Thymoquinone Pre-and Post-Treatment Effects on Ehrlich Ascites Carcinoma Induced Albino Mice: An in vivo study.

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ABSTRACT

Background: Chemoprevention is an effective approach towards cancer regulation. For that, it is important nowadays to investigate more active and influential natural products, which have a powerful capability to fight cancer cells. Thymoquinone was the selected natural product for the current study to investigate its probable anticancer property in Ehrlich ascites carcinoma (EAC) inoculated mice within both cases of pre-and post-treatment. Objective: The present research was assumed to evaluate both the protective and therapeutic anti-cancer influence of Thymoquinone (TQ) against Ehrlich ascites carcinoma (EAC) inoculated mice. Experimental mice were split into five sets: 1) control group, 2) TQ group: mice were orally injected with (10mg/kg) for four weeks. 3)EAC group: mice were intramuscularly inoculated with EAC (2.5x10^6) cells once, 4) TQ+EAC group: mice were previously orally injected with TQ two weeks before experiment, then were intramuscularly inoculated with EAC, 5) EAC+TQ group: mice were injected with EAC and TQ at the same time till last day of experiment. All animals’ liver and tumors were collected and examined histologically (H&E), histochemical (Feulgen stain reaction), ultra structurally (transmission electron microscope) and immunohistochemically (PCNA antigen & Ki-67 antigen). Results: All results demonstrated that treatment with TQ showed alike features like normal and TQ groups through varying degrees. In addition, there was a great superiority of TQ pre-treated group than post-treated one. Conclusion: this study evidently illustrated that Thymoquinone could perfectly prevent and/or protect cancer incidence, effectively retarding tumor progress and so, it is significantly recommended to be considered as a part of our daily diet. Overall, extra evaluation of Thymoquinone is strongly recommended in clinical fields to outline its possible value to be utilized as an innovative complementary remedy for cancer diseases.

INTRODUCTION

Cancer is considered a common reason of death all over the world (Global Burden of Disease Cancer and others, 2015). EAC is one of the major suggested tumor modeling in experimental research. It is defined as an undifferentiated carcinoma with great transplantable ability, quick proliferation, tinier life period, one hundred percent malignancy. EAC does not possess tumor-specific transplantation antigen (TSTA) (Ozaslan and others, 2011). Routine medications have been unsuccessful or even non-influential for intact cells, so, from this point, exploiting of natural agents as an alternative cancer therapy is believed to be valuable for cancer management and destruction. Chemoprevention is considered to be a promising approach for cancer inhibition today, that is identified as utilizing of natural agents (individually or combination) to restrain cancer progression. Recently, the use of freshly synthesized small particles, the combination of naturally occurring substances or therapeutic modalities to inhibit cancer recurrence and/or progress has become broadly established(Apryshko and others, 2005; Stojkovic and Radacic, 2002; Xie and others, 2006). Thymoquinone comprises the principal component...
of *Nigella sativa* seeds. It exhibited anti-neoplastic, anti-neoplastic, and anti-inflammatory actions in both *in vivo* and *in vitro* (Asaduzzaman Khan and others, 2017; Gali-Muhtasib and others, 2006).

**MATERIALS AND METHODS**

2.1. Animals:

Fifty male Swiss albino mice, 6-8 weeks age, weighing 19±2 g, were purchased from Theodore Bilharz Research Institute, Egypt. Animals and their care were in conformity with the National Institutes of Health (NIH) policies for laboratory animals nursing and use. Mice were accommodated in metabolic cages under well-organized environmental circumstances (25°C and a 12 h light/dark cycle). Animals were always supplied with tap water and standard diet.

2.2. Materials:

Thymoquinone was purchased from Sigma – Aldrich Company Ltd. (The old Brickyard, New Road, Gillinham, United Kingdom). EAC was kindly provided by The Pharmacology and Experimental Oncology Unit of the National Cancer Institute, Cairo University, Egypt, and was preserved through continuous inoculation of (2.5 x10⁶) cells intraperitoneally. (Badr El-Din et al., 2007) (Badary and others, 2007). The ascetic fluid was gathered via an insulin syringe from the intraperitoneal cavity.

2.3. Experimental design:

Fifty mice were separated into 10 mice for each cage. Group (1): was considered as the typical control. Group (2): based on LD50 (TQ) was administrated to mice orally via gavage after being dissolved in bi-distilled water at a dose of 10 mg/kg/day for 4 sequential weeks (Al-Ali and others, 2008; Badary and others, 1998; Nagi and others, 1999). Group 3: (EAC) 0.2mL Ehrlich Ascites Carcinoma (EAC) (2.5x10⁶ cells) were intramuscularly injected at the lower limb’s right thigh only once at the beginning of the experiment. Group (4): (TQ+EAC) animals were pre-treated with TQ (10 mg/kg/day) orally for 2 weeks before intramuscularly inoculated with 0.2mL (EAC) (2.5x10⁶ cells) similarly then left to the end of the experiment. Group (5): (EAC+TQ) mice were intramuscularly injected alike with 0.2mL (EAC) (2.5x10⁶ cells) then they received TQ (10 mg/kg/day) orally after 8 days for a whole month. All mice were sacrificed and samples were collected after 4 weeks.

2.4. Histopathological investigation:

All liver and tumor samples were collected, washed three times cautiously via saline solution, then fixed in 10% prepared formalin solution, processed and converted to paraffin blocks. A thickness of 4-μm sections were cut. Haematoxylin and eosin stained slides were used for the histological examination.

2.5. Electron Microscopy:

The common technique for transmission electron microscope for specimen preparation was applied (Kasas and others, 2003) consequently, samples examination were performed through an electron microscope (TEM, 100CX, JEOL, USA) (Kasas and others, 2003).

2.6. Histochemical Investigations (*Feulgen reaction for DNA*):

Further sections were stained by Periodic acid-Schiff stain for DNA detection (Garvin and others, 1976).

2.7. Immunohistochemical investigation:

Slides with tissues were deparaffinized, hydrated in descending sequences of alcohol, and kept in antigen retrieval (slides were boiled at 98°C in 10 mmol/L sodium citrate buffer for 20 minutes). To block endogenous peroxidase sections were treated with 3% H₂O₂. Monoclonal antibody mouse MAb anti-rat Ki-67 and mouse MAb anti-PCNA were used. Consequently, sections were processed classically according to (Horiguchi and others, 2007). Assessment of Ki-67 and PCNA labeling index was dependent upon the staining intensity and the proportion of +ve cells, where 5-8 areas per specimen were randomly chosen. Results were represented by the mean number of counted +ve cells per particular area.
2.8. Statistical analysis:
Data were represented as the mean ± SE. Statistics were estimated with SPSS program. Significant variances between groups were calculated by means of One-way ANOVA test and when P-value < 0.05, this reflected significant variation.

3. Results:
3.1. Body weight changes:

Figures (i) represented the effect of TQ administration on body weight (g), tumor weight (g) in Ehrlich Solid Carcinoma bearing mice.

![Body weight (g) and tumor weight (g) in control and different treated group.](image)

**Fig.(i):** Body weight (g) and tumor weight (g) in control and different treated group.

3.2. Body weight (BW):

Figure (i) showed the effect of TQ administration in initial, final and net final BWs changes in Ehrlich solid carcinoma (ESC) bearing mice.

As shown in figure (i), TQ injected animals displayed insignificant variation from control in body weight gain. Whereas EAC inoculated group showed significant decrease in body weight gain compared to normal one. This decrease was overcome in (TQ+EAC) injected mice when compared to ESC group. (EAC+TQ) group revealed also a considerable rise in body weight compared to EAC inoculated one but with lower degree than those of (TQ+EAC) group.

3.3. Histological investigations:

Light microscopic examination of the liver of control animals treated with saline solution only showed healthy hepatic construction which was composed of hepatic lobules with radiating cords that are made up of hepatocytes extending from the portal region to central vein. Sinusoids located between cords and lined with endothelial cells. Space of Disse was localized between the sinusoid and the hepatocyte. Von Kupffer cells were projected into the sinusoid lumen Figure 1a.

Examination of liver sections from Thymoquinone treated animals displayed classical hepatic structure resembling those of control animals Figure 1b.
Ehrlich tumor cells inoculated into animals induced intramuscular tumors at the injected spot. These masses were protruding and showed quick progression associated with various inflammatory reaction and lymphocytic response demonstrating a constant proliferation action Figure 1g.

In the specimen examined after one month of subcutaneous inoculation with a single dose of Ehrlich tumor cells led to severe pathological alterations including perivascular round cell infiltration. Hydropic degeneration (Oedema) was also seen in severely injured hepatocytes, large numbers of liver cells had undergone degeneration, such as; focal necrosis, congestion in the central vein, and infiltration of lymphocytes was obviously seen Figures 1c, 1d.

Abnormalities of the hepatocyte nuclei like multinucleated giant cells, pyknosis, and karyorrhexis were also noted. Activated Kupffer cells and hepatic cirrhosis were obviously seen among expanded sinusoidal spaces. EAC inoculated animals pre-treated with TQ group failed to develop solid tumors. Remarkable restoration of normal cell structure was noticed. The hepatocytes were regular, with normal size, normal homogenous cytoplasm with a rounded nucleus. Cells nuclei size was alike that of control one, also cells containing two nuclei were significantly less than in EAC group. An absence of cirrhosis, bile duct proliferation, and carcinoma was noticed. The hepatotoxicity produced by ESC cells were almost eliminated by the protective effect of TQ extract administration Figure 1e. In ESC inoculated animals post-treated with TQ, solid tumor size were reduced about 69.47%. However, there was no effect on skin and revealed by normal hair follicles and skin. Increased necrosis as well as a noticeable reduction in mitosis occurrence indicating slow development. There was an evident drop in tumor size and appeared growing slow and broken into parts after treatment with TQ. Fewer sinusoids dilation and reduction of leucocytic infiltration were also significant. However, Kupffer cells still active. In addition, few foci spots of inflammatory and/or necrotic cells can be seen. Chromatinolysis, pyknosis, and apoptosis were the major nuclear modifications detected within mice. This point to a limited prevention of ESC cells by TQ extract Figure 1f.
Fig. 1: Photomicrograph of liver and tumor sections in mice stained with Hx&E (a,b) control and TQ treated groups respectively illustrating normal hepatocyte (HC), blood sinusoid (BS), central vein (CV), normal endothelial cells (EC) and Kupffer cells (KC). (c,d) Ehrlich inoculated group showed great injured hepatic showing focal necrosis (N), Pyknotic nucleus (PK), leukocytic infiltration (LI), portal vein (PV), activated Kupffer cells (KC), karyorrhexis (*), and feathery degeneration (arrows) hemorrhage (arrows) respectively, (e) TQ pre-treated EAC inoculated group with dilated portal vein (PV) with bile duct (BD), Kupffer cell (KC), pyknotic nuclei (PK), binucleated hepatocyte (arrows), and necrosis (arrow heads), (f) Post-treated EAC inoculated group demonstrated, giant cell (arrow), dilated blood sinusoid (BS), activated Kupffer cell (KC) and karyorrhexis (Kr), (g,h) A longitudinal section of Ehrlich tumor bearing mouse thigh muscle group and TQ post-treated EAC inoculated group tumor respectively showing focal necrosis (N), and muscle fibers (M), invaded by sheets of tumor cells (arrow heads), and the last photo displayed almost normal muscle fibers (M) pressing on tumor cells (arrow heads). (X 400).

3.4. Histochemical investigations:

The use of Feulgen histochemical technique was for the assessment of apoptosis. Control group showed a higher content of coarse chromatin that appeared magenta red in color inside the nuclei of cells indicates the presence of DNA with evident mitotic figures in some cells. The cytoplasm showed a negative Feulgen reaction indicating the absence of any detectable DNA inclusion Figure 2a.

In TQ treated mice showed similar results as that of control group Figure 2b. Hepatocytes of Ehrlich inoculated mice showed a marked decrease in the DNA content seemed faintly stained suggesting that the DNA has probably degraded Figure 2c,2d.

The EAC group previously treated with TQ liver sections had a moderate reaction than the control group, but more than the EAC inoculated rats post-treated with TQ Figure 2e. The EAC treated with TQ previously showed a rise in colored DNA constituent parts. The inflammatory cells included nuclei were densely stained Figure 2f.
Fig. 2: Photomicrograph of liver mice stained with Feulgen reaction (a,b) control and TQ treated group showed normal content of DNA in the hepatocytes indicated by intense magenta colour. (c,d)EAC inoculated group displaying weak Feulgen reaction indicating the depletion of DNA containing nuclei of hepatocytes, giant cell (arrow), pyknosis (arrowhead), (e) TQ pre-treated EAC inoculated animals illustrating strong Feulgen reaction indicating the increment of DNA containing hepatocyte nuclei (f) TQ post-treated EAC inoculated animals displaying moderate DNA content within hepatocytes’ nuclei as indicated by the moderate magenta red colour. (X 400).

3.5. Electron microscopic investigations:

In electron microscopic preparations, liver of control animals revealed normal hepatocytes with large nucleus that contains large amount euchromatin spread between heterochromatin that was dispersed near the margins of the nuclear cover and within the nuclei. A distinct spherical and centrally located nucleolus was also present. Numerous rounded and elongated mitochondrial profiles with well-developed transversal cristae were located. Well-designed rough endoplasmic reticulum carried ribosomes was noticed in the cytoplasm. Microvilli projected into the bile canaliculi lumen. Space of Disse was facing the blood sinusoids and contained microvilli. There were several types of healthy junctions. Endothelial and Kupffer cells were lined the sinusoidal walls Figure 3a.

Electron microscopic examination liver samples of Thymoquinone treated mice demonstrated similar features as those of the control group Figure 3b.

In contrast to the previous results, ultrastructural observations carried out on liver tissues of EAC inoculated animals revealed several morphological alterations like irregular envelope and there were nuclei were with evident chromatolysis. In other instances, pyknotic and apoptotic nuclei with irregular nuclear membrane were
observed. The presence of "microvilli" appeared clearly all over most the nuclei of this group. There were rough endoplasmic reticulum wrapping atrophied mitochondria.

Fat droplets occurred isolated in the cytoplasm. There were also excessive amount of red blood cells referring to congested blood vessels. Cell junctions were rarely found, and the space of Disse lost its microvilli. Also, constricted bile canaliculi lacking their microvilli were detected Figures 3c,3d. Certain tumor cells possessed elevated quantities of glycogen; others were having curved nuclei surrounding the lipid to some extent Figure 3g.

Ultrastructural micrographs for liver of animals pre-protected with TQ against Ehrlich inoculated animals showed remarkable improvement. The nuclei were almost alike those of normal mice. Mitochondria, rough and smooth endoplasmic reticulum in most instances were as normal as those of control outlines. Bile canaliculi raised in size and were completely established. On the other hand, blood sinusoids have the characteristic appearance of those of control counterpart as well as Kupffer cells Figure 3e.

Examination of animals post-treated with Thymoquinone displayed partial amelioration with few pathological alterations compared to the previous group. However, there were some shrinkage in some nuclei accompanied by irregular nuclear membrane. Kupffer cells were still large and activated. In few spots, the space of Disse was not fully formed. There were also some degenerated mitochondria with few or incompletely cristae and less abundant and dilated rough endoplasmic reticulum. In some foci, some junctions between cells were noticed clearly and karyorrhexis was obviously seen Figure 3f. All these features confirmed that the tumor still present and was not totally damaged. There was major variation in the shape of isolated tumors of this group. Certain apoptotic bodies that revealed membrane blebbing went through a process of cytoplasmic breakup. In addition, some foci displayed pyknotic or karyolitic nuclei. Cytoplasm contained malformed mitochondria, plentiful vacuoles and tiny profiles of endoplasmic reticulum were evident Figure 3h.
Fig. 3: Electron photomicrograph of liver mice (a,b) control and TQ treated groups respectively illustrating the presence of intact central and spherical nucleus (N), nucleolus (NU), rough endoplasmic reticulum (RER), numerous mitochondria (M), lysosome (Ly), microvilli (Mv), and endothelial cell (EC), and Kupffer cell. (c,d) Ehrlich carcinoma inoculated mice showing irregular shape and large nuclei (N) with nucleoli (NU), mitochondria (M), extensive rough endoplasmic reticulum (RER), space of Disse (SD), endothelial cell (EC), red blood cells (RBCs), pyknotic nuclei (PN), vacuole (V), fat droplets (FD) and Microvilli (arrow), collagen fibers (C) and Microvilli(arrow). (e) TQ pretreated EAC inoculated mice glycogen (G). (f) TQ post-treated EAC inoculated mice karyorrhexis (Kh), dilated rough endoplasmic reticulum (RER) wrapped the altered mitochondria (M), vacuole (V) and necrosis (arrows).(g,h) EAC solid tumor, EAC solid tumor post-treated with TQ, microvilli (arrow), fat droplet (FD), fragmented nuclei (*) (X, 2000,2500,1500,2500,5000,2000, 4000,4000) respectively.
3.6. Immunohistochemical observations:

Ki-67, PCNA immunoreactivity was contained in the nucleolus and nucleus. Its appearance was assessed according to the percentage of cells that were definitely stained by the antibody. In control and TQ treated animals liver section had limited number of Ki-67, PCNA positive cells, EAC group displayed enormous nuclear expression. Nevertheless, pre-EAC treated group indicated remarkable reduction in Ki-67 and PCNA labeling index compared to both EAC and/or TQ post-EAC treated groups as shown in Figures 4, 5, 6,7 and table 1.

Fig. 4: Photomicrograph of mice of Ki-67 immunostained liver (a,b) control and TQ treated group respectively displaying weak positive stained nuclei. (c,d) EAC inoculated group showing strong range of positive stained nuclei. (e) TQ pre-treated EAC inoculated group illustrated negative stained nuclei. (f) TQ post-treated EAC inoculated animals displaying weak positive reaction. (X 200).
Fig. 5: Photomicrograph of mice of PCNA immunostained liver (a,b) control and TQ treated group respectively displaying weak positive stained nuclei. (c,d) EAC inoculated group showing strong range of positive stained nuclei. (e) TQ pre-treated EAC inoculated group illustrated negative stained nuclei. (f) TQ post-treated EAC inoculated animals displaying weak positive reaction. (X 200).

Table 1: Liver Ki-67 and PCNA labeling index as a result of EAC and/or Thymoquinone treatment:

<table>
<thead>
<tr>
<th>Groups</th>
<th>Antigens</th>
<th>EAC</th>
<th>TQ</th>
<th>Control</th>
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<tbody>
<tr>
<td>EAC+TQ</td>
<td>23.17±0.54&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>9.17±0.48&lt;sup&gt;b&lt;/sup&gt;</td>
<td>72.83±0.83&lt;sup&gt;*&lt;/sup&gt;</td>
<td>2±0.37&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>TQ+EAC</td>
<td>27.33±0.56&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>9±0.37&lt;sup&gt;b&lt;/sup&gt;</td>
<td>78±0.63&lt;sup&gt;*&lt;/sup&gt;</td>
<td>2±0.37&lt;sup&gt;b&lt;/sup&gt;</td>
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Data are expressed as mean±SE, (a) significant compared to the control P<0.05, (b) significant compared to EAC, P<0.05.
4. Discussion:

This article was performed to assess the promising curative and protective influence of TQ on EAC cells inoculated animals. Toxicologically, the liver has a particular interest since hepatocytes exhibiting altered morphological and biochemical properties in response to exposure to hepatocarcinogens (Maronpot and others, 1987). Ehrlich ascites carcinoma (EAC) is considered the popular experimental destructive kind of tumor modeling which easily and rapidly develops after 14 days maximum, achieving great dimensions within few days (Carneiro and others, 2013; Portilho and others, 2011). Chemoprevention offers a tremendous hopeful approach for cancer prevention. Innovative chemo preventive substances of the natural source have been represented by vegetables and fruits to be studied owing to high constituents of bioactive compounds (Rafter, 2002) and their assorted pharmacological properties which are cytotoxic and chemo preventive effects against cancer (Dahiru and others, 2005; Gupta and others, 2004).

*Nigella sativa* seeds become a useful natural compound (Vihan and Panwar, 2006). Its remedial effects are attributed to Thymoquinone's (The master component of *Nigella sativa*) that is considered a hopeful natural product within *in-vivo* and *in-vitro* studies and numerous-cancer reports (Asaduzzaman Khan and others, 2017; Gali-Muhtasib and others, 2004; Hasan and others, 2013).
In the present study, EAC inoculated group exhibited a notable diminution in body weight gain compared to normal mice. This finding matches with that of (Badr El-Din and others, 2008; Miranda-Vilela and others, 2011). Body weight fall resulted from the decrease in growth caused by food intake drop and the tumor heaviness with its enormous development degree as well. The drop of tumor volume and body weight gain are the standard measures for assessing the efficiency of any drug against tumors (Miranda-Vilela and others, 2014). Liver tissues of TQ-treated rats showed the normal construction of the liver like that of the control animals demonstrating that TQ treatment did not cause any side effect or hepatotoxicity and this is in agreement with (Attalla F. El-kott and others, 2013; Mohamed and others, 2010). On the other hand, (Sayed-Ahmed and others, 2010) were in disagreement with who mentioned that liver of rats that administered TQ showed a mild degree of hepatic injuries.

EAC subcutaneous inoculated animals displayed increased necrosis, leucocytic infiltration, pleomorphic nuclei and cytoplasmic degeneration that are in agreement with earlier findings (Islam and others, 2013). Moreover, Leucocytic infiltration was believed to be a remarkable reaction of the body organs to stand up for any harmful influences (Sakr, 1999). Certain authors proposed that the cytoplasmic vacuolization is an outcome of marked disruption in the metabolism of fat and lipid. Shrinkage necrosis is an expression of programmed cell death in response to pathological changes (Wyllie and others, 1980). Tumors that comprised necrotic regions, included lysosomes and leucocytes which produces enzymes of elevated lysosomal hydrolase actions (Chen and others, 1996). TQ can maintain the standard performance of liver towards the aggressive action of EAC through the restraining of hepatic enzyme release.

Liver of ESC bearing mice presented lots of alterations caused by lysosomal enzymes and hydration (Morgan and others, 1960; Musa and others, 2004a). ESC cells infiltration probably attributed to tumor proliferation then invade the nearby organs (Chakraborty and others, 2007). Besides, EAC-injected mice revealed huge blocks of tumor pushing and transferred to the muscle tissue, these were in accordance with (Alshaymaa and others, 2012). Liver samples of EAC and TQ-injected group exhibited lack of classic hepatic architecture, these modifications likely as a result of cytoplasmic degeneration (Hashimoto and others, 1995), or perhaps due to mitochondrial degeneration. Necrosis usually includes clusters of nearby cells and causes permanent impairment of cell organelles, (Fukuda and others, 1993).

It is well known that the histological grade of tumor is an evaluation of the level of differentiation of a tumor and pointed to the tumor’s aggressiveness (Weigelt and Reis-Filho, 2009). Tumor histopathological and morphometric results indicated that TQ was effective in reducing tumor aggressiveness and in provoking tumor degradation. The presence of pleomorphic microvilli around the nuclei of most hepatocytes was considered to be a cancerous marker (Fukushima and others, 1981). Ultrastructural investigation of EAC inoculated animals revealed destruction and/or absent of microvilli in space of Disse and bile canaliculi. A decrease in microvilli count has been distinguished in hepatitis inoculated mice and after CCl4 administration (Reynolds, 1963; Sherlock and Dooley, 1997).

Fatty acids and other ingredients constituents of TQ are possibly responsible for the anti-tumor action of TQ and the mechanism was proposed by (Ando and others, 1970). Originally, the tumor cell membrane was ruined by fatty acid action (Das, 1989). (Morgan and others, 1960) stated that fatty acid length suppressed tumor progression in vitro. Comparable research was done on *Nigella sativa* seeds (Salomi and others, 1992). Our investigations reinforced that TQ possesses scavenging influence against oxidative stress induced through EAC. Modern studies discusses that ethanolic extract of *Nigella sativa* seeds has antitumor property(Musa and others, 2004b). On the other hand, the present data revealed that TQ extract administration improved liver injuries induced by EAC cells that is represented in light and electron microscopy micrographs and these results goes parallel to that obtained by (Fatma and others, 2011; Jafri and others, 2010; Li and others, 2010).

Some of the hepatic cells displayed reduction instead of swelling which is a distinct characteristic of apoptosis. Apoptosis is the programmed cell death that prompted by specific enzymes which destroyed DNA (Kumar and others, 2009). This typical phenomenon serves to eliminate undesirable cells during development (Fadeel and Orrenius, 2005). Beginning of apoptosis happens through signals from two separate pathways: the intrinsic or (the mitochondrial) pathway and the extrinsic or (death receptor–initiated) pathway. Both of them stimulate caspases, that are cell death mediators (Danial and Korsmeyer, 2004).

These results were supported by cytological examination that revealed a noteworthy rise in number of mitotic cells of positive control group. Conversely, the cytological examination of EAC cells in both preventive and protective mice, revealed a significant decrease in number of mitotic cells compared to positive normal one. Moreover, there was insignificant variance between preventive and protective mice in the appearance and morphology of ESC cells. This was in agreement with (Simoni and others, 2001)Arindam et al. (2003) who identified apoptotic features as well. Also (Pauloin and others, 2008) reviewed that during early apoptosis, cells presenting severe compressed chromatin, splitting nuclei, and conserved cytoplasmic organelles, conversely, in late apoptosis cells presented alike features, but lacking nuclear cover and most organelles. In addition to significant increase in necrotic spots, TQ treatment stimulated necrosis in the marginal regions of tumors. These regions are of penetrative neoplastic feature (Gali-Muhtasib and others, 2005; Kaseb and others, 2007; Roepke...
and others, 2007). TQ is probably slightly toxic to intact cells. Moreover, these results suggested that TQ stops tumor progression and could be considered as a cancer therapy prospective remedy candidate (Assaf and others, 2017; Mostofa and others, 2017). TQ pre-treated mice displayed a negligible pathological features to a great degree compared to that of control mice’s.

Feulgen reaction proposed 75 years ago is a kind of histochemical reactions commonly applied in biology. It is a good indicator of DNA state providing significant information about biological behavior of cells. DNA staining strength is relational to the DNA intensity which is a good indicator of liver cell apoptosis (Chieco and Derenzini, 1999). DNA content of control cells using Feulgen reaction appeared magenta red in color and sections examined after Ehrlich carcinoma cells inoculation showed a marked decrease in the DNA content. These findings were consistent with (Musa and others, 2004b). Many factors have been suggested to interfere with the reduction of DNA; one of the main factors is lysosomal particles and their contents of hydrolytic enzymes. Once the lysosomal membranes are disrupted under any pathological conditions, these enzymes become free in the cytoplasm causes obvious degradation of DNA and proteins. This concept confirms the findings of (Awasthi and others, 1984). Furthermore, (Musa and others, 2004b) investigated the effective action of Nigella sativa seeds in inhibiting the tumor cells propagation and reducing the mitotic index in the treated mice. These results confirmed that were achieved by (Salomi and others, 1992). Besides, liver samples revealed rearrangement of most diploid cells, lack of aneuploid cells with rise in proliferation index and an elevation in DNA content as well as. These outcomes probably owing to TQ and its ingredients’ antioxidant features. This study revealed that TQ could prevent the development of tumor and shows deletion in tumor progress.

Regarding to Ki-67, (Ito and others, 1999) stated that Ki-67 expression levels are important in assessing carcinogenicity of chemicals in rat, and its existence through different phases of the cell cycle considering it as a perfect indicator for neoplasia. Our investigations were matched with the results of (Pizem and others, 2001) who mentioned that Ki-67 and PCNA were valuable for proliferation assessment of hepatic cells. They showed a statistically considerable connection between Ki-67 and PCNA proliferative indexes in HCC in addition to affirmative connection between Ki-67 index and tumor stage. Besides, the investigation of (Stroescu and others, 2008) exhibited that immunostaining of HCC for Ki-67 was correlated with increased mitotic action.

**Conclusion:**

The outcomes of the current investigation are encouraging, and Thymoquinone has revealed a remarkable drop in cell propagation, DNA production, and elongation of lifespan of the animals with the Ehrlich ascites tumor. Overall, extra evaluation of Thymoquinone is strongly recommended within clinical and scientific fields to outline its possible value to be utilized as an innovative complementary remedy for cancer diseases.

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**Conflict of interest:**

All authors declare no conflict of interests.

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