INTRODUCTION

Methotrexate (MTX) is a chemotherapeutic agent widely used for treating cancer, psoriasis, dermatomyositis, sarcoidosis, and rheumatoid arthritis [1]. MTX mainly interferes with the growth rate of highly reproducing cells such as cancer cells, bone marrow cells, etc. However, the drug efficiency of MTX is limited and thus chronic high-dosage administration leads to hematopoietic suppression, hepatotoxicity including cellular necrosis and cirrhosis of the liver [2-4]. Studies recommend that MTX-induced liver toxicity is associated with high production of ROS (reactive oxygen species) and failure of antioxidant defense system due to reduction in glutathione (GSH) supply [5]. High concentration of ROS, in turn, enhances the lipid peroxidation level as well unsaturated fatty acid accumulation in cell membranes.

Asparagus racemosus of family Liliaceae is a prevalent consuming vegetable plant grown in almost all fractions of the continents Asia, Australia, and Africa. The root of this plant possesses certain biological properties such as anti-diabetic, anti-arthritis, anti-cancer, anti-fungal, etc [6,7]. This plant has already been used as a remedy for diseases such as diarrhea, dysentery, and nervous breakdown [8]. As far as concern, very less information is available on the hepatoprotective role of A. racemosus on the induced liver injury. The utilization of herbal extracts has cure for many diseases without any toxic side effects and thus have a high success rate. Hence, to evaluate the efficiency of the protective role of A. racemosus, the present study has been conducted where an aqueous extract of the plant is administered to female Wistar albino rats with MTX-induced hepatic damage.

METHODS

Chemicals

The tablets of MTX were commercially acquired from the pharmacy and were used by dissolving in normal saline solution. All the chemicals and reagents used for conducting the experiments were analytically graded.

Extract preparation from plant root

The roots of the medicinal herb A. racemosus were procured from the localities of Vellore around August, 2017 and authentication done from Horticultural Research Station, Yercaud, Tamil Nadu, India. The fresh roots collected were cleaned up well and shade dried for a week at 45°C. The thoroughly dried roots were powdered finely by multiple grinding and was mixed in water. The mixture was kept at room temperature for overnight; and on next day, the mixture is filtered separately using filter paper for collecting the root extract. The extract solution is then dried completely at 45°C for a time span of 4–5 h in a hot air oven. The dried extract was then collected in vials and stored in 4°C for the present experimental study. The rats were kept in normal room temperature (24±2°C) and contained 12 h:12 h light and dark cycles. They were provided with usual laboratory diet.

Preparation of animals for experiment

A total of 24 female Wistar albino rats with weights differing in the span of 150–200 g resorted for the present experimental study. The rats were kept in normal room temperature (24±2°C) and contained 12 h:12 h light and dark cycles. They were provided with usual laboratory diet. The approval for the experimental protocol was given by the Animal Ethical Committee, (VIT/IAEC/14Nov/08) VIT, Vellore.

Experimental design

The animals were eratically allocated into four groups with six rats in each group. The total time span for the study was 14 days. Group I containing positive control animals were tended with normal saline solution orally for entire 14 days of study. The animals in Group II and IV, the levels were restored near to control, which supported the protective role of A. racemosus against MTX-induced hepatic damage.

ABSTRACT

Objectives: Various clinically available drugs along with the beneficial action also have drastic side effects due to chronic exposure. In liver, these resulting side effects can be available production of reactive oxygen species, which will further lead to oxidative stress and hepatotoxicity. Therefore, as a preventive measure, the protective role of herbal extracts is being evaluated due to its high success rate and low toxic effects. The primary aim of this study was to evaluate the efficiency of the protective role of Asparagus racemosus is evaluated and studied against methotrexate (MTX)-induced hepatic damage in male Wistar albino rats.

Methods: The course of the study was for 14 days. During this experimental study, the animals were categorized into four groups with six rats per group. Group I (positive control) which was treated with normal saline, Group II (negative control) with MTX 20 mg/kg of body weight on 12th day, Group III with A. racemosus 300 mg/kg of body weight + MTX 20 mg/kg on 12th day, and Group IV with A. racemosus 100 mg/kg of body weight + MTX 20 mg/kg on 12th day. On 14th day, the animals were sacrificed, and histopathological as well as antioxidant assays were performed.

Results and Conclusion: Assays revealed high lipid peroxidation level and low antioxidant levels in Group II. Meanwhile, in Group III and IV, the levels were restored near to control, which supported the protective role of A. racemosus against MTX-induced hepatic damage. Histopathology evaluation also supported the above-mentioned findings.

Keywords: Hepatotoxicity, Lipid peroxidation, Methotrexate, Reactive oxygen species, Asparagus racemosus.

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it served as the negative control. The MTX was given to the animals in the mode of intraperitoneal injection. A low dosage of the aqueous root extract of *A. racemosus* with the concentration of 100 mg/kg of body weight was given orally to the Group III animals. Group IV animals were considered as high dosage group and were administered with aqueous root extract of *A. racemosus* with the concentration of 300 mg/kg of body weight. On 12th day, Group II and Group IV were injected with MTX intraperitoneally in the concentration of 20 mg/kg of body weight. The dosages were provided to animals based on Devi et al.

The body weights of the animals were estimated on the first and last days of the experimental studies. On 15th day, 24 h later to last dosage, the animals were sacrificed; and in vials, the blood was collected through heart puncture. Later, the liver tissue sample was carefully excised from the animal and blotted dried after washing in normal saline solution to clean off the stains of blood. The dried liver samples were then weighed and homogenized for carrying out the histopathological and other in vivo antioxidant assays.

**Estimation of biochemical parameters**

The blood samples collected in vials from animals were allowed to develop clots in normal environmental conditions for few minutes and later centrifuged to separate out the serum from blood clots at 3000 rpm for 10 min at a temperature of 4°C. The serum thus separated was used in evaluating the levels of various marker enzymes alkaline phosphatase (ALP) and alkaline transferase (ALT) in the liver. It was carried out with the help of commercially available kits for estimating the marker enzymes from Autospan Diagnostics Pvt. Ltd., India. The total protein content in the blood serum was calculated based on the protocol by Lowry et al. (1951).

**Homogenate solution preparation**

Liver homogenate solution of concentration 10% was prepared using 0.1M ice-cold phosphate buffer with pH 7.4 and 1:17% KCl. A part of the homogenate thus obtained was used in estimating lipid peroxidation and reduced GSH status. The remaining part of homogenate solution was subjected to centrifugation at 4°C for 10 min at 10,000 rpm. The supernatant solution separated out was utilized in conducting antioxidant enzymatic assays as follows: Catalase (CAT) and superoxide dismutase (SOD).

**Assessment of activity of GSH peroxidase (GPx)**

With the help of electron donor, GSH, GPx prevents free radical formation by reducing peroxyl radicals. GPx assay was carried out based on the protocol developed by Rotruck et al. To 0.4 mL phosphate buffer with pH 7.0 and 0.4M, 0.1 mL of homogenate supernatant was added along with 0.1 mL sodium azide, 2.5 mM H$_2$O$_2$, 0.2 mL of 4 mM GSH, and 0.2 mL distilled water. 10% TCA was added to the mixture after incubating for 10 min at room temperature, followed by centrifugation for 10 min at 1500 rpm. From supernatant thus obtained, 1 mL was mixed with 0.5 mL of DTNB (Ellman’s Reagent) and about 4 mL of 0.3M disodium hydrogen phosphate buffer. The absorbance value for the resulting mixture was taken at 412 nm, and the results were expressed as µg of GSH utilized/min mg protein.

**Assessment of lipid peroxidation**

Estimating the quantity of end product of lipid peroxidation, malondialdehyde (MDA), can detect the extent of oxidative stress happened quantitatively. Hence, it acts as the biomarker in estimating the cell oxidative stress levels. Lipid peroxidation estimation was carried out according to the protocol of Ohkawa et al. (1971). To 1 mL of tissue homogenate, 1 mL TCA (10%) was added at incubated for 10 min at room temperature, followed with centrifugation at 4°C for 15 min at 2500 rpm. To 1 mL of TBA (0.67%), 1 mL of supernatant was added and kept in water bath for 20 min. The mixture tubes were then cooked under tap water and later 1 mL of distilled water and 5 mL n-butanol: pyridine was added separately to each tube. The mixture was again centrifuged and the butanol layer formed on the top was collected, and the absorbance reading was taken at 532 nm. The result calculated was denoted as n moles MDA formed/mg protein.

**Assessment of SOD**

The enzyme SOD acts as an antagonist for free radicals which are highly reactive like hydroxyl radicals by converting it into oxygen and H$_2$O$_2$. The protocol by Marklund and Marklund et al. (1974) for auto-oxidation method of pyrogallol was used to determine enzymatic activity. To 0.5 mL of tissue homogenate supernatant, 0.15 mL chloroform and 0.25 mL absolute ethanol were added, and kept in a shaker for 15 min followed by centrifugation. 2 mL Tris HCL with pH value 8.2 and 0.1M was added to 0.5 mL of supernatant acquired along with 1 mL of distilled water and 0.5 mL solution of pyrogallol. The absorbance reading for the final mixture is taken at 420 nm with 1 min interval for a total of 3 min. The results were expressed as U/mg protein.

**Assessment of CAT activity**

CAT enzyme helps in the conversion of H$_2$O$_2$ developed during toxic reactions inside the body, to water, and oxygen. To 0.1 mL of tissue homogenate, 1.9 mL of phosphate buffer along with 1 mL of H$_2$O$_2$ was added. The protocol followed was based on the method of Beers and Sizer et al. (1952). The readings of the absorbance were taken at 240 nm for 3 min with an interval of 1 min. The results of enzyme activity were expressed in mM H$_2$O$_2$ decomposed/min/mg protein.

**Histopathological evaluation**

The liver samples resected from the animals of each group were stored in a neutral formalin buffer (10%). Using an automated tissue processor, the liver tissue samples were processed and later fixed in wax. With the help of Leica microtome RM 2155, the tissue sections were sliced on a thickness of 5 µm and later stained with Eosin and Hematoxylin (E and H). The slides after staining were visualized under a light microscope.

**Statistical evaluation**

The result data were expressed in mean ± standard error mean, followed by the statistical analysis using ANOVA and Dunnett’s test. Statistical significance was considered at p<0.05. Statistical significance studies were executed using Graphpad Instat Software Inc., of version 3.06, San Diego, USA.

**RESULTS**

**Body and organ weight**

The variations between initial and final body weight as well as changes in liver weight are shown below in Table 1. In Group II, the animals body weight got increased a little bit due to the treatment of MTX compared to the normal controls of Group I whereas in both Group III and Group IV due to the restoration action of aqueous root extract of *A. racemosus*, the body weight is more or less similar to Group I. The liver weight in Group II, Group III, and Group IV did not show any significant difference. Hence, it can be concluded that pretreated with aqueous root extract of *A. racemosus* has no effect on organ weight.

**Biochemical estimation**

The levels of liver marker enzymes and serum total protein are depicted in Figs. 1-3. In Group II which was treated with MTX, the levels of liver marker enzymes ALT (Fig. 1) and ALP (Fig. 2) were increased drastically (p<0.05) compared to the normal control of Group I. This elevation depicts the presence of liver damage caused by the induced MTX toxicity. *A. racemosus* treated Group III and IV before MTX treatment showed a decrease in the levels ALT (Fig. 1) and ALP (Fig. 2) marker enzymes, which indicate that the aqueous root extract of *A. racemosus* has counteracted the toxicity action of MTX and restored back the level to normal.

The levels of serum total protein (Fig. 3) also decreased drastically in MTX administered Group II whereas the serum protein levels were restored close to normal in Group III and IV due to the administration of aqueous root extract of *A. racemosus* when compared with the normal control Group I.

**Antioxidant assays**

Table 2 shows the variations in levels of lipid peroxidation, antioxidant enzymes as follows: SOD, CAT, and GSH peroxidizes (GPx).
Group II clearly showed the elevated levels of lipid peroxidation due to the action of MTX administered whereas the levels got reduced in Group III and IV due to the pretreatment with *A. racemosus* when compared to the positive control Group I. Both Group III and IV gave results close to the Group I, which suggests the restoration role of aqueous root extract of *A. racemosus*.

The antioxidant SOD, CAT, and GPx enzyme levels significantly got reduced in MTX pretreated Group II when compared to normal control Group I. This shows the toxic action of MTX which resulted in the impaired action of antioxidant enzymes due to antioxidant depletion. In Group III and IV, the antioxidant enzyme levels were increased near to the normal level when compared to control Group I. This suggests the protective role of the aqueous extract of *A. racemosus* against the induced toxicity.

**Histopathological Examinations**

Fig. 4a depicts the normal control Group I which showed normal histology. Fig. 4b depicts the MTX pretreated Group II which showed damaged histology due to dilated and congested central veins as well as increase in the number of Kupffer cell in the sinusoids. It also showed chronic inflammatory cell infiltration, increase in eosinophil, and destroyed periportal hepatocytes. Fig. 4c depicts the *A. racemosus* extract treated Group III where more normal hepatocytes and few hepatocytes show necrosis were observed. Similarly, Fig. 4d of *A. racemosus* treated Group IV also showed minimal inflammatory cell infiltration in the portal tract.

**DISCUSSION**

The liver is the main xenobiotic metabolism site, and hence, increased the amount of toxicants hinder the normal function of the liver by reducing the antioxidant reserve [9]. One of the factors which increase the level of toxicity is the chronic overdose of the drug. Chemotherapeutic drugs are considered to have acute side effects in different organ systems. Studies revealed that chronic overdose intake of MTX results in liver injury [10,11]. MTX inhibits the cytoplasmic NADP dehydrogenases and NADP malic enzymes which are required to maintain the reduced state of GSH reductase [12,13]. Reduction in cellular GSH, in turn, reduces the effectiveness of antioxidant defense of the liver against the increased ROS and thereby leading to hepatotoxicity [14,15].

The interpreted data from this study suggest that the levels of lipid peroxidation were increased significantly whereas other antioxidant enzyme levels including GSH were decreased in the negative control group compared to the positive control. The levels ALT and ALP, which are the markers of hepatic injury, were also elevated in the negative group. The reduced levels of these antioxidant enzymes were restored back again, and the ALT and ALP levels were normalized in groups.

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**Table 1: The activity of methotrexate on body weight and organ weight of the experimental groups of animals with or without the treatment of aqueous root extract of *Asparagus racemosus***

| Groups                  | Initial body weight (mg) | Final body weight (mg) | Percentage increased (%) | Absolute organ weight (g) |
|-------------------------|--------------------------|------------------------|--------------------------|--------------------------|
| Group I (control)       | 178.3±3.28               | 190.66±2.96            | 6.47±0.27                | 0.17±0.0014              |
| Group II (negative)     | 230±16.64                | 242.66±15.98           | 5.29±0.62                | 0.176±0.0048             |
| Group III (low dosage)  | 172.6±5.20               | 183.33±4.91            | 5.83±0.32                | 0.174±0.0053             |
| Group IV (medium dosage)| 192.3±19.42              | 202.66±20.53           | 5.08±0.21                | 0.170±0.0006             |

The result values are represented as mean ± standard error mean; n=6 in each group; *p<0.01 and **p<0.05 considered statistically significant. Comparisons are as follows: (a) Group II when compared with normal group. (b) Group III and IV when compared with negative control Group II. Statistical analysis using ANOVA, followed by Dunnett’s comparison test. *A. racemosus: Asparagus racemosus*
Table 2: Action of MTX on the antioxidant enzyme parameters and lipid peroxidation in the liver organs of experimental animals with or without A. racemosus treatment

| Parameters                          | Group I (control) | Group II (negative) | Group III (low dosage) | Group IV (high dosage) |
|-------------------------------------|-------------------|---------------------|------------------------|------------------------|
| Lipid peroxidation (n moles/mg protein) | 0.04±0.005        | 0.072±0.0007a**     | 0.040±0.006b**         | 0.051±0.0036b**        |
| CAT (j mol H2O2 consumed/min mg protein) | 0.13±0.0008b      | 0.115±0.0007a*      | 0.103±0.0066b*         | 0.127±0.004b*          |
| SOD (U/mg protein)                  | 3.83±0.067        | 1.81±0.307a*        | 3.31±0.224b**          | 3.023±0.342b**         |
| GSH peroxidase (µg of GSH used/min/mg protein) | 23.991±0.090      | 14.87±1.070a*       | 20.76±1.15b**          | 21.88±1.17b**          |

The result values are represented as mean ± standard error with n=6 in every group. **p<0.01 and ***p<0.05 considered statistically significant. Comparisons are as follows: a - Group II when compared with normal group. b - Group III and IV when compared with negative control Group II. Statistical analysis using ANOVA, followed by Dunnett’s comparison test. GSH: Glutathione, A. racemosus: Asparagus racemosus, CAT: Catalase, SOD: Superoxide dismutase

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REFERENCES

1. Bayram M, Ozogul C, Dursun A, Ercan ZS, Isik I, Dilekko E. Light and electron microscope examination of the effects of methotrexate on the endosalpinx. Eur J Obstet Gynecol Reprod Bio 2005;120:96-103.
2. Tobias H, Auerbach R. Hepatotoxicity of long term methotrexate therapy for psoriasis. Arch Intern Med 1973;132:359-61.
3. Hytiroglou P, Tobias H, Saxena R, Abramiou M, Papadimitriou C, Theise ND, et al. The canals of hering might represent a target of methotrexate hepatic toxicity. Am J Clin Pathol 2004;121:324-9.
4. Barak AJ, Tuma DJ, Beckenhauer HC. Methotrexate hepatotoxicity. J Am Coll Nutr 1984;3:93-6.
5. Cetiner M, Sener G, Sehirli AO, Demiralp EE, Erkan F, Sirvanci S, et al. Taurine protects against methotrexate-induced toxicity and inhibits leukocyte death. Toxicol Appl Pharmacol 2005;209:39-50.
6. Edenharder R. Antinutritagenic activity of vegetable and fruit extracts against in vitro benzo(a)pyrene. Z Gesamte Hyg 1990;36:144-8.
7. Shimooyamada M, Suzuki M, Sonta H, Maruyama M, Okubo K. Antifungal activity of the saponin fraction obtained from asparagus and its active principle. Agric Biol Chem 1990;54:2553-7.
8. Nadkarni AK. India Materia Medica. Bombay: Popular Prakashan; 1976. p. 151-5.
9. Asha DS, Blossom B. Hepatoprotective role of Spirulina fusiiformis on methotrexate induced liver injury in Wistar rats. Res J Biotechnol 2016;11:58-64.
10. Sener G, Demiralp EE, Cetiner M, Erkan F, Yegen BC. β-glucan ameliorates methotrexate-induced oxidative organ injury via its antioxidant and immunomodulatory effects. Eur J Pharmacol 2006b;542:170-8.
11. Urzu S, Tahan V, Aygun C, Eren F, Unluoguzel G, Yokuş M, et al. Role of ursodeoxycholic acid in prevention of methotrexate-induced liver damage. Dig Dis Sci 2008;53:1071-7.
12. Babiai RM, Campello AP, Carnieri EG, Oliveira MB. Methotrexate: Pentose cycle and oxidative stress. Cell Biochem Funct 1998;16:283-93.
13. Babar AJ, Tuma DJ, Beckenhauer HC. Methotrexate hepatotoxicity. J Am Coll Nutr 1984;3:93-6.
14. Sabina EP, Samuel J, Ramya SR, Patel S, Mandal N, Pranatharthiharan P, et al. Hepatoprotective and antioxidant potential of Spirulina fusiiformis on acetaminophen induced hepatotoxicity in mice. Int J Integr Biol 2009;6:1-5.
15. Kumar RS, Daniel JA, Jayachristy SA, Devi SA. Hepatoprotective role of Abutilon indicum on Lead induced liver injury in Wistar rats. Int J Pharm 2016;10:36-9.
16. Om FR, Kumar MR, Mani TT, Niyam KS, Kumar BS, Phaneendra P, et al. Hepatoprotective activity of “Asparagus racemosus root” on liver damage caused by paracetamol in rats. Indian J Novel Drug Deliv 2011;3:112-7.
17. Suchismita R, Shrabani P, Shreya ML, Koushik D, Arpita P, Animesh S, et al. Phytochemical analysis, antimicrobial activity and assessment of potential compounds by thin layer chromatography of ethanol fraction of Asparagus racemosus roots. Int J Pharm Pharm Sci 2014;6:367-70.
18. Rachana G, Bhavna S, Nilesh G, Ghanshyam P, Bhavna P. Isolation and characterization of Shatavarin IV from root of Asparagus racemosus wild. Int J Pharm Pharm Sci 2015;7:362-5.