Antioxidant and immunomodulatory effect of AKSS16-LIV01 – a multi herbal formulation against ethanol induced liver dysfunction in mice

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Abstract

Background: Liver complication arises commonly due to high alcohol consumption rate. Majority of the people residing in both developed and under developed countries consuming alcohol face various liver complications such as liver fibrosis, fatty liver, liver cirrhosis and even hepatocellular carcinoma. Invention of safe and symptomatic medication to overcome this situation is a new challenge worldwide. The main objective of the study is to deliver a safe and symptomatic medication to reduce the ethanol induced liver dysfunction.

Methods: In this study we have developed a multi herbal formulation (AKSS-16-LIV01) which minimised liver damage against various toxicants. Swiss albino mice were divided into seven groups where ethanol induced damage was observed for weeks followed by sanative response observation by our herbal formulation. The groups are normal control group, ethanol treated group (50% v/v), AKSS16-LIV01 low dose (75 mg/kg/day) pre-treated group, AKSS16-LIV01 middle dose (150 mg/kg/day) pre-treated group, AKSS16-LIV01 high dose (300 mg/kg/day) pre-treated group, Sylimarin pre-treated group (100 mg/kg/day) and only AKSS16-LIV01 (300 mg/kg/day) treated group.

Results: The results potrayed significant elevation of various biochemical parameters, lipid profile parameters, lipid peroxidation, nitric oxide (NO) concentration, nitric oxide synthase level and pro inflammatory cytokines level i.e. tumor necrosis factor (TNF-α) and transforming growth factor (TGF-β1) in the ethanol induced mice. On the other hand serum total protein, total albumin, albumin globulin ratio and level of tissue antioxidant enzymes activity (SOD, CAT, GSH and GPx) were significantly reduced by ethanol. Dose depended therapeutic application of the formulation (AKSS16-LIV01) significantly suppressed all the relevant above parameters and protected the liver from ethanol induced fibrogenesis. Apart from this gross morphology of the liver, H&E liver histology and massontrichrome&serius red examination of the liver section strongly supported the hepatoprotive effect of the formulation as compared with standard drug Sylimarin. Result of the study implies that developed multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg/day gave the best optimum response to reduce the ethanol intoxication.

Conclusion: Result clearly depict that AKSS16-LIV01 may be a safe and nontoxic medication which protect the liver against ethanol induced oxidative injury and maintained pro inflammatory cytokines level in the future.

Keywords: Multi herbal formulation, Liver damage, Liver function test, Lipid profile, Oxidative stress, Liver histology

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Introduction

Our body continuously exposed with various harmful toxicants as a result liver detoxifying the toxicant and maintains cellular homeostasis [1, 2]. People in both developed and underdeveloped countries when consume excesses alcohol or taken alcohol on a continuous basis leading to alcoholic liver disease (ALD). Individuals those are suffering with ALD facing lots of liver complications such as fatty liver disease, hepatic fibrosis, hepatic cirrhosis and hepatic cellular carcinoma even death [3, 4]. World health organization (WHO) published a report that stated death due to the liver failure hepatic cirrhosis and hepatic cellular carcinoma even death [3, 4]. Scientific study stated that consumption of ethanol elevate the ratio of NADH/NAD⁺ in the liver cell which create disruption of oxidation of fatty acids in mitochondria developed liver dysfunction [6, 7]. Other causes of liver disease deposition of lipid molecule in the liver cells which stop the normal hepatic functions. Alcohol enhances the transportation process of lipids towards the liver from the small intestine which elevate the fatty acids mobilization from adipose tissue, taken up by the liver [8]. This causes damage of the liver cell membrane which releases the transaminases enzymes (AST and ALT) in the blood stream. On the other hand this damaged cell membrane also release alkaline phosphatase (ALP) and gamma glutamyltransferase (GGT) which indicate hepatic damage and inhibit the intercellular homeostasis [9–12].

Superoxide (O₂⁻), hydroxyl radical (·OH), hydroxyl ion (OH⁻) and hydrogen peroxide (H₂O₂) are the common reactive oxygen species (ROS) were generated in the liver cell [13] when exposed with certain chemicals, environmental pollutants, xenobiotics etc. Firstly, these free radicals are generated from the oxidation procedure within the cell developed oxidative stress which destabilizes the normal cellular homeostasis [14, 15]. Secondly, Chronic intake of alcohol generates ROS via cytochrome P450 2E1 in the liver cell producing DNA damage, loss of membrane integrity, amino acid oxidation and inactivation of specific enzymes through oxidation of their cofactors [16, 17]. Important antioxidant enzymes like super oxide dismutase (SOD), catalase (CAT), glutathione (GSH) and glutathione peroxidase (GPx) mainly converts these harmful radicals through a series of biochemical reaction into oxygen and water molecule and protect the hepatic cell from the oxidative stress injury. It is evident that excessive intake of alcohol may develop cellular oxidative stress; indirectly disrupting cellular antioxidant defense system, which eventually produces cell death (apoptosis) and tissue damage. On the other hand nitric oxide (NO) is an important mediator of many physiological functions responsible for pathogenesis of many diseases. Chronic alcohol consumption increases nitric oxide (NO) levels which may lead to toxicity by peroxynitrite and destroy the membranous integrity. Free radicals i.e. reactive oxygen species (ROS) and reactive nitrogen species (RNS) when generates excessively in cell may not be neutralized in the system and cannot be eliminated from the body [18].

From the ancient time people from various countries depends on traditional system of medicine for curing the diseases as a safe and symptomatic medications. In the modern world Indian medicinal herbs and medicinal spices have been extensively used as an alternative medicine because of their promising medicinal property and lesser side effects in comparison to the allopathic drugs [19]. These medicinal herbs and spices are enriched with various constituents likes polyphenols, flavonoids, alkaloids, glycosides, tannins, proteins, amino acids, saponin etc., which play a key role in cellular protection from the toxicants. Herbal formulations composed of some medicinally sound plants, now have garnered greater interest throughout the world due to its synergistic action. Recently people have a greater interest in herbal medicines because of their lesser side effect in clinical experience, pronounced effectiveness, safe for long term use and relatively low cost [20–22].

Currently throughout the globe fight against various liver dysfunctions such as fatty liver, liver fibrosis, liver cirrhosis etc. through symptomatic and safe medication is a new challenge. Presently, there is no effective treatment for hepatic dysfunctions. These herbs are natural resources of antioxidants that serve as the first line of defense against free radical damage and are considered to be important in maintaining optimum health and hygiene. Our previous reports stated that the formulation have no adverse side effect and no toxicity in mice [23, 24]. So, in the present study we formulated a new novel phytomedicine (AKSS16-LIV01) composed of six indigenous medicinal herbs and three medicinal spices those were mentioned in Ayurveda. Commercially available formulations composed of more than 12 medicinal herbs without medicinal spices. On the other hand in composition of the formulation the concentration of the individual herbs is less in comparison to marketed products. So, the developed formulation is an unique one and produced better therapeutic effects on animal models. Therefore, the present study was undertaken to evaluate the ethanol induced oxidative stress and hepatic injury of adult mice and its deleterious recovery by the application of unique novel multi herbal formulation (AKSS16-LIV01). The results of this study are expected to supply a transparent picture about the role of our newly formulated AKSS16-LIV01 in ethanol-induced hepatic damage, and should shed light on an achievable ethno-botany driven solution for serious liver problems.
Materials and methods

Chemicals
Trichloroacetic acid (TCA), thiobarbituric acid (TBA), ethanol, TRIS buffers were purchased from SISCO laboratories, India. PBS buffer (pH 7.2) was taken from Sigma-Aldrich, Germany. All the biochemical kits (ALT, AST, GGT, ALP, Cholesterol, Triglyceride, Phospholipid, HDL, LDL etc.) were procured from Span Diagnostics, Surat, India. Hydroxyproline assay kit was procured from Bio Vision, Milpitas, CA. Antioxidant kits (SOD, CAT, GSH and GPx) were obtained from Boehringer, USA. ELISA kit TNF-α and TGF-β₁ were procured from Sigma Aldrich, USA. All others reagents utilized in this study are laboratory grade.

Collection and authentication of the herbs
In this study all the required medicinal herbs and medicinal species used for the preparation of novel formulation were collected from registered (authorized by West Bengal AYUSH department) medicinal herbs supplier. The plants and spices were identified and authenticated by renounced taxonomist, department of Pharmaceutical Technology, Jadavpur University, Kolkata, India and properly kept as voucher specimen (Table 1). The plants and plant parts used in extract preparation are listed in Table 2.

Preparation of extract
At the onset of the experiment collected plants and spices were cleaned by double distilled water until and unless those are properly cleaned. The cleaned spices and plants were air dried through normal temperature under sunlight. After that the samples were kept in calibrated and validated hot air oven at 75 °C for 20 min and 55 °C for 30 min. To obtain fine powder the plant parts and the spices were ground by a blade mill. A standard established protocol was followed for the preparation of extract with slight modification. After collection of the final extract, it was stored at 4 °C for further use. The extract was prepared with the ratio of 2:2:5:5:5:2:1:1:1 based on the individual plants ingredients. Individual yield of the each extract were Tinospora sinensis (Lour.) Merr. 11.80%, Terminalia chebula Retz. 10.52%, Azadirachta indica A. Juss. 9.75%, Andrographis paniculata (Burm.f) Nees 12.52%, Aloe barbadensis Mill. 11.48%, Curcuma longa L. 14.06%, Trigonella foenum-graecum L. 15.05%, Piper nigrum L. 13.60% and Elettaria cardamomum (L.) Maton 12.64%. After the final collection of the working extract we obtained 92% of the plant extract which is commercially sound.

Preparation of Tinospora cordifolia extract
The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55°C for 30 min. Two hundred grams of Tinospora cordifolia powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35 °C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4 °C for further use. The percentage yield of the extract is 11.80%.

Preparation of Terminalia chebula extract
The collected plants were identified and authenticated by a well-known botanist. Plants were cleaned with double distilled water. After that the sample was dried under sunlight. The plants were kept in a hot air oven at 75 °C for 20 min and 55 °C for 30 min. Two hundred grams of Terminalia chebula powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35 °C in a rotary evaporator.

Table 1 List of ingredients and their voucher specimen number

| Sl. No. | Plant Ingredients | Voucher specimen number |
|---------|-------------------|-------------------------|
| 1.      | Tinospora sinensis (Lour.) Merr. | JU/ACK/16/151 |
| 2.      | Terminalia chebula Retz. | JU/ACK/16/152 |
| 3.      | Azadirachta indica A. Juss. | JU/ACK/16/153 |
| 4.      | Andrographis paniculata (Burm.f) Nees | JU/ACK/16/154 |
| 5.      | Aloe barbadensis Mill. | JU/ACK/16/155 |
| 6.      | Curcuma longa L. | JU/ACK/16/156 |
| 7.      | Trigonella foenum-graecum L. | JU/ACK/16/157 |
| 8.      | Piper nigrum L. | JU/ACK/16/158 |
| 9.      | Elettaria cardamomum (L.) Maton | JU/ACK/16/159 |
Preparation of Azadirachta indica extract

Azadirachta indica was collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55°C for 30 min. Five hundred grams of Azadirachta indica powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35°C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4°C for further use. The percentage yield of the extract is 9.75%.

Preparation of Andrographis paniculata extract

The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55°C for 30 min. Five hundred grams of Andrographis paniculata powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35°C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4°C for further use. The percentage yield of the extract is 12.52%.

Preparation of Aloe barbadensis miller extract

The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55°C for 30 min. Five hundred grams of Aloe barbadensis miller powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35°C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4°C for further use. The percentage yield of the extract is 11.48%.

Preparation of Curcuma longa extract

The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55°C for 30 min. Two hundred grams of Curcuma longa powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35°C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4°C for further use. The percentage yield of the extract is 14.06%.
20 min and 55 °C for 30 min. One hundred grams of *Trigonella foenum-graecum* powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35 °C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4 °C for further use. The percentage yield of the extract is 15.05%.

**Preparation of Piper nigrum extract**
The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55 °C for 30 min. One hundred grams of *Piper nigrum* powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process repeated four times, finally supernatant was evaporated under reduced pressure at 35 °C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4 °C for further use. The percentage yield of the extract is 13.60%.

**Preparation of Elettaria cardamomum extract**
The plants were collected, identified and authenticated by an experienced botanist. Then the plants were cleaned with double distilled water and dried under sunlight. The plants were kept in a hot air oven at 75°C for 20 min and 55 °C for 30 min. 100 gmo of *Elettaria cardamomum* powder were mixed with 1200 ml methanol and sonicated for 30 min using an ultrasonic bath at room temperature, centrifuged at 3000 rpm for 15 min and the supernatant was collected, process was repeated four times, finally supernatant was evaporated under reduced pressure at 35 °C in a rotary evaporator. Three milliliters of methanol was used to reconstitute the residue. The extract was filtered using Whatman filter paper and kept at 4 °C for further use. The percentage yield of the extract is 12.64%.

**Phytochemical screening**
Various essential plant secondary metabolites such as sterols and triterpenes, Mg$^{2+}$ turning test of flavonoids, alkaloids, saponins, glycosides, tannins, phenolic content, total flavonoids content in the developed multi-herbal formulation (AKSS16-LIV01) were detected through quantitative analysis with slight modification as described by Evans and Gueverra [25–27]. Detailed procedure as follows:

**Test for tannins**
0.30 g grinding powder sample was weighed into a test tube and boiled for 10 min in a water bath containing 30 ml of water. Standard filter paper was used for filtration. To 5 ml of the filtrate was added 3 drops of 0.1% ferric chloride. A brownish green or a blue black colouration showed positive test [28].

**Test for Saponin**
Distilled water (30 ml) was added to grinding powder samples (0.30 g) and boiled for 10 min in a water bath and filtered using Whatman filter paper number 42 (125 mm). A mixture of distilled water (5 ml) and filtrate (10 ml) was agitated vigorously for a stable persistent froth. The formation of emulsion on addition of three drops of olive oil showed positive result [29].

**Test for steroid**
Weighed 0.03 g sample into a beaker was mixed with 20 ml of ethanol; the component was extracted for 2 h. Five milliliters of plant extract (ethanolic) was taken and 2 ml acetic anhydride added within it. After that 2 ml of concentrated tetraoxosulphate (VI) acid was added to obtained the colour. A violet to blue or rather bluish green colour change in sample(s) indicated the presence of steroids [30].

**Test for Terpenoids**
Powder sample (0.30 g) was weighed into a beaker with 30 ml ethanol and component extracted for 2 h. Two milliliters of chloroform and 3 ml of concentrated tetraoxosulphate (VI) acid was taken and mixed vigorously. Then added 5 ml of plant extract to it. A reddish brown colour in the sample indicates the positive results and presence of terpenoids [31].

**Test for flavonoids**
0.30 g powder was taken in a cleaned beaker and added 30 ml of distilled water. The mixture was stayed for 2 h and filtered with standard filter paper number. Ten milliliters of the filtrate extract (aqueous) was taken and 5 ml of 1.0 M dilute ammonia solution added to it. After that 5 ml of concentrated tetraoxosulphate (VI) acid was added. Appearance of yellow colouration which disappeared on standing shows the presence of flavonoids [32].

**Test for alkaloids**
Two grams powder sample was in a cleaned conical flask for extraction by using 5% tetraoxosulphate (VI) acid (H$_2$SO$_4$) (20 ml) in 50% ethanol by boiling for 2 min and
filtered through Whatman filter paper. The filtrate was made alkaline using 5 ml of 28% ammonia solution (NH₃) in a separating funnel. Two sets of chloroform (5.0 ml) was for further solution extraction where chloroform solution was extracted with two 5 ml portions of 1.0 M dilute tetraoxosulphate (VI) acid. This final acid extract was then accustomed perform the subsequent test: zero.5 ml of Dragendorff’s chemical agent (Bismuth iodide solution) was mixed with 2 ml of acid extract and precipitated orange color infers the presence of organic compound [33].

Test for glycoside
To 2.00 g of sample 20 ml of water was supplementary, heated for 5 min on a water bathtub and filtered through Gem paper (12.5 cm). The subsequent tests were administered with the filtrate: (a) zero.2 ml of Fehling’s solutions A and B was mixed with 5 ml of the filtrate till it became base-forming (tested with acid-base indicator paper). A brick-red colouration on heating showed a positive result. (b) Instead of water, 15 ml of 1.0 M sulphuric acid was used to repeat the above test and the quantity of precipitate obtained compared with that of (a) above. High precipitate content indicated the presence of glycoside while low content showed the absence of glycoside [34].

Test for phenolic content
One millilitre of plant sample was taken in a test tube few drops of 10% ferric chloride solution was added to it. Violet colour appeared which indicated the presence of phenolic compounds [35].

Animals
Adult male swiss albino mice weighing 26 g ± 3 g were obtained from our CPCSEA registered central animal house facility. The animals were divided into seven experimental groups with 10 animals in each group. For acclimatization mice were kept in the environment controlled animal room for 1 week before the onset of the experiment. The animals were maintained at 12 h light/dark cycle with constant temperature (22 ± 2 °C) and humidity (54 ± 4%). Standard pellet diets (Procure from Hind liver India Limited, Mumbai) were given to the animal with and water ad libitum. The whole experimental procedure were carried out according to the new revised guidelines (2018) of CPCSEA, Ministry of Agricultural and Animal Husbandry, Govt. of India, New Delhi and approved by the Institutional Animal Ethics Committee (IAEC), Jadavpur University having approval number (IAEC/PHARM/1503/03/2019 dated 30.11.19).

Experimental protocol
Swiss albino mice were divided into seven groups where ethanol induced damage was observed for weeks followed by sanative response observation by our herbal formulation. The groups are normal control group, ethanol treated group (50% v/v), AKSS16-LIV01 low dose (75 mg/kg/day) pre-treated group, AKSS16-LIV01 middle dose (150 mg/kg/day) pre-treated group, AKSS16-LIV01 high dose (300 mg/kg/day) pre-treated group, Sylmarin pre-treated group (100 mg/kg/day) and only AKSS16-LIV01 (300 mg/kg/day) group. A detail of the protocol was given in Table 3.

Body weight gains and feed efficiency
Routine body weights of the each animal were measured and recorded in every week from the initial day to the final day of experiment and then determined the body weight alteration. Regular food consumption was calculated by measuring food residue on the basis food given at the fix time. Feed conversion was obtained by dividing total feed intake by weight gain.

Blood collection
After the experimental period 200 μL of blood sample were collected into micro-centrifuge tubes with and without EDTA (2%) from the retro orbital plexus of the mice. Blood collected from animals were taken in heparinized tube. The tubes were stay for 3 h in 45° angle at room temperature (27 °C). After that the blood samples were centrifuged at 3500 g for 15 min to obtained serum. Developed light yellow colour serum was collected with caution and stored – 4 °C for further biochemical analysis.

Table 3 Experimental Design

| Groups | Treatment |
|--------|-----------|
| I      | Normal control received only the vehicle (1 ml/kg olive oil twice a week for 8 weeks) |
| II     | Received Ethanol (50% v/v) daily for 8 weeks |
| III    | Treatment with ethanol (50% v/v) for 4 weeks and then pre-treated with herbal formulation AKSS16-LIV01 (75 mg/kg bw/day) for next 4 weeks |
| IV     | Treatment with ethanol (50% v/v) for 4 weeks and then pre-treated with herbal formulation AKSS16-LIV01 (150 mg/kg bw/day) for next 4 weeks |
| V      | Treatment with ethanol (50% v/v) for 4 weeks and then pre-treated with herbal formulation AKSS16-LIV01 (300 mg/kg bw/day) for next 4 weeks |
| VI     | Treatment with ethanol (50% v/v) for 4 weeks and then pre-treated with Silymarin standard hepatoprotective drug at a dose (100 mg/kg bw) for next 4 weeks |
| VII    | Treated with multi herbal formulation (MHF) AKSS16-LIV01 (300 mg/kg bw/day all over the experiment). |
Hematological parameters
Blood samples were taken from the retro orbital plexus of experimental mice and collected in heparinized tubes for determination of haematological parameters. Blood parameters were studied in this experiment i.e. haemoglobin (Hb), mean corpuscular volume (MCV), reticulocyte, hematocrit, mean corpuscular hemoglobin concentration (MCHC), total red blood cell (TRBC), mean corpuscular hemoglobin (MCH), total white blood cell (TWBC), platelets, and differential count using Sysmax-K1000 automatic Cell Counter.

Assessment of liver function parameters
All biochemical tests were carried out by using commercial kits with little modification of manufacturer’s instruction. Liver function test (LFT) parameters like different aminotransferase (AST & ALT), [36, 37] alkaline phosphatase (ALP), [38] gamma-glutamyl transferase (GGT), [39] globulin, [40] bilirubin (total & direct) [41] were analysed using biochemical assay kits (Merck, India) with manufacturer’s instruction. Total protein concentration was determined in the serum by the method of Lowry et al. [42].

Assessment of serum lipid profile
Serum lipid profile like Cholesterol, [43] Triglyceride, [44] Phospholipids, [45] Free fatty acids, [46] LDL-cholesterol [47] and HDL-cholesterol [48] were measured using enzymatic calorimetric kits (ELITech Diagnostic, France) according to manufacturer instructions.

Preparation of tissue homogenate
Prior to tissue biochemical analysis, 100 mg/mL of whole liver was homogenized in 50 mM phosphate buffer (pH 7.0). After homogenization the homogenate was centrifuged at 11000 rpm for 12 mins and the supernatant was collected and used for different parameters. Protein concentrations of liver supernatant were determined [49] using commercially available kit (Span Diagnostics Ltd., India) following procedure prescribed by manufacturer.

Hydroxyproline assay
Hydroxyproline assay to quantify collagen content were performed by established protocol. In details, 10 mg of freshly collected liver was homogenized in 100 mL of sterile MQ water followed by hydrolysis in 12 N HCl (100 ml) at 120 °C. Four hours later 5 mL of tissue lysate became transferred to a 96-well plate. The whole sample was incubated at 37 °C for 18 h to evaporate the acid. Test Samples have been incubated with including equal amounts of chloramine T and Ehrlich’s reagents for 40 min at 70 °C. Absorbance was recorded at 560 nm by using an ELISA plate reader (Synergy BioTek, Winooski, VT). Three to five mL blood sample was required to conduct the whole study.

Assessment of lipid peroxidation, NO and iNOS
The level of lipid peroxidation (MDA content) was measured from tissue sample and serum sample according to the method of Ohkawa [50] with slight modification. The level of NO and iNOS activity were determined from the liver homogenate through ELISA method using the commercial kit [51]. The detailed methods are as follows:

Lipid peroxidation (LPO)
The measurement was carried out using lipid peroxidation (MDA) assay kit (Sigma-aldrich Ltd., UK) in accordance to the manufacturer’s instructions. Here lipid peroxidation is determined by the reaction of malondialdehyde (MDA) with thiobarbituric acid (TBA) as a reagent to form a colouring product, proportional to the MDA present. To form the MDA-TBA adduct, the TBA solution (600 mL) was added into each sample and incubated at 95 °C for 60 min, after cooling to room temperature in an ice bath for 10 min. Two hundred milliliters of each mixture was transferred into a 96-well plate for analysis. The absorbance was measured at 532 nm [50].

NO activity
Nitrite is estimated through established “Griess Reaction” method. In this method, two-step diazotization reaction during which acidified NO$^-$ produces a nitrosating agent, which chemically reacts with sulphamic acid to supply the diazonium ion. This ion is then coupled to N-(1-naphthyl) ethylenediamine [NEDD] to make the chromophoric azo-derivative which absorbs light at 540 nm. In brief equal volumes of tissue sample and Griess reagent (5% phosphoric acid containing 0.1% NEDD, 1% sulfanilamide) was mixed and incubated in dark for 10 min at room temperature. After that the reaction mixture is measured at 540 nm. The concentration of nitrite in the sample can be determined from a sodium nitrite (NaNO$_2$) standard curve [51].

iNOS activity
Forty microliters of Reaction Mixture (mixture contains 10 μL Diluted NOS Cofactor 1 + 20 μL NOS Cofactor 2 (1X) + 5 μL NOS Substrate + 5 μL Nitrate Reductase) was taken in sample, standard and control tube and mix properly. Incubate it at 37 °C for 1 h. Ninety microliter of NOS Assay Buffer was added to sample, standard and control tube. Mixed well and then add 5 μL of Enhancer to all. All the tubes were kept in room temperature for 10 min after proper mixing. Finally 50 μL of Griess Reagent 1 and 50 μL of Griess Reagent 2 were then added
to all with proper mixing and incubated all the tubes at room temperature for 10 min. Measured the OD at 540 nm and calculated the reading through the standard curve [51].

**Assessments of antioxidant enzymes**

Tissue antioxidant enzymes activities were measured according to standard protocol with slight modification. These are as follows:

**Determination of superoxide dismutase (SOD)**

The activity of superoxide dismutase (SOD) was measured according to a well-established reported method [52]. In brief, 2.5 mL reagent solution (xanthine 0.3 mM, EDTA 0.65 mM, 140 μM NBT), sodium carbonate 0.4 M, and bovine albumin (35 mg/30 mL) was added to 0.1 mL sample and 50 μL xanthine oxidase (10 μL in 2 M ammonium sulphate), incubated at 25 °C for 20 min and mixed with 0.1 mL 8 M copper chloride. The developed color was measured at 560 nm and calculates the result with using co-factor.

**Determination catalase (CAT)**

Catalase activity was measured according to the method of Maehly (1955) [53, 54]. Briefly, on the aftermath of the addition of 5 μL liver homogenate to 0.995 mL 30 mM H₂O₂ solution in potassium phosphate buffer (pH 7.0), change in absorbance at 240 nm up to 1 min to determine catalase activity. The enzymatic activity was expressed as U/mg protein.

**Determination of glutathione (GSH) and GPx content**

Reduced glutathione (GSH) was determined from liver homogenate spectrophotometrically according to Ellman’s method [55]. GPx activity was assayed using a modified method of Hafeman al [56].

**Determination of TNF-α and TGF 1β**

Levels of TNF and TGF in the liver were measured following the procedure provided with the purchased kit [57]. Protein levels were also measured according to the literature provided with the kit.

**TGF-beta 1**

All reagents, samples and standards were freshly prepared according to manual. Ten microliters balance solution was added to each 100 μL tissue sample and mixed with caution. One hundred microliters standard and sample was added to specific selected well. Then 50 μL of 1x HRP-conjugate was added to each well except blank. Mixed gently and incubated for 60 min at 37 °C. Aspirated the liquid from each well and washed carefully. Approximately 350 μL of 1x wash buffer was added through multi-channel pipette. All the tubes were kept for 15 min before completely aspirating. After the remaining wash, aspirating eliminated any closing Wash Buffer after which inverted the plate and tapped towards smooth absorbent paper. 50 μL of substrate A and 50 μL of substrate B was added with gentle shaking. All the wells were kept in the dark for 15–20 min at 37 °C. Then 50 μL of stop solution was added to each well. The blue color changed to yellow immediately. Blank contained only 100 μL of PBS (0.02 mol/L pH 7.0–7.2). Optical density of each well immediately determined at 450 nm using a microplate reader.

**Hematoxylin and eosin staining**

Freshly liver sample was taken in a 4% formaldehyde saline solution and stored 72 h. The tissue was embedded and prepares paraffin blocks. Five micron thin sections were deparaffinised and washed with water. After washing tissue sections were treated with Mayer’s hematoxylin solution and kept for 5 min. Observed the staining quality under microscope. If staining was perfect then counter stained with 1% eosin for 2 min. The sections stained by the eosin and used dehydrated alcohol to remove any residual stain [58]. Mount the section with Canada balsam by using cover slip. Visualized the liver section under microscope and taken photographs through camera (Olympus BX51 fluorescence microscope) for further histological scoring analysis.

**Sirius red staining**

After the experimental period liver slices were fixed in 10% neutral buffered formalin for overnight and then transferred to 70% ethanol before imbedding in paraffin blocks. A paraffin-embedded liver tissue block was taken and cut into 5 mm thick sections. Paraffin-free sections were soaked for approximately 1 h in Pico-Sirius red solution (Abcam, Cambridge, MA) followed by a short rinse with acetic acid (0.05%). Liver sections were dehydrated by washing with absolute alcohol [59]. Sections were observed with a light microscope (Olympus BX51 fluorescence microscope).
Statistical analysis
Two-way analysis of variance (ANOVA) followed by Dunnett’s multiple comparison tests were executed for comparison of different parameters between the groups using a computer program GraphPad Prism (version 5.00 for Windows), GraphPad Software, California, USA. *p*-values lower than 0.05 (*p*<0.05) were considered statistically significant.

Results

Determination of phytochemical constituents
Table 4 represents qualitative analysis of the plants secondary metabolites in water extract of multi herbal formulation (AKSS16-LIV01) revealed the presence of sterols in trace amount. After the closing wash, aspirating eliminated any last Wash Buffer after which inverted the plate and tapped towards smooth absorbent paper. Triterpenes weren’t detected within the chloroform, ethanol, methanol and aqueous extract (Table 4). Saponins were detected in trace amount, flavonoids, alkaloids and glycosides were found to be moderately plethoric and therefore the presence of sterols was found to be abundant. This implies that a lot of secondary metabolites were found within the liquid extract that could be a sensible sign to determine the extract as a drug.

Determination of physical morphology, body weight, liver weight, liver index and food consumption
Figure 1 showed that chronic administration of ethanol retarded the overall growth and development (Fig. 1C) of the mice in compared with control untreated animal. Novel herbal formulation (AKSS16-LIV01) recovered the normal growth. Interestingly ethanol intoxication decreased gross body weight (Fig. 2B), food consumption (Fig. 2C) as well as daily water intake. Therapeutic treatment with AKSS16-LIV01 (150 & 300 mg/kg) prevent the toxic effects of ethanol and maintained the normal increasing body weight pattern. Moreover significantly increased liver weight and liver index (Fig. 2A) by ethanol intoxication was normalized through natural therapy (Fig. 2).

Table 4 Qualitative analysis of the phytochemical constituents of multi herbal formulation (AKSS16-LIV01)

| Phytochemicals | Ethanol | Methanol | Chloroform | Aqueous |
|----------------|---------|----------|------------|---------|
| Sterols        | (+)     | (+)      | (+)        | (++)    |
| Triterpenes    | (+)     | (+++)    | (+++)      | (++++)  |
| Flavonoids     | (+)     | (+)      | (+)        | (++)    |
| Alkaloids      | (+)     | (+)      | (+)        | (++)    |
| Saponins       | (+)     | (+)      | (+)        | (++)    |
| Glycosides     | (+)     | (+)      | (+)        | (++)    |
| Tannins        | (+)     | (+)      | (+)        | (++)    |

[+]= traces, (+)= moderate, (+++)= abundant, (−)= absence of constituents

Determination of serum AST, ALT, ALP, GGT and total protein level
Result presented in Fig. 3 indicated that levels of serum enzymes such as AST, ALT, ALP and GGT were significantly elevated (*P*<0.001) in ethanol intoxicated mice compared with normal control mice. However in dose dependent study serum enzymes like AST, ALT, ALP and GGT were significantly reduced in multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg (*P*<0.05), 150 mg/kg (*P*<0.05) and 300 mg/kg (*P*<0.001) respectively compared with ethanol treated mice. Moreover, Serum total protein level was significantly decreased (*P*<0.001) in ethanol treated mice compared with normal control mice. Pre-treatment with multi herbal formulation (AKSS16-LIV01) significantly increased the serum total protein level at a dose of 75 mg/kg (*P*<0.05), 150 mg/kg (*P*<0.05) and 300 mg/kg (*P*<0.001) respectively compared with ethanol treated mice. Multi herbal formulation (AKSS16-LIV01) showed 12% better protective effect compared with standard drug silymarin. Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone showed no such toxic effect in mice compared with control mice.

Determination of serum BUN, total bilirubin, direct bilirubin, albumin level and albumin-globulin ratio
Result presented in Table 5 indicated that levels of serum biochemical hepatotoxic marker such as blood urea nitrogen (BUN), total bilirubin and direct bilirubin were significantly elevated (*P*<0.001) in ethanol intoxicated mice compared with normal control mice. However in dose dependent study serum enzymes like BUN, total bilirubin and direct bilirubin were significantly reduced in multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg (*P*<0.05), 150 mg/kg (*P*<0.05) and 300 mg/kg (*P*<0.001) respectively compared with ethanol treated mice. Moreover, Serum total albumin level and albumin-globulin ration was significantly decreased (*P*<0.001) in ethanol treated mice compared with normal control mice. However in dose dependent study serum enzymes like BUN, total bilirubin and direct bilirubin were significantly reduced in multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg (*P*<0.05), 150 mg/kg (*P*<0.05) and 300 mg/kg (*P*<0.001) respectively compared with ethanol treated mice. Multi herbal formulation (AKSS16-LIV01) showed 10% better protective effect compared with standard drug silymarin. Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone showed no such toxic effect in mice compared with control mice.

Determination of liver non enzymatic markers
The total cholesterol, triglycerides, phospholipids, free fatty acids, LDL level of untreated ethanol control group
was significantly higher ($P < 0.001$) and HDL level significantly lower ($P < 0.001$) than the control groups presented in Table 6. In contrast, the levels of cholesterol, triglycerides, phospholipids, free fatty acids and LDL of the 75 mg/kg and 150 mg/kg body weight of multi herbal formulation (AKSS16-LIV01) with ethanol groups were significantly lower ($P < 0.05$) and HDL level significantly higher ($P < 0.05$) than the ethanol control group. Administration of multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg body, significantly normalise ($P < 0.001$) the levels of cholesterol, triglycerides, phospholipids, free fatty acids and LDL and significantly increase HDL level compared to control.

**Study of Morphological Parameters**

| Parameters          | Normal | ETHO | ETHO + AKSS-15-75 | ETHO + AKSS-15-150 | ETHO + AKSS-15-300 | Silimarlin | AKSS-16-300 |
|---------------------|--------|------|-------------------|--------------------|-------------------|------------|-------------|
| Final Body Weight (g) | 36.52±1.18 | 27.94±1.22 | 34.81±2.15       | 35.01±1.34         | 34.08±2.21       | 38.26±1.17 | 40.62±1.26  |
| Liver Weight (g)     | 1.94±0.82  | 3.50±0.28   | 2.25±0.47        | 1.98±0.18          | 1.86±0.64        | 2.03±0.64  | 1.99±0.52   |
| Liver Index          | 4.02±0.12  | 7.72±0.15   | 5.62±0.16        | 5.65±0.17          | 4.95±0.17        | 5.22±0.18  | 5.51±0.24   |

**Fig. 2** Study of the morphological parameters. A Table showed summary of the body weight, liver weight and liver index in mice. B Day wise body weight, C Mean food consumption. The values are expressed as the mean ± SEM. Significantly different from control *$p < 0.001$ and significantly different from ethanol$^*$ $p < 0.05$, **$p < 0.001$ using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test.
0.001) the deleterious effect (Table 6) caused by ethanol. Multi herbal formulation (AKSS16-LIV01) showed 15% better protective effect compared with standard drug silymarin. Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone showed no such toxic effect in mice compared with control mice.

### Determination of Haematological parameters

The hematopoietic system is one of the target organs of ethanol toxicity. The results concerning hematologic parameters depicted the Table 7 showed a significant ($p < 0.001$) decline in total erythrocyte count, total leukocyte count, hemoglobin concentration, Mean corpuscular haemoglobin concentration, neutrophil content and monocyte content in the ethanol treated animals. On the other hand, reticulocyte content content insignificantly increased in ethanol treated group, when compared with control animals. Gradually all the above mentioned parameters recover in the dose dependent AKSS16-LIV01 treated groups as compared with standard drug silymarin.

### Determination of liver hydroxyproline level

Table 8 showed the hydroxyproline level of control and experimental group. In our study, long term ethanol intoxication produced deleterious effects which was clearly indicated when we found that hydroxyprolein level significantly elevated ($P < 0.001$) in experimental mice with compared with normal untreated group. Pre-treatment

### Table 5 Effect of AKSS16-LIV01 on liver function test parameters across the groups in chronic ethanol-induced hepatic damage in mice

| Parameters       | Normal       | Ethanol       | Ethanol + AKSS16-LIV01 (75) | Ethanol+AKSS16-LIV01 (150) | Ethanol+AKSS16-LIV01 (300) | Ethanol+Silymarin (100) | AKSS16-LIV01 (300) |
|------------------|--------------|---------------|----------------------------|-----------------------------|----------------------------|--------------------------|----------------------|
| BUN (mg/dl)      | 0.41 ± 0.02  | 0.72 ± 0.04*  | 0.58 ± 0.04*                | 0.67 ± 0.02*                | 0.46 ± 0.02**              | 0.54 ± 0.03              | 0.46 ± 0.03*         |
| Total Bilirubin  | 0.12 ± 0.2   | 0.62 ± 0.11*  | 0.22 ± 0.09*                | 0.34 ± 0.08*                | 0.16 ± 0.09**              | 0.24 ± 0.08*             | 0.19 ± 0.11*         |
| Direct Bilirubin | 0.06 ± 0.001 | 0.33 ± 0.07*  | 0.19 ± 0.002*               | 0.09 ± 0.003*               | 0.09 ± 0.002**             | 0.11 ± 0.005*            | 0.07 ± 0.002*        |
| Alb (gr/dL)      | 3.48 ± 0.186 | 1.97 ± 0.036* | 2.85 ± 0.12*                | 3.70 ± 0.11*                | 4.02 ± 0.14**              | 3.32 ± 0.15*             | 3.16 ± 0.13*         |
| Alb/globulin     | 1.18 ± 0.141 | 0.54 ± 0.013* | 0.81 ± 0.091*               | 0.91 ± 0.096*               | 1.21 ± 0.095**             | 1.04 ± 0.091*            | 0.98 ± 0.135*        |

The values are expressed as the mean ± SEM. Significantly different from control $p < 0.001$ and significantly different from ethanol $p < 0.05$, $^* p < 0.001$ using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test.
with multi herbal formulation (AKSS16-LIV01) significantly reduce at a dose of 75 mg/kg (P < 0.05), 150 mg/kg (P < 0.05) and 300 mg/kg (P < 0.001) respectively compared with ethanol treated mice (Group-II). Multi herbal formulation (AKSS16-LIV01) showed 15% better protective effect compared with standard drug silymarin (Group-VI). Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone (Group-VII) showed no such toxic effect in mice compared with control mice.

### Determination of NO and iNOS

In this study result showed that in Table 8 ethanol intoxication significantly elevated (P < 0.001) NO content and iNOS level in mice as compared with normal untreated group. Pre-treatment with multi herbal formulation (AKSS16-LIV01) significantly reduce at a dose of 75 mg/kg (P < 0.05), 150 mg/kg (P < 0.05) and 300 mg/kg (P < 0.001) respectively compared with ethanol treated mice (Group-II). Multi herbal formulation (AKSS16-LIV01) showed 12% better protective effect compared with standard drug silymarin (Group-VI). Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone (Group-VII) showed no such toxic effect in mice compared with control mice.

### Determination of MDA and ROS level

Result presented in Table 8 and Fig. 4 indicated that level of lipid peroxidation (MDA level) and the tissue ROS content

| Parameters | Normal | Ethanol | Ethanol + AKSS16-LIV01 (75) | Ethanol+AKSS16-LIV01 (150) | Ethanol+AKSS16-LIV01 (300) | Ethanol + Silymarin(100) | AKSS16-LIV01 (300) |
|------------|--------|---------|-----------------------------|-----------------------------|-----------------------------|--------------------------|---------------------|
| Cholesterol (mg/dL) | 80.17 ± 5.402 | 138.99 ± 6.23 | 91.25 ± 4.29 | 114.77 ± 2.37 | 93.86 ± 3.45 | 85.11 ± 2.98 | 88.05 ± 3.16 |
| Triglyceride (mg/dL) | 42.55 ± 3.56 | 75.66 ± 4.28 | 68.61 ± 4.24 | 47.98 ± 1.97 | 37.25 ± 1.87 | 44.75 ± 3.25 | 51.23 ± 3.01 |
| Phospholipids (mg/dL) | 78.48 ± 5.29 | 141.51 ± 5.14 | 91.72 ± 2.66 | 98.51 ± 3.28 | 82.69 ± 4.87 | 88.03 ± 2.84 | 92.37 ± 3.16 |
| Free fatty acids (mg/dL) | 14.07 ± 0.79 | 34.87 ± 1.87 | 24.33 ± 1.91 | 21.22 ± 1.69 | 14.09 ± 1.22 | 19.27 ± 2.01 | 20.14 ± 1.96 |
| LDL-cholesterol (mg/dL) | 38.11 ± 1.91 | 76.94 ± 1.77 | 68.61 ± 1.77 | 47.98 ± 1.65 | 36.85 ± 1.25 | 42.52 ± 2.28 | 43.61 ± 1.88 |
| HDL-cholesterol (mg/dL) | 19.52 ± 0.88 | 9.21 ± 0.68 | 16.25 ± 0.62 | 16.24 ± 0.41 | 10.28 ± 0.28 | 21.57 ± 0.99 | 15.28 ± 0.55 |

The values are expressed as the mean ± SEM. Significantly different from control *p < 0.001 and significantly different from ethanol p < 0.05, **p < 0.001 using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test

| Parameters | Normal | Ethanol | Ethanol + AKSS16-LIV01 (75) | Ethanol+AKSS16-LIV01 (150) | Ethanol+AKSS16-LIV01 (300) | Ethanol + Silymarin(100) | AKSS16-LIV01 (300) |
|------------|--------|---------|-----------------------------|-----------------------------|-----------------------------|--------------------------|---------------------|
| Hb (g %)   | 12.1 ± 1.05 | 9.03 ± 0.89 | 12.0 ± 1.02 | 11.05 ± 0.99 | 12.51 ± 0.95 | 10.96 ± 0.74 | 11.21 ± 0.82 |
| RBC (x 10^6 cm^-2) | 10.8 ± 0.82 | 8.1 ± 0.71 | 10.5 ± 0.77 | 9.44 ± 0.71 | 10.02 ± 0.85 | 9.85 ± 0.79 | 9.62 ± 0.84 |
| RT (%)     | 2.7 ± 0.12 | 4.9 ± 0.26 | 2.6 ± 0.14 | 3.1 ± 0.14 | 2.8 ± 0.15 | 3.0 ± 0.12 | 3.6 ± 0.16 |
| HCT (%)    | 34.6 ± 0.48 | 39.4 ± 0.55 | 34.1 ± 0.44 | 35.8 ± 0.51 | 34.9 ± 0.56 | 34.4 ± 0.51 | 35.1 ± 0.77 |
| MCV (μm^3) | 37.8 ± 0.32 | 31.0 ± 0.68 | 36.7 ± 0.29 | 36.5 ± 0.44 | 35.9 ± 0.79 | 36.2 ± 0.43 | 35.5 ± 0.36 |
| MCH (pg)   | 21.2 ± 0.15 | 22.2 ± 0.14 | 22.8 ± 0.23 | 21.1 ± 0.12 | 21.4 ± 0.11 | 21.2 ± 0.14 | 21.1 ± 0.12 |
| MCHC (%)   | 41.2 ± 1.06 | 32.4 ± 0.95 | 40.2 ± 1.07 | 37.1 ± 0.92 | 39.6 ± 0.87 | 38.6 ± 0.99 | 36.2 ± 0.91 |
| Platelets  | 6.5 ± 0.02 | 5.5 ± 0.03 | 6.5 ± 0.04 | 5.8 ± 0.05 | 6.1 ± 0.07 | 5.5 ± 0.05 | 5.4 ± 0.06 |
| WBC (x 10^5 cm^-3) | 9.2 ± 0.09 | 12.4 ± 0.11 | 9.1 ± 0.08 | 10.8 ± 0.12 | 9.2 ± 0.11 | 10.1 ± 0.13 | 10.7 ± 0.11 |
| Lymphocyte | 74 ± 2.98 | 79 ± 3.04 | 72 ± 2.54 | 73 ± 3.06 | 74 ± 2.58 | 72 ± 3.08 | 71 ± 3.11 |
| Neutrophil | 26 ± 1.12 | 15 ± 0.49 | 24 ± 1.09 | 20 ± 0.56 | 25 ± 0.69 | 24 ± 0.51 | 21.52 ± 0.20 |

Data are expressed as mean ± standard deviation (N = 6). Hb: Haemoglobin, RBC: Read Blood corpuscle, RT: Reticulocyte, HCT: Haematocrit, MCV: Mean corpuscular volume, MCH: Mean corpuscular haemoglobin, MCHC: Mean corpuscular haemoglobin concentration, WBC: White Blood corpuscle

The values are expressed as the mean ± SEM. Significantly different from control *p < 0.001 and significantly different from ethanol p < 0.05, **p < 0.001 using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test
were significantly elevated \((P < 0.001)\) in ethanol intoxicated mice compared with normal control mice. However in dose dependent study MDA and ROS content were significantly reduced in multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg \((P < 0.05)\), 150 mg/kg \((P < 0.05)\) and 300 mg/kg \((P < 0.001)\) respectively compared with ethanol treated mice. Multi herbal formulation (AKSS16-LIV01) showed 12–13% better protective effect compared with standard drug silymarin. Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone showed no such toxic effect in mice compared with control mice.

### Determination of TNF-α and TGF 1β

Inflammation is commonly associated with liver dysfunction and fibrosis during chronic liver injury. The values of proinflammatory cytokines like TNF-α were determined from the hepatic tissue sample. Table 9 shows significant increase \((p < 0.001)\) of TNF-α in ethanol treated mice when compared to control untreated animals. The elevated level of TNF-α was significantly inhibited \((p < 0.05, p < 0.001)\) by the application of multi herbal formulation (AKSS16-LIV01) in dose dependent manner 150 and 300 mg/kg). This observation was comparable to that of the standard drug silymarin.

### Table 8 Effect of AKSS16-LIV01 on liver Lipid peroxidation, NO, iNOS levels and Hydroxyproline concentration in chronic ethanol-induced hepatic damage in mice

| Parameters                    | Normal | Ethanol | Ethanol + AKSS16-LIV01 (75) | Ethanol+AKSS16-LIV01 (150) | Ethanol+AKSS16-LIV01 (300) | Ethanol + Silymarin(100) | AKSS16-LIV01 (300) |
|-------------------------------|--------|---------|-----------------------------|-----------------------------|-----------------------------|-------------------------|---------------------|
| Lipid Peroxidation (nm/100 g tissue) | 68.16 ± 3.52 | 182.16 ± 3.09 | 92.35 ± 2.14 * | 77.92 ± 1.47 † | 62.58 ± 2.52 †† | 82.57 ± 1.67 † † | 66.57 ± 0.99 * |
| NO (μmol/mg protein)          | 0.64 ± 0.004 | 2.29 ± 0.002 a | 102.77 ± 0.005 * | 0.84 ± 0.004 † | 0.69 ± 0.006 †† | 0.88 ± 0.007 † † | 0.65 ± 0.002 a |
| iNOS (U/mg protein)           | 0.32 ± 0.003 | 1.11 ± 0.002 a | 0.62 ± 0.005 † | 0.46 ± 0.006 †† | 0.38 ± 0.003 † † | 0.45 ± 0.005 † † | 0.39 ± 0.003 a |
| Hydroxyproline (μg/g)         | 0.42 ± 0.004 | 0.99 ± 0.004 a | 0.76 ± 0.006 † | 0.55 ± 0.003 †† | 0.47 ± 0.002 † † | 0.56 ± 0.004 † † | 0.44 ± 0.005 a |

The values are expressed as the mean ± SEM. Significantly different from control \#\(^p < 0.001\) and significantly different from ethanol \*\(^p < 0.05, \**\(^p < 0.001\) using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test.
Therapeutic application of AKSS16-LIV01 at a dose of 300 mg/kg showed optimum protective potential against ethanol induce liver intoxication. TGF-1β is the major profibrogenic cytokine. As shown in Table 9 significantly increased level of TGF-1β was observed in ethanol treated mice when compared to control untreated animals. On the other hand, treatment with AKSS16-LIV01 both 150 mg/kg and 300 mg/kg significantly decreased (p < 0.05, p < 0.001) the level of TGF-1β when compared with ethanol treated animals. This observation was comparable to that of the standard drug silymarin. Therapeutic application of AKSS16-LIV01 at a dose of 300 mg/kg alone showed optimum protective potential against ethanol induce liver dysfunction.

**Determination of SOD, CAT, GSH and GPx level**

Result depicted in Fig. 4 indicated that levels of different antioxidant enzymes such as SOD, CAT, GSH and GPx were significantly reduce (P < 0.001) in ethanol intoxicated mice compared with normal control mice. However in dose dependent study the above antioxidant enzymes i.e. SOD, CAT, GSH and GPx were significantly reduced in multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg (P < 0.05), 150 mg/kg (P < 0.05) and 300 mg/kg (P < 0.001) respectively compared with ethanol treated mice. Multi herbal formulation (AKSS16-LIV01) showed 12–13% better protective effect compared with standard drug silymarin. Administration of Multi herbal formulation (AKSS16-LIV01) at a dose of 300 mg/kg alone showed no such toxic effect in mice compared with control mice.

**Histopathological examination**

Histologic examination Fig. 5 shows normal morphological architecture under light microscope of H&E in the control group. In the ethanol treated intoxicated groups (50:50 v/v) necrosis, hyperemia, vacuolar degeneration and infiltration of the inflammatory cells were observed which indicated hepatocellular damage (Fig. 5). In this regards, administration of multi herbal formulation (AKSS16-LIV01) at a dose of 75 mg/kg showed less damages compared with ethanol intoxicated mice (Fig. 6). Mild necrosis vacuolar degeneration and infiltration of the inflammatory cells were observed in the group 4 (150 mg/kg). In the group 5 (300 mg/kg) only mild vacuolar degeneration and infiltration of the inflammatory cells were seen which may indicate that the treatment of multi herbal formulation (AKSS16-LIV01) was very effective (Fig. 5). Histopathologic parameters of the liver tissues were graded in Fig. 5. Masson trichrom and sirius red staining photographs showed that liver’s normal architectures completely massed by chronic administration of ethanol and deleterious effect was completely restored by AKSS16-LIV01 (300 mg/kg) which fully supported the above results.

**Discussion**

People all over the globes widely consume alcoholic drinks and consequently suffer from various diseases such as liver cirrhosis, liver fibrosis, fatty liver and, hepatic cell carcinoma (liver cancer). The condition produces hepatic dysfunctions that alter the body’s normal homeostasis [60–62]. From ancient times, Indian medicinal herbs and spices are very useful for liver complications by boosting the antioxidant system and make a balance between antioxidants and prooxidants. These herbs and spices are enriched with polyphenols, flavonoids, tannins, proteins, amino acids, saponin, etc. which play a key role in liver protection against various toxicants. It also increases the body’s antioxidant ability which further increases the immune power for fighting against disease. In the present study, we developed a multi herbal formulation (MHF) composed of six Indian medicinal herbs and three Indian medicinal spices (AKSS16-LIV01) have cumulative actions in comparison with the single herbal extract. We try to establish its antioxidant and immune-suppressive effect against ethanol intoxication and find out a new safe and symptomatic medication in liver dysfunction.

The preclinical and clinical studies have already established that ethanol is a potent hepatotoxican which produces severe liver complications [63]. Liver damage by ethanol is closely associated with the generation of reactive oxygen species (ROS) such as peroxide, singlet oxygen, superoxide anion, hydroxyl ions which elevates MDA (malondialdehyde), NO (nitric oxide) and iNOS (nitric oxide synthase), suppress the cellular integrity. Ethanol intoxication showed the function of the various antioxidant enzymes gets severely reduced causing cell

| Parameters | Normal | Ethanol | Ethanol + AKSS16-LIV01 (75) | Ethanol+AKSS16-LIV01 (150) | Ethanol+AKSS16-LIV01 (300) | Ethanol + Silymarin (100) | AKSS16-LIV01 (300) |
|------------|--------|---------|-----------------------------|-----------------------------|-----------------------------|--------------------------|---------------------|
| TNF-α (pg/mg protein) | 1411.91 ± 206.53 | 2344.77 ± 371.98<sup>a</sup> | 1636.77 ± 220.97<sup>b</sup> | 1572.25 ± 236.93<sup>c</sup> | 1437.11 ± 285.83<sup>d</sup> | 1478.26 ± 288.17<sup>e</sup> | 1422.87 ± 198.54<sup>f</sup> |
| TGF-1β (pg/mg protein) | 485.90 ± 91.34 | 1237.05 ± 125.78<sup>a</sup> | 702.38 ± 98.77<sup>b</sup> | 598.11 ± 212.45<sup>c</sup> | 474.03 ± 102.76<sup>d</sup> | 552.03 ± 203.84<sup>e</sup> | 492.11 ± 88.16<sup>f</sup> |

The values are expressed as the mean ± SEM. Significantly different from control *p < 0.001 and significantly different from ethanol *p < 0.05, **p < 0.001 using analysis of variance (ANOVA) followed by Dunnett’s Multiple Comparison Test
apoptosis [64, 65]. Increased and decreased enzymatic and non-enzymatic markers of serum were also associated with this condition. Alcoholic liver disease (ALD) was normally found in liver histology, which disrupts the normal liver architecture and reduces regular functions. Our developed formulation AKSS16-LIV01 was enriched with antioxidants that could revert and lower the free radicals level. It had shown that the beneficial effects of this phytomedicine in preventing the ethanol-induced hepatotoxicity caused by free radicals.

In the present study, we evaluated the ameliorative effects of AKSS16-LIV01 against ethanol-induced hepatotoxicity in the animal model. Dose-dependent administration of ethanol increased the gross liver weight and liver-body weight ratio and decreased the cumulative body weight, which caused the changes in the behavior of mice. Administration of AKSS16-LIV01 with three specific doses gradually normalized the changes. Ethanol intoxication elevated the concentrations of key cellular enzymes like AST, ALT, ALP, and GGT present in the liver cells that leak into the serum during liver damage [66–70]. This happens because of a higher concentration of alcohol dehydrogenase in the liver, which catalyzes alcohol to its corresponding aldehyde [71]. Therefore, administration of AKSS16-LIV01 at a dose of 150 mg and 300 mg body weight could help to normalize the AST, ALT, ALP, and GGT enzyme levels. So, the developed multi herbal formulation AKSS16-LIV01 could reduce the level of these enzyme markers.

These results are also consistent with the protective effects of the developed formulation correlating with its antioxidant ability against alcohol-induced hepatocyte cells of the liver [72–74]. Our results also showed that administration of ethanol increased the serum essential enzymes blood urea nitrogen (BUN), total and direct bilirubin which was reverted with the treatment of AKSS16-LIV01. On the other hand, it was found that ethanol toxicity reduces the body’s different protein concentration and breaks the normal homeostasis. Our results showed that reduced total protein, albumin, and albumin globulin ratio were normalized by AKSS16-LIV01.

Ethanol administration increased serum total cholesterol (TC), total triglyceride (TC), phospholipids, LDL,
VLDL, and HDL levels of non-enzymatic markers which caused liver damage. This damage is attributed to the higher concentration of alcohol dehydrogenase enzyme which catalyzes alcohol to aldehyde and accumulation of export type proteins due to inhibition of the secretion of the proteins from the liver of alcoholics [75]. Both doses of AKSS16-LIV01 restored the higher level of lipid profile parameters in a dose-dependent manner to normal levels [74, 76, 77].

It is reported that in cirrhotic mice caused by ethanol produced significantly higher reactive oxygen species (ROS) and malondialdehyde (MDA) as compared to the control animals [1, 2]. In this study our result also supported the above reports. On the other hand the levels of MDA and ROS content were markedly low in the AKSS16-LIV01 treated group (Table 7). It is predicted that polyphenol, flavonoids, and other essential constituents rich multi herbal formulation (MHF) inhibit lipid peroxidation and reactive oxygen level in experimental mice [78]. During this study, the antioxidant system of liver fibrotic mice was extensively impaired, causing a high level of MDA and ROS. However, in this study higher levels of liver nitrite/nitrate indicating significantly increased production of hepatocellular NO content in response to chronic alcohol administration via induction of inducible nitric oxide synthase (iNOS) [79]. Nitric oxide (NO) and its metabolite i.e. peroxynitrite (ONOO⁻) penetrate the cell membranes through anion channels, which generates nitration of tyrosine and inactivation of biologically important proteins and enzymes [80]. In this study treatment with AKSS16-LIV01 decreased the levels of nitrite/nitrate significantly which was altered by ethanol intoxication. Increasing nitrites/nitrates production in alcoholic mice restored after receiving AKSS16-LIV01, which might be mainly due to inhibition of the hepatic cytosolic iNOS enzyme activity by the newly developed novel multi herbal formulation [81].

Oxidative stress (OS) induced by ethanol intoxication causes liver ailment in alcoholism [82, 83] and produced various health complications. Indian traditional healthcare system is extensively used for a long time to cure liver dysfunctions and augment body’s immune system. Two lines of oxidative defense (first and second order) protect cells against oxidative stress. Various scavenging
enzymes such as SOD, CAT, GPx, and GSH are neutralized and save the cell against oxidative injury. Superoxide ions were scavenged by SOD and catalase converts hydrogen peroxide (H₂O₂) to water.

In this study chronic administration of alcohol reduced CAT, SOD, GPx as well GSH level in hepatic cell in mice. Oxidative inactivation of the enzyme is the main causes of decreased SOD activity responsible for cell damage. This condition generates massive reactive oxygen species (ROS) and break cellular homeostasis [84]. Reduced glutathione (GSH) is one of the major non-protein thiol and takes a pivotal role in coordinating the antioxidant defense process. Normal cell structure & functions through its detoxification process requires equilibrium of GSH and GPx [85]. Glutathione (GSH) in combination with glutathione peroxidase (GPx) helps to metabolize hydrogen peroxide (H₂O₂) to water via biochemical pathways, thereby protecting cells against oxidative injury. Administration of our developed multi herbal formulation against ethanol intoxicated mice normalized these essential antioxidant enzymes concentrations in the liver tissue. The formulation thus protects the liver against ethanol-induced redox healing.

TNF-α is a key factor that initiates a cascade of immune responses involving the induction of cytokines after liver damage. In the damaged liver, predominantly Kuffer cells and infiltrating macrophages and neutrophils will produce TNF-α. TNF-α plays a dichotomous role in the hepatic tissue, where it not only induces hepatocyte proliferation, apoptosis, and inflammation but also is known to suppress collagen α1 gene expression. Our results showed that the administration of multi herbal formulation (AKSS16-LIV01) with ethanol reduced the level of TNF-α in the liver as compared to ethanol administration alone. This result suggested that AKSS16-LIV01 is capable to suppress TNF-α production against ethanol-induced liver damage.

Hepatic stellate cells (HSC) activation was triggered by TGF-β1, which was released from Kuffer cells as well as oxidative stress caused by ethanol. TGF-β1 regulates the production, degradation, and accumulation of the extracellular matrix (ECM) in liver fibrosis [86]. TGF-β1 leads fibrogenesis through the autocrine and paracrine effects of HSC. Our results showed that the administration of AKSS16-LIV01 with ethanol reduced the level of TGF-β1 in liver tissue as compared with ethanol administration alone. This suggested that the ameliorative effect of AKSS16-LIV01 on ethanol-induced hepatotoxicity was associated with their abilities to inhibit HSC activation by reducing TGF-β1 production.

Histopathological examination was an indication of hepatic damages after administration of ethanol. Chronic administration of ethanol developed cellular necrosis, vacuolar degeneration, hyperemia and infiltration of the inflammatory and lymphocyte cells [87–90]. The histopathology of the liver confirmed the protective effect of multi herbal formulation (AKSS16-LIV01). Application of the newly developed unique formulation restored the hepatic damage in mice and inhibits the damaging effect of ethanol. The efficacy of the developed formulation was depicted by the reduction of livers damages such as necrosis, hyperemia, vacuolar degeneration, and infiltration of the inflammatory cells; the effects were especially evident at the dose of 300 mg/kg. All the above study was compared with standard drug silymarin and it is showed that AKSS16-LIV01 has more potent hepatoprotective effect instead of standard drug.

Conclusion

The newly developed multi herbal formulation (AKSS16-LIV01) has shown protection by ethanol-induced liver injury by ameliorating oxidative stress in mice. The protective action was evidenced by liver histopathological studies, various hepatic enzymes, reduced lipid peroxidation, NO levels, and elevated antioxidant status. Apart from this AKSS16-LIV01 can suppress pro-inflammatory cytokines TNF-α and inhibit HSC activation by reducing profibrogenic cytokines TGF-β1. The presence of various constituents such as tannins, polyphenols, flavonoids in AKSS16-LIV01 could be contributed to the above mechanism.

Acknowledgments

The authors are thankful to Prof. S K Pal, Senior Professor Department of Chemical, Biological & Macromolecular Sciences S N Bose National Centre for Basic Sciences J D Block, Sector-III Salt Lake City for his guidance and valuable suggestion during this investigation.

Authors’ contributions

Soumendra Darbar (SD) and Atiskumar Chattopadhyay (AKC): Both authors conceived and designed the experiment. Soumendra Darbar and Srimoyee Saha (SS): Both authors conducted the animal and biochemical experiments, analysed the samples and compiled the data. SD, SS, AKC and Kausik Sankar Pramanik (KP): All the authors wrote and revised the manuscript. The author(s) read and approved the final manuscript.

Funding

There is no funding support for this article.

Availability of data and materials

Research data and materials can be provided on request.

Declarations

Ethics approval and consent to participate

The animals were maintained according to the guidelines recommended by Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA), Chennai, India which was approved by the Institutional Animal Ethics Committee (IAEC) (Approval No. AEC/PHARM/1503/03/2019 dated 30.11.19). All procedures complied with the Declaration of Helsinki, as revised in 1996.

Consent for publication

All authors totally agreed for the publication of this research.

Competing interests

The authors declare they have no competing interest.
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