels in different groups under study. Data are expressed as
 136 mean \pm SE of 10 observations. *Significant difference from control group at $P < 0.05$. #Significant
 137 difference from Vepsid group at $P < 0.05$. Where G1, control group; G2, rosemary group; G3, Vepsid
 138 group; G4, co-treated Vepsid with rosemary group; G5, post-treated Vepsid with rosemary group.

139 **Table 1:** Comet assay parameters obtained by image analysis in liver cells of all groups after
 140 prevention experiment.

Group	Tailed %	Untailed %	Tails length μm	Tail DNA%	Tail moment
G1	4	96	1.79 \pm 0.35 ^d	1.68	3.04
G2	1.5	98.5	1.34 \pm 0.10 ^d	1.39	1.86
G3	19	81	6.55 \pm 0.34 ^a	5.05	33.08
G4	9	91	3.73 \pm 0.13 ^c	3.02	11.26
G5	12	88	4.62 \pm 0.21 ^b	3.70	17.09

141 Different superscript letters in the same column of tail length showed significance difference at $P <$
 142 0.05. Where G1, control group; G2, rosemary group; G3, Vepsid group; G4, co-treated Vepsid with
 143 rosemary group; G5, post-treated Vepsid with rosemary group.

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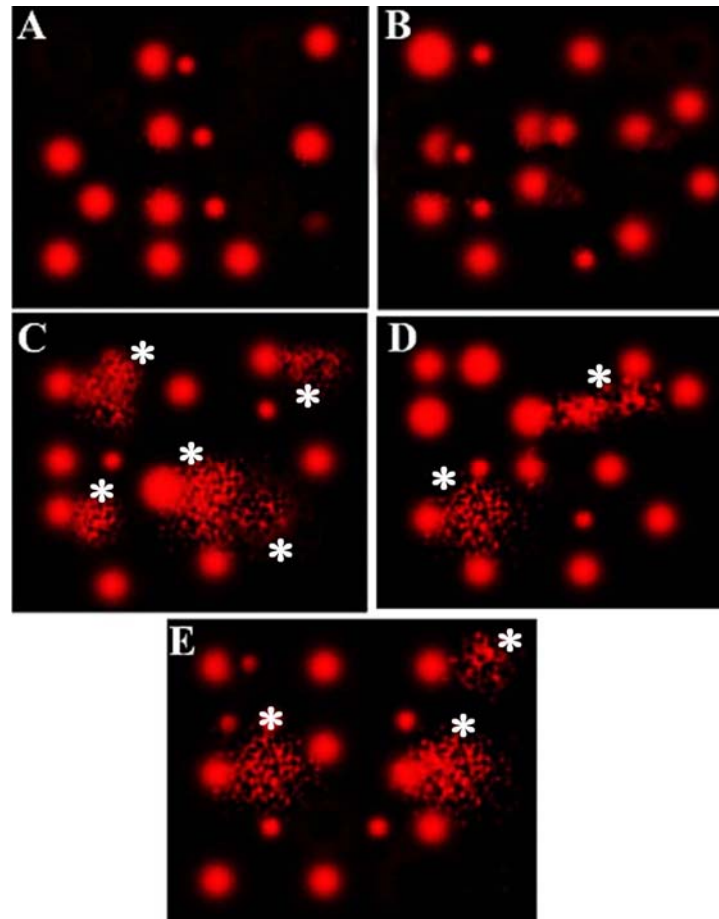
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147 **Figure 3:** Photomicrographs representation of DNA damage in liver tissues, using comet assay, in
 148 normal control group (A), rosemary group (B), Vepsid group (C), co-treated Vepsid with rosemary
 149 group (D), and post-treated Vepsid with rosemary group (E).

150

151 3.4. DNA damage in kidney tissues

152 A comet assay was performed to assess DNA damage in kidney of rats after treatment by rosemary
 153 and/or Vepsid as compared to normal control. The results of comet assay were shown in Figures (4)
 154 and Tables (2). Administration of Vepsid (G3) led to significant increase in kidney DNA damage ($P <$
 155 0.05) that was indicated by increase in tail length, tail DNA% and tail moment as compared to normal
 156 control (G1) and rosemary (G2) groups (Table 2 & Figure 4). This increased kidney DNA damage
 157 was reduced after administration of rosemary when co-treated with Vepsid (G4), or post-treated after
 158 Vepsid (G5) for four weeks with lowest damage in G4. On the other hand, no significant difference
 159 in kidney DNA damage (tail length) was observed between normal control (G1) and rosemary treated
 160 groups (G2).



162

163 **Figure 4:** Photomicrographs representation of DNA damage in kidney tissues, using comet assay, in
 164 normal control group (A), rosemary control group (B), positive control group (C), co-treated Vepsid
 165 with rosemary group (D), and post treated Vepsid with rosemary group (E).

166 **Table 2:** Comet assay parameters obtained by image analysis in cells of all groups after treatment
 167 experiment.

Group	Tailed %	Untailed %	Tails length μm	Tail DNA%	Tail moment
G1	1.5	98.5	1.36 ± 0.11^d	1.46	1.99
G2	3	97	1.48 ± 0.12^d	1.60	2.37
G3	16	84	5.70 ± 0.35^a	4.71	26.85
G4	7	93	3.11 ± 0.17^c	2.28	7.09
G5	11	89	4.47 ± 0.12^b	3.51	15.69

168

169 Different superscript letters in the same column of tail length showed significance difference at $P < 0.05$. Where G1, control group; G2, rosemary group; G3, Vepsid group; G4, co-treated Vepsid with
 170 rosemary group; G5, post-treated Vepsid with rosemary group.

171 **4. DISCUSSION**

172

173 Chemotherapy involves the use of chemical agents to stop the growth and eliminate cancer cells even
174 at distant sites from the origin of primary tumor [6]. However, it does not distinguish between a
175 cancer and normal cells, and eliminates not only the fast-growing cancer cells but also other fast-
176 growing cells in the body, including, hair and blood cells. The current study aimed to study the
177 protective and ameliorating effects of rosemary extract against liver toxicity induced by Vepsid in
178 male albino rats.

179 Chemotherapy-induced hepatotoxicity is a common cause of abnormal liver function test in patients,
180 this hepatotoxicity are usually begins with vague clinical symptoms such as fatigue, anorexia, nausea,
181 dark urine, right upper quadrant discomfort and jaundice. In the current study; a significant increase in
182 serum ALT, AST, ALP and a significant decrease albumen and total proteins indicated the liver
183 toxicity were detected after the treatments of rats with Vepsid as compared with control. This result is
184 in harmony with Abouzeinab [30]; Nasr [31]; Abdel-Wahhab et al. [32] and Basuony et al. [6] who
185 reported that, Cisplatin administration induced an increase in ALT, AST, ALP and decrease albumen
186 and total proteins. Also; this current result is in harmony with Tousson et al. [3,4] who reported that;
187 methotrexate-induced hepatic and renal toxicity in male rats and the increased in liver function
188 associated with free radicals trigger cell damage through binding to cellular macromolecules. Similar
189 findings were reported by Juma [33] and McDonald et al. [34] who reported that; cyclophosphamide -
190 Induced hepatotoxicity in human liver. Elevated levels of serum ALT and AST enzymes are
191 indicative of cellular leakage and loss of functional integrity of cell membranes in the liver [12,24].
192 The estimation of these enzymes in the serum is a useful quantitative marker for the extent and type of
193 hepatocellular damage [7,8,35].

194 Urea and creatinine are nitrogenous end products of metabolism. Urea is the primary metabolite
195 derived from dietary protein and tissue protein turnover [27,36,37]. Creatinine is the product of
196 muscle creatine catabolism. Both are relatively small molecules (60 and 113 daltons, respectively)
197 that distribute throughout total body water. The rationale for the use of creatinine or urea
198 measurement to assess renal function is that plasma/serum levels of both reflect glomerular filtration
199 rate (GFR), the parameter that defines kidney function for the clinician. Irrespective of its cause,
200 kidney disease is associated with decrease in GFR, and the severity of kidney disease correlates
201 closely but inversely with GFR [38].

202 Chemotherapy-induced renal tototoxicity is a common cause of abnormal kidney function test in
203 patients and in animal models. Renal injury may follow treatment with anticancer drugs and lead to
204 glomerular, tubular dysfunctions, or any combination of these [39]. Nephrotoxicity is an unusual side
205 effect of chemotherapy in general. Most chemotherapy drugs target pathways that are essential to
206 dividing cells. In the current study; serum creatinine, urea, potassium and chloride ions levels were
207 significantly ($P<0.05$) increase in treated rats with Vepesid as compared to control group. In contrast;
208 a significant ($P<0.05$) decrease in serum sodium and calcium ions levels were detected in treated rats
209 with Vepesid as compared to control group. Mechanisms of anticancer drug-induced renal disorders
210 generally include a varying degree of prerenal hypoperfusion, intrinsic renal damage, renal tubular
211 obstruction, and damage to the microvascular structure of the kidneys [40]. Our result is agree with
212 Tousson et al. [1] who find that MTX increased urea and creatinine activities which induced renal
213 toxicity. Our result is agreed with Basuony et al. [6] who reported that; Cisplatin induced renal
214 toxicity in rats. On the other hand, our results are disagreement with Cetiner et al. [41]. Our result is
215 agreed with Beyer et al. [42] who reported that; High-dose carboplatin, etoposide and ifosfamide
216 induced renal toxicity in human. Also; our result is agreed with Al-Ameri [43] who reported
217 Etoposide induced kidney toxicity, electrolytes changes and injury. Chemotherapy-induced
218 nephrotoxicity is a major cause of morbidity and mortality among cancer patients. Therefore,
219 assessing baseline renal function before initiation of therapy and during therapy, adjusting drug
220 dosages, avoiding nephrotoxic drug combinations, and correcting the extracellular fluid volume
221 depletion is essential in the cancer patients [44].

222 Treatment of rats with Vepesid and rosemary revealed a significant decrease in creatinine, urea,
223 potassium and chloride ions levels and a significant ($P<0.05$) increase in sodium and calcium ions

224 levels when compared with treated rats with Vepesid indicated that rosemary has renal protective
225 against chemotherapy. The topoisomerase II inhibitor etoposide is an antineoplastic drug that has been
226 widely used to couple DNA damage to apoptosis [45]. Topoisomerase II is a nuclear enzyme that
227 functions during both DNA replication and transcription [46]. Topoisomerase II prevents “knots”
228 from forming in DNA by allowing the passage of an intact segment of the helical DNA through a
229 transient double strand break [47]. Topoisomerase II inhibitors such as etoposide stabilize the
230 complex formed by topoisomerase II and the 5'-cleaved ends of the DNA, thus forming stable
231 (nonrepairable) protein-linked DNA double strand breaks [47]. Cells are apparently able to recognize
232 such DNA damage and, in turn, to eliminate the injured cells by apoptosis. In the current study,
233 treatment of rats with Vepesid led to significant increase in liver and kidney DNA damage ($P < 0.05$)
234 that was indicated by increase in tail length, tail DNA% and tail moment as compared to normal
235 control and rosemary groups. This increased kidney DNA damage was reduced after administration of
236 rosemary when co-treated with Vepesid (G4), or post-treated after Vepesid (G5) for four weeks with
237 lowest damage in G4. Our results agree with Tousson et al. [10] who reported that; Etoposide
238 induced DNA damage in testicular tissues.
239

240 **5. CONCLUSION**

241
242 Our recommendation is Etoposide treatments induced changes in liver and kidney functions and DNA
243 damage. Physicians should be aware of Etoposide a differential diagnosis for hepatic and renal with
244 an unknown etiology. Rosemary has a promising role and it worth to be considered as a natural
245 substance for protective the liver and kidney toxicity induced by Vepesid chemotherapy.
246

247 **Conflict of interests**

248 The authors declare no conflict of interest.
249

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