INTRODUCTION

Cancer is a major health hazard in the world and claims over six million lives every year (Abdullaev, 2001). Cancer, a term comprising over 100 types of malignancy, is one of the major burdens of chronic disease in the world. It is very difficult to cure this disease primarily due to its multifactorial etiology. Early in the 20th century, only cancers small and localized enough to be completely removed by surgery were curable. Later, radiation was used after surgery to control small tumor growths that were not surgically removed. Finally, chemotherapy was added to destroy small tumor growths that had spread beyond the reach of the surgeon and radiotherapist [1,2]. Plant derived substances have recently become of great interest in the development of novel drugs [3]. The medicinal value of plants lies in the chemical substances that produce a definite physiological action on the human body. Many of the indigenous medicinal plants are used as spices [4]. With the aim of searching novel compounds without undesirable side effects, we focused on natural medicines. Plants are reported to have a long history in the treatment of cancer [5]. The use of plants and plant-based products for cancer treatment is rapidly growing in medical practices [6].

Scientific advancements have led to the exploration of many such plants chemically for its constituents and their impact in treating several types of diseases. As little attention has been devoted to understand the functional role of ethno-medicinally important plant proteins in relation to its therapeutic use and considering the potential advantage of using plant proteins as drugs (due to nontoxic nature), a thorough scientific research is attempted. Proteome analysis deals with determination of biological roles and functions of identified proteins [7].

This led us to choose the medicinal plant named as Tabernaemontana divaricata (Td) (Tamil-Nandhiyar vattam). This plant has been used in the traditional medicine of the Indian Ayurvedic system belonging to Apocynaceae family which is a small shrub, about 54 cm high, with large shiny leaves, crepe jasmine flowers, and appearing sporadically all year. It is a rich source of secondary metabolites with various pharmacological properties. It has been used in the folk medicine for anti-infection, antitumor, and antioxidative effect and the effect in neuronal activity (Pratchayasakul et al., 2008) [8]. The objective of this study is to evaluate the antitumor activity of protein extract being reported to validate the traditional use of the crude drug through in vivo evaluation. Our results suggest that whole plant extracts are promising anticancer agent.

METHODS

Collection and authentication of plant

The fresh leaves of Td (Apocynaceae) were collected from Coimbatore district, India. Taxonomic authentication was done by Dr.G.V.S. Murthy Taxonomist, TNAU, Coimbatore, Tamil Nadu, India, and the authentication number BSI/SRC/5/23/2015/Tech/2083.

Preparation of protein extract of Td leaf (TdPf)

Protein was extracted by recrystallization of ammonium sulfate. 20% fresh leaves of Td were taken and homogenized with phosphate-buffered saline buffer pH 7.2 and were centrifuged for 5000 rpm for 10 minutes. Pellets were discarded and supernatant was saved. To the supernatant add 10-100% ammonium sulfate and was centrifuged at 10,000 rpm, 4°C for 10 minutes. The supernatant was discarded and the pellet was suspended with dialysis membrane for salting out. The crude extract was kept at −20°C.
Chemicals
The chemicals and solvents used in the study were of the highest purity and analytical reagents grade. The chemicals were purchased from SD Fine Chem, Himedia and Sigma, India.

Experimental animals
About 70-day-old Swiss albino mice weighing 20±2 g were used for this study. The mice were procured from the Amala cancer Research Institute Kerala, India. The mice were divided into five groups (six mice per group) and maintained in polyacrylic cages at a temperature of 25±2°C, suitable humidity, dark/light cycle, with feed and water (ad-libitum). The mice were acclimatized to laboratory conditions for 7 days before the commencement of the experiment. The animal care and handling was done according to the regulations of Council Directive CPCSEA No: (IAEC.2015.BT:04) about good laboratory practice on animal experimentation. All animal experiments were performed in the laboratory according to the ethical guidelines suggested by the International Animal Ethics Committee.

Induction of lymphoma
Dalton’s lymphoma ascites (DLA) cells were obtained from Amala Cancer Research Institute, Kerala, India. The cells were maintained in vivo in Swiss albino mice by intraperitoneal transplantation of 1 × 10⁶ cells/mouse. The DLA cells aspirated from the peritoneal cavity of the mice were washed with saline and given intraperitoneally to the experimental animals to develop ascitic tumors.

Experimental design
A total of 30 Swiss albino mice were divided into five groups (n=6) and given food and water ad libitum. The vehicle and extract were administered intraperitoneal for 15 and 30 days. Group A served as control group; Group B mice induced with paraffin; Group C mice induced with silymarin, Group D mice induced with TdPf alone with effective concentration 50% value of 52 μg/kg body weight from the next day of induction for 15 and 60 days; Group E mice intraperitoneal administered with DLA and protein fraction of Td for 15 and 60 days. Group F received DLA cancer cells alone (i.e.) 10⁶ cells/ml. After the treatment period, six animals of each group were sacrificed by cervical dislocation to measure antitumor activity. The blood was collected from the animals by heart puncture. From the collected blood sample, the serum was separated and the liver marker enzyme was estimated from the animals by heart puncture. About 70-day-old Swiss albino mice weighing 20±2 g were used for this study. The mice were procured from the Amala cancer Research Institute, Kerala, India. The mice were divided into five groups (six mice per group) and maintained in polyacrylic cages at a temperature of 25±2°C, suitable humidity, dark/light cycle, with feed and water (ad-libitum). The mice were acclimatized to laboratory conditions for 7 days before the commencement of the experiment. The animal care and handling was done according to the regulations of Council Directive CPCSEA No: (IAEC.2015.BT:04) about good laboratory practice on animal experimentation. All animal experiments were performed in the laboratory according to the ethical guidelines suggested by the International Animal Ethics Committee.

Preparation of tissue homogenate
The liver was removed quickly and washed in ice cold isotonic saline. The liver was washed with saline and given intraperitoneally to the experimental animals to develop ascitic tumors. The liver was removed quickly and washed in ice cold isotonic saline. The liver was washed with saline and given intraperitoneally to the experimental animals to develop ascitic tumors. The liver was removed quickly and washed in ice cold isotonic saline. The liver was removed quickly and washed in ice cold isotonic saline. The liver was removed quickly and washed in ice cold isotonic saline. The liver was removed quickly and washed in ice cold isotonic saline. The liver was removed quickly and washed in ice cold isotonic saline. The liver was removed quickly and washed in ice cold isotonic saline.

Analytical methods

Estimation of the liver marker enzymes such as aspartate transaminase (SGOT), alanine transaminase (SGPT), and alkaline phosphatase (ALP) was carried out in the serum samples using Standard Kit method (Genei diagnostics).

Antioxidant assays
The liver homogenate was used to analyze the enzymatic antioxidant activities by superoxide dismutase (SOD), catalase (CAT), glutathione (GSH) peroxidase (GPx), and nonenzymatic antioxidants such as vitamin A, vitamin E, and reduced GSH were evaluated in the liver tissue homogenate using Standard Kit methods.

Histopathological study
The liver tissue samples collected were fixed in 10% formalin solution. After fixation, the tissue samples were embedded in paraffin and sections cut at 5 μm to later be stained with hematoxylin and eosin. The sections were then examined under light microscope and photographed (Standish et al, 2006).

Statistical analysis
The results obtained were reported as mean ± standard deviation. One-way and two-way analysis of variance was performed to analyze statistically significance of the data using Agres Statistical Package.

RESULTS
Estimation of liver marker enzymes in serum samples
Graph 1a-c shows the activities of marker enzymes in control and lymphoma bearing animals, respectively. In lymphoma bearing animals, the activities of the aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were significantly (69.40±2.48; 82.87±4.98) increased, whereas ALP were found to be significantly (56.51±3.81; 40.02±1.28; 8.97±1.33) increased when compared to the control mice (p<0.05). All the marker enzymes were found to be significantly increased in the serum of tumor bearing animals when compared to normal control mice. On TdPf treatment, the activities of all these enzymes were significantly brought back to near normal levels.

Estimation of lipid peroxidation
This study shows that a significant increase in lipid peroxidation (Graph 2) was observed in DLA induced mice.

Estimation of antioxidant activity

Enzymatic antioxidant
DLA cells induced mice showed a significant (p<0.05) decreased levels of SOD (0.83±0.03), CAT (1.93±0.03), and GPx (0.99±0.01). The treatment of TdPf alone at 5.2 μg/kg body weight showed a significant increased levels of SOD (1.74±0.46), CAT (1.94±0.03), and GPx (1.09±0.01) when compared to the normal mice group and also highly significant in 60 days of treated animals (p<0.05) (Table 1).

Estimation of nonenzymic antioxidants
Table 2 exhibits the levels of nonenzymatic antioxidants in the liver of control and experimental animals. From the Table 2, it is found that the level of vitamin A, vitamin E, and GSH was significantly reverted back to near normal level when treated with the plant extract when compared to Group III treated (standard antioxidant).

Histopathological observations in the liver of TdPf treated mice were comparable to that of controls and silymarin without any structural changes in the liver morphology.

Plate 1a-j indicates the histopathological observation of DLA bearing mice, (a) Untreated; (b) paraffin oil 15 days observation; (c) silymarin (standard drug); (d) protein extract (TdPf); (e) protein extract + DLA; (f) paraffin oil 60 days observation; (g) silymarin 60 days observation; (h) plant extract 60 days; (i) plant extract + DLA 60 days; (j) DLA 15 days observation.

Hepatic cells of DLA induced mice showed large confluent areas of hepato cellular necrosis with peripheral rim of surviving cells with focal steatosis and balloon degeneration. This could be due to the formation of highly reactive radicals because of oxidative threat caused by DLA.
However, the treatment with TdPf exerted a significant inhibition of metastasis in the liver indicating their antimetastatic activity which could be comparable to that of silymarin, the standard drug used for comparison. However, TdPf was found less efficient in preventing metastasis at abdominal muscle when compared to silymarin (Plate 1c, d, h and i).

**DISCUSSION**

Many natural products have served as anticancer agents in the treatment and also as lead compounds for further research. Many plants are used to treat tumors in the Indian traditional system of medicine, but most of the plants have not been scientifically evaluated (Pushpangadan and Subramonian, 1998). Hence, enormous scope exists for identifying potent anticancer plants. Based on our earlier studies, this work was aimed to evaluate the TdPf for *in vivo* antitumor properties. ALT catalyses the conversion of alanine to pyruvate and glutamate and is released in a similar manner. Elevated levels of serum enzymes are indicative of cellular leakage and loss of functional integrity of cell membrane in the liver [9]. A similar significant decrease in AST, ALT, and ALP was observed by Santhi and Annapoorani [10] by the administration of silymarin to carbon tetrachloride (CCl₄) induced hepatotoxicity in mice. Heba et al. [11] also reported that rats fed with powder of *Ziziphus spina-christi* fruit decrease the activities of AST, ALT, and ALP in CCl₄ induced rats in a dose-dependent manner. A similar significant increase in the activity of AST in the serum of ELA induced mice was reported by Nalini et al. [12].

Elevated lipid peroxidation and a poor antioxidant system have been reported in cancer patients. Altered activities of enzymatic antioxidants are reported during carcinogenesis or after tumor formation. Hence, the elevated lipid peroxidation in the circulation of cancer animals is due to a poor antioxidant defense mechanism. A decrease in SOD and CAT activities described in tumors is regarded as a marker of malignant transformation. Lowered activities of SOD and CAT were reported in several cancers [14].

Excessive generation of reactive oxygen species has been considered as hallmark in several cancers, including Dalton’s ascitic tumor model [15]. Antioxidant enzymes that scavenge intermediates of oxygen reduction provide a primary defense against free radicals in situ. It is well known that SOD, CAT, and GPx also play important roles as protective enzymes against LPO in tissues [16]. Several investigators reported that reduced activities of SOD, CAT, and GPx in tum or-bearing animals may be due to a down-regulation of SOD and CAT genes induced by certain hormonal factors or ROS themselves (Quan et al., 2011).

Antioxidants are important substances with the ability to protect the body from damage caused by free radical-induced oxidative stress. A variety of free radical scavenging antioxidants exists within the body. Vitamin A has been associated with a decreased risk of human cancer and has protective effects in animal models of carcinogenesis. Vitamin A is known to be an important natural antioxidant capable of countereacting oxygen free radicals and exerting a protective antioxidant...
effect [17]. Santhi et al. [18] reported that Cynodon dactylon leaf protein increases the level of vitamin A in ELA induced mice. Parks and Traber [19] reported that vitamin E is one of the most important free radical scavenging chain breaking antioxidant within the biomembrane. A significant increased level of vitamin A and E in the liver and kidney of rats induced with ammonium metavanadate toxicity by the prior treatment with green tea was reported by Soussi et al. [20]. The GSH is a powerful nucleophile, critical for cellular protection such as detoxification from reactive oxygen species, conjugation and excretion of toxic molecules, and control of the inflammatory cytokine cascade [21]. Bigoniya and Rana [22] reported that saponin fraction of Euphorbia neriifolia has increased the levels of hepatic GSH in CCl₄ induced hepatotoxicity in rats.

**CONCLUSION**

This study was conducted to evaluate the antitumor activity of TdPf offered protective effect against DLA tumor by their in vivo antioxidant and antitumorigenic potential. The extract treatment at the dose of 52 µg/kg inhibited the tumor activity by the serum biochemical
assessment. The extract also restored the hepatic lipid peroxidation and free radical scavenging enzyme GSH as well as other antioxidant enzymes such as SOD, CAT, and GPx in tumor-bearing mice to near normal levels. Our results suggest that whole plant extracts are promising anticancer reagents.

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REFERENCES