Chemical constituents of sandbox tree \((Hura crepitans \text{ Linn.})\) and anti- hepatotoxic activity of the leaves and stem bark extracts

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The use of natural products derived from plants for therapeutic purpose is as ancient as human civilization. Sandbox tree, \(Hura crepitans \text{ L.} \) (Euphorbiaceae) is one of such plant and has been reported to have many ethnomedicinal applications especially as antimicrobial, anti-inflammatory and antihepatotoxic effects. This recent study was designed to determine the anti-hepatotoxic activity of the ethylacetate soluble fraction of the leaves and stem bark of \(H. \) crepitans and to isolate secondary metabolites. Chromatographic technique was used for isolation and Ultraviolet-Visible (UV), Infra-red (IR) and Nuclear Magnetic Resonance (NMR) spectroscopies were used for structural elucidation. Antihepatotoxicity study was carried out using carbon tetrachloride (CCl\(_4\)) induced rat model and biochemical parameters: alanine aminotransferase (ALT), aspartate aminotransferase (AST), L-\(\alpha\)-glutamyltransferase (GGT), urea and creatinine (CREA) were assayed on the serum. Phytomicrographs of the liver samples were also taken and analyzed. Our present study showed that biochemical studies of blood samples of CCl\(_4\) treated rats with value 105.0±0.001 AU inALT showed significant increase in the level of serum enzyme activities reflecting liver injury but, 69.0±13.23 AU for leaves and 53.3±2.52 AU for bark (p<0.05) indicated protection of hepatic cells. AST, GGT, urea and CREA also reduced significantly. Daphnane diterpenes, daphnetoxin acid and huratoxin were isolated from \(H. \) crepitans in this recent study along with apocynin and methylpentadecanoate. Conclusion: \(H. \) crepitans significantly reduced the level of biochemical parameters indicating protection against hepatocellular injury. Isolates obtained from this plant could also serve as lead compounds in therapy of diseases involving hepatic injury.

Key words: Daphnane diterpenes, apocynin, methylpentadecanoate, hepatoprotective, histopathology.

INTRODUCTION

Mineral, plant and animal products were the main sources of drugs for a long time and the use of these natural products with therapeutic properties is as ancient as human civilization (Hamburger and Hostettman, 1991; Simson and Ogorzaly, 1995). But recently, interest in therapeutic use of natural products, especially those
derived from plants deepened because report has it that certain percentage of drugs prescribed worldwide and those considered as basic and essential by the World Health Organization (WHO) are exclusively of plant origin or many are synthetic drugs obtained from lead compounds (Hamburger and Hostettman, 1991; Simson and Ogorzaly, 1995; Yue-Zhong, 1998). Examples of important drugs obtained from medicinal plants are morphine and codeine from Papaver somniferum, artemisinin from Artemesia annua, vincristine and vinblastine from Catharanthus roseus, digoxin from Digitalis spp., quinine and quindine from Cinchona spp., and atropine from Atropa belladona. The importance of these natural products cannot be overemphasized as these secondary plant metabolites can be lead compounds, allowing for the design and rational planning of new drugs, development and the discovery of new therapeutic properties not yet attributed to known compounds. In addition, compounds such as physostigmine, cannabinoids and yohimbine obtained from plants, are important tools used in pharmacological, physiological and biochemical studies (Hamburger and Hostettman, 1991; Simson and Ogorzaly, 1995; Yue-Zhong, 1998; Ayuveda and Verpoorte, 2005; Calixto, 2000; Cragg et al., 1997; De Pasquale, 1984; Vulto and Smet, 1988).

Sandbox tree, Hura crepitans L. (Euphorbiaceae), a plant common to the tropics has been reported to have many ethnomedical applications especially as antimicrobial, anti-inflammatory and antineutropotoxic effects (Burkill, 1994). Adedire and Ajayi (2003) reported the insecticidal activity of the seed oil. Huratoxin was isolated from the milky sap of H. crepitans along with hexahydrohuratoxin and keto-enal. Crepitin is also a toxaalbumin derived from H. crepitans and has been shown to be toxic (Jaffé, 1943; Jaffé and Seidi, 1969). Hurain, the proteolytic enzyme of the sap of H. crepitans, has also been reported to be activated by Fe²⁺ and inhibited by N⁵⁰⁰⁰, Zn²⁺, Cd²⁺ and Cl⁻ ions; its activity diminished in the presence of trypsin inhibitors from soy bean, black bean and Ascaris lumbricoides (Kawazu, 1972; Wehner et al., 1931). Lectins which are carbohydrate-binding proteins, specific for sugar moieties were isolated from the latex of H. crepitans L. and Euphorbia characias L. (Mediterranean spurge) (Barbieri et al., 1983). Recently, phytochemicals such as glycosides, alkaloids, flavonoids, tannins was reported in the seed oil (Fowomola and Akindahunsi, 2007; Oderinde et al., 2009a, b). The presence of some fatty acids, sterols and acyclic aliphatic ketones in the root of H. crepitans has also been reported (19) while antimicrobial activity of the extracts from leaves and stem bark; chemical composition and antimicrobial activity of essential oil of H. crepitans leaf (Oloyede and Olatinwo, 2014) have also been investigated. Not much report of the nature of secondary metabolites and pharmacological activity of the aerial parts of H. crepitans L. (leaf, stem or stem bark) was found in literature. The objective of this research work therefore was to isolate, purify and characterize the chemical constituents in H. crepitans leaves and stem bark and to screen the ethylacetate soluble fraction for hepatoprotective activity against carbon tetra-chloride-induced liver damage in rats since it has been established in a separate assay that H. crepitans ethylacetate fraction has antioxidant activity when compared to other fractions (Oloyede and Olatinwo, 2014). Plants with antioxidant activity have been observed to have protective effect against hepatic injury (Oloyede et al., 2011). Hepatotoxicity implies chemical-driven liver damage. Liver plays a central role in transforming and clearing chemicals and is susceptible to the toxicity from these agents. There are different kinds of disease conditions that affect the liver, some of which are liver cirrhosis, hepatitis, liver cancer amongst others. Chemicals that cause liver injury or damage are called hepatotoxins and more than 900 drugs have been implicated in causing liver damage. Hepatotoxic substances that are commonly used for in vivo and in vitro assays are CCl₄, vinyl chloride, bromobenzene and ethanol (Sharma et al., 1991; Subramonium and Pushpangadan, 1999).

MATERIALS AND METHODS

Plant collection and identification

Fresh leaves and stem bark of H. crepitans were identified and authenticated by Mr E. Donatus of the Herbarium unit of Department of Botany and Microbiology of the Faculty of Science University of Ibadan and were collected in September 2009 at the Botanical Gardens, University of Ibadan, Ibadan, Oyo state, Nigeria. This plant was also compared and found to be the same with a Herbarium specimen labelled FRIN 10565 at Forestry Research Institute of Nigeria, Ibadan, Oyo state, Nigeria.

Animals

Adult Wistar (albino) rats of average weight 160 g (14 week old) bred in the animal house of the Department of Biochemistry, University of Ibadan were used. The animals were divided into 8 groups. Five animals were used per group. The rats were kept at 27 ± 2°C for 1 week before and during the experiments and were fed with balanced livestock feeds from Pfizer, Plc and water, ad libitum. The animal experimental protocol was approved by the Institutional Animal Ethics Committee.

Chemicals and reagents

Hexane, ethyl acetate (EtOAc), methanol (MeOH), butanol (BuOH), chloroform, hydrochloric acid, naphthal, bismuth nitrate, potassium iodide, sodium hydroxide, copper acetate, NaOH, sodium chloride, copper sulphate pentahydrate, ferric chloride, conc. tetraoxosulphate (VI) acid, conc. HCl, ammonia solution, sodium potassium tartrate, potassium chloride, glacial acetic acid, Molisch reagent, Dragendorff's reagent, disodium hydrogen phosphate, dihydrogen potassium phosphate, iodine crystals, copper acetate, carbon tetrachloride were all BDH (England) chemicals and solvents (distilled prior to use), silica gel 70-230
microns (Merck, Germany), and Randox Laboratory kits for biochemical assays (Randox, UK).

**Equipment and apparatus**

Soxhlet apparatus and condenser (Pyrex, UK), Mettler analytical balance H80 (Mettler, UK), Water Bath (Gallenkamp, UK), Rotavapor R100 (Buchi, England), silica gel GF254 ( precoated aluminum sheets · Merck Germany), Centrifuge (Gallenkamp, UK), pH meter (Jenway model, UK), Astel Hearson Oven (Gallenkamp, UK), Teflon Homogenizer (Gallenkamp, UK), UV-Visible spectrophotometer (UVD-2960 model equipped with a UVWIN software version LABOMED INC, USA), IR Spectrophotometer (Perkin Elmer FT-IR system, Spectrum BX model of IR Spectrophotometer, England), Varian-Mercury NMR Spectrophotometer (England) operating at 200 MHz for H and 50 MHz for C nuclei.

**Sample preparation, extraction and fractionation procedure**

The leaves and stem bark of *H. crepituans* were weighed separately and air-dried for 4 and 6 weeks respectively until the weight was constant and then pulverized using mill machine at the Wood extraction laboratory, Department of Chemistry, University of Ibadan. The pulverized dried leaves (3.5 kg) and bark (2.5 kg) were extracted with 6.0 L and 5.0 L of distilled methanol respectively. Methanol was used to mimic local gin (alcohol) used in preparation of the plant *H. crepituans* in folklore medicine. Soxhlet apparatus was used for extraction since it has the tendency of extracting large amount of plants giving high yield within the shortest possible time. The extracts were collected and concentrated with the aid of Bucchi rotavapor at 37°C and partitioned into various fractions with distilled water, hexane, ethylacetate and butanol in other to separate plant constituents based on polarity. The EtOAc and BuOH fractions of leaves and EtOAc fraction of bark obtained above were subjected to column chromatography using silica gel of mesh size 70-230 μ (activated in an oven at 105°C for 1 h) as adsorbent. Slurry of adsorbent was prepared by mixing activated silica gel with n-hexane in optimum proportion (ratio of silica gel to extract was 30:1) and wet-packed into the column gently to avoid air-bubbles and cracking which could affect the resolution. Ethylacetate and butanol fractions (15 g each) of the leaves and bark were pre- adsorbed in silica gel and dried before loading onto the column. Hexane was used to wash the packed column prior to analysis. The loaded sample was then covered with pure white sand and small piece of cotton wool. The top of the column was usually filled with enough solvent/mixture of solvents to avoid cracking. The elution started with 100% non-polar hexane after which the polarity of the solvent(s) was increased gradually with EtOAc and methanol in a gradient elution chromatographic technique. 25 ml of each of the eluents was collected. The purity of the fractions was assessed by thin layer chromatographic analysis (TLC) using pre-coated aluminum TLC plates. The spotted plate was developed in different solvent systems depending on the degree of polarity of each constituent at room temperature. If not coloured after development, it was visualized in the iodine tank. Solvent systems used were Hexane/EtOAc (1:1), (3:1), (1:3), (4:1); Hexane/Methanol (1:1), (1:3), (3:1), (1:4); Hexane/Ethanol (1:1), (1:2), (2:1), (1:3), (4:1), (1:4) and Hexane/EtOAc/Methanol (1:1:1), (3:1:1), (4:1:1), (2:1:4).

**Elution of fractions**

A total of 126 ethylacetate fractions of leaves were collected, analyzed using TLC and pooled according to Rf values; 1-5 (H1), 6-

**Analysis of the isolates**

Spectroscopic techniques; UV, IR and NMR were used for structural elucidation. Samples (0.01% w/v) were scanned between 190 nm and 400 nm with the aid of UVD-2960 model UV/Visible spectrophotometer. Data from chart/recorder gave a graph of Absorbance against wavelength (nm). Vmax (cm⁻¹) from IR data obtained from a Perkin Elmer FT-IR system, Spectrum BX model of IR Spectrophotometer and using the KBr disc method also confirmed the structures. The 1H NMR and 13C NMR spectra of the pure compounds were determined using a 200 MHz machine for 10% (w/v) solutions in deuteriodimethylsulphoxide (DMSO) as internal standard. Pulse irradiation technique employed was FT NMR at ambient temperature. Chemical shifts are reported in ppm. Also anti-hepatotoxicity study was carried out on the ethylacetate soluble fractions of the leaves and stem bark using carbon tetrachloride induced rats.

**Experimental design for antihepatotoxicity study**

Rats which were bred to mature body mass were randomly assigned to eight groups with each group containing five rats (n=5). Animal experimental conditions are as described above. The animals were administered 100 mg/kg body mass of extracts (vehicle for the extracts was corn oil) and toxicant (CCl₄) intraperitonially. Group 1 served as control (corn oil), Group 2 (CCl₄ Only), Group 3 (CCl₄ + HCEt (leaves) pre-treatment), Group 4 (CCl₄ + HCEt (bark), pre-treatment), Group 5 (HCEt (leaves), Group 6 (HCEt (bark)), Group 7 (CCl₄ + KV pre-treatment) and Group 8 (KV) where HCEt is ethylacetate fraction of *H. crepituans* and KV is Kola viron (standard). CCl₄ was administered 3 times in a week for two weeks. Extracts and control were administered 5 times in a week for two weeks for those without pre-treatment while those pre-treated were administered within 5 times in a week for three weeks. The dose regimen and animal experimental conditions are in line with standard procedure for antihisapoptotic screening (Recknagel, 1983; Chattopadhyay, 2003; Mankani et al., 2005).

**Biochemical assay**

**Determination of haematological parameters**

The male albino rats were sacrificed by cervical dislocation after an overnight fasting. Blood samples were collected by cardiac puncture technique into clean dry centrifuge tubes and allowed to clot by standing for 30 min and later centrifuged at 3000 g for 15 min.
The supernatant (serum) was collected and stored in the refrigerator for preservation prior to estimation of various biochemical parameters. Also collected was the liver, rinsed in ice-cold 1.15% potassium chloride (KCl), blotted on filter paper and weighed appropriately. The liver tissues were then macerated, rinsed and homogenized in 4 volumes of homogenizing buffer (pH 7.4) using a Teflon homogenizer. The homogenates were spun for 10 min at 10000 r/min using cold centrifuge at 4°C. Aliquots of the supernatant obtained (the post-mitochondrial fraction) was stored at 200°C while the liver and serum for biochemical assay were kept at ice-cold temperature.

Histopathology studies

The liver for histopathology was stored in 10% formalin and then fixed in bovine solution; processed for paraffin embedding following the standard micro technique. Sections of liver were stained with haematoxylin-eosin and were observed microscopically for any histopathological changes. Photomicrographs of the liver samples were taken and analyzed (25-29). The following biochemical parameters: Alanine Aminotransferase (ALT), Aspartate Aminotransferase (AST), L-α-Glutamyltransferase (GGT), Urea and Creatinine (CREA) were assayed on the serum according to the methods described in the kits manufactured by Randox Laboratories Ltd, United Kingdom. In ALT determination, α-oxoglutarate reacted with L-alanine to give L-glutamate and pyruvate and ALT was measured by monitoring the concentration of pyruvate hydrazone formed with the 2, 4- dinitrophenylhydrazine. Transaminase activities in some sera were stimulated by high concentrations of aldehydes, ketones or oxo acids. Measurement against a serum blank instead of a reagent blank eliminated this effect. Also for AST determination, α-oxoglutarate reacted with L-aspartate to give L-glutamate and oxaloacetate, AST was measured by monitoring the concentration of oxaloacetate hydrazone formed with 2, 4- dinitrophenylhydrazine. L-α-Glutamyltransferase (α-GT/GGT) however, involved a reaction of L- α- glutamyl-3-carboxy-4-nitroanilide with glycylglycine in the presence of α-GT to give L-α- glutamylglycylglycine and 5- amino-2-nitrobenzoate. CREA, in alkaline solution reacted with picric acid to form a colored complex. The amount of the complex formed was directly proportional to the creatinine concentration and was so determined. Lastly, Urea in serum was hydrolyzed to ammonia in the presence of urease. Ammonia was then measured photometrically by the Berthelot’s reaction (Smna, 1972).

Determination of protein

Protein concentration of the various samples was determined by means of the Biuret method as described by Gornal et al. (1949) with some modification. Biuret reagent contained CuSO₄, KI and potassium sodium tartrate (KNaC₅H₄O₆. 4H₂O). Proteins form a colored complex with cupric ions (Cu²⁺) in an alkaline solution. Potassium iodide (KI) was added to the reagent in order to prevent the precipitation of Cu²⁺ ions as cuprous oxide.

Procedure for protein determination in the samples

The post mitochondrial fractions of the supernatants were diluted 5 times with distilled water so as to increase the sensitivity range of Biuret method. Diluted sample (1 ml) was added to 3 ml of Biuret reagent. The mixture was incubated at room temperature for 30 min after which the absorbance was read at 540 nm using distilled water as blank. The protein content of the samples was extrapolated from the standard Bovine Serum Albumin (BSA) curve (Said and Husein, 2009; Smna, 1972; Gornal, 1949).

Data analysis

Graphs were plotted from the experiment carried out and Absorbance Vs UI values given in the kit served as the calibration curve. ALT, AST, GGT, Urea and CREA activities in the serum were then obtained by extrapolation from the graph. The mean value ± SEM was calculated for each parameter and analyzed separately using ANOVA followed by Dunnett’s t test.

RESULTS AND DISCUSSION

Secondary plant metabolites

Secondary metabolites found in H. crepitan leaves and stem bark extracts were alkaloids, flavonoids, tannins, steroids and phenolic compounds. Five pure compounds were obtained and recrystallized but four were fully characterized. They are daphnetoxin acid, huratoxin, apocynin and methylnepentadecanoate. Daphnetoxin; a daphnane diterpene and apocynin though are known compounds are newly reported in H. crepitan. These compounds were identified by their UV, IR, ¹H and ¹³C - NMR. The spectra data obtained are in complete agreement with literature values. Structures of the compounds reported here are shown in Figure 1.

1-(4-hydroxy-3-methoxyphenyl) ethanone (Apocynin; Acetovanillone) (Ola 1):

UV [EtOH]nm (log á): 264 (0.423), 301 (0.162); IR (KBr) Vmax cm⁻¹: 3440.21 (O-H), 1750.14 (C=O), 1601-1487(C= aromatic); ¹H NMR (200 MHz; Acetone): ppm: 9.6 (O=CH), 6.75-7.39 (3H, m, aromatic), 3.2-4.6 (3H, OCH₂), 2.6 (3H, CH₃), ¹³C NMR (50 MHz, Acetone) ppm: 206.2 (C=O), 104.0 (C-unsaturated ring system), 58.6 (O-CH₃), 28 (CH₃); Molecular formulae: C₉H₁₀O₃, Mpt 112-115°C.

Huratoxin (Ola 2)

UV [EtOH]nm (log á): 290.65 (0.257), IR (KBr) Vmax cm⁻¹: 3433.26 (O-H), 2928.72 (C-H), 1653.97 (C=O), 1508.13 (C=C alkene), 1458.02 (C-H bending), 1261.68 (C-O alcohol), 1024.37 (C-O ether), 995.46 (C-H, ring), 827.00 - 632.26 and 383.39-370.49 (fingerprint region, C-H aliphatic , C – O, C – C); ¹H NMR (200 MHz; DMSO): ppm: 8.8 (1H, m, cycloalkene-H₁), 6.6-7.3 (3H, m, OH), 5.2-5.5 (6H, m, alkeine) 3.3-4.2 (10H, m, cyclic), 0.4-2.8 (28H, m, polymethylene CH /terminal CH3); ¹³C NMR (50 MHz, DMSO) ppm: 207.190 (C-3, C=O), 177.275(C-1), 174.933 (C-15), 172.473 (C-2), 145.984 (C-3’), 145.823 (C-4’), 138.839 (C-5), 109.173(C-12), 108.968 (C-16), 79.259 (C-8), 77.472(C-2’), 56.387(C9,13,14, C-3-O, bridged), 53.356 (C- 4, 5,20, 3C-O, (OHI)), 51.921 (C-6, 7), 36.649(C-10), 32.974(C-11), 30.602(C-18), 30.206(C-
17), 29.825-24.935 (C-6' - 13'), 19.019(C-1’), 18.814(C-19), 14.451-14.012 (C-14’) Molecular formulae C29H32O9, Mpt 217-219°C.

Daphnetoxin acid (Ola 3)

UV [EtOH]nm (log à): 295.07 (0.257); IR (KBr) νmax cm⁻¹: 3400.14 (O-H stretch, broad), 2907.06 (C-H, saturated), 1712.97(C=O), and 1508.13 (C=C), 1475.02 (C-H bending), 1361.68 (C-O, alcohol), 1234.37 (C-O, ether), 995.46, 827.00, 752.26 (C-H, aromatic); ¹H NMR (200 MHz; DMSO): C9,13,14, C-O, bridged ppm, J/Hz: 12.85 (OH, carboxylic acid), 7.763 (5H, cycloalkene), 6.104 (CH₂, ethylene, cyclopentene), 4.6 - 3.8 (OH, alcohol), 3.653- 3.362 (H, methine, cyclopentene, cyclohexane), 2.913 - 2.491 (H, oxiran, saturated ring system), 1.589-1.137 (H, methyl); ¹³C NMR (50 MHz, DMSO) ppm: 173.146 (C=O, C- 21), 171.301(C-2’), 162.530 (C-15), 150.143(C-2), 144.872(C-3’), 136.994(C-7’), 136.291(C-5’), 135.398(C-4’), 135.427(C-6’), 132.859(C-1), 128.984(C-1’), 128.736(C-16), 123.054(C-13), 104.429 (C-9), 97.152(C-14), 94.252(C-6), 89.069(C-7), 52.639(C-20), 51.277(C-5), 40.763(C-4), 39.928(C-11), 39.519(C-12), 38.584(C-3), 38.260(C-8), 22.534(C-10), 17.305(C-19), 11.903(C-17), 10.556 (C-18) Molecular formulae: C29H2O9, Mpt 193-196°C.

Ola 4 is a dark brownish solid substance; UV [EtOH]nm (log à): 290.65 (0.38125); IR (KBr) νmax cm⁻¹ : 3399.29 (O-H stretch, broad), 1700.18 (C=O),1653.82 - 1507.35 (C=C), 1457.81 (C-H, bend), 1030.85 (C-O), 665.89,432.89, 373.41, 361.91(fingerprint region, C-H aliphatic , C = O, C – C). No enough data to characterize it but chemical test confirmed it was a flavonoid.

Methylpentadecanoate (Ola 5)

dark brownish solid; UV [EtOH]nm (log à): 300.34 (0.374); IR (KBr) νmax cm⁻¹: 2914.08 (C-H), 1710.02 (C=O), 1076.23 (C-O), ¹H NMR (200 MHz; DMSO): ppm, J/Hz: 3.2-3.6 (3H,s, -CH -O), 2.5 (2H,s, -COCH-), 0.8-1.6 (-CH-, Aliphatic -CH- unresolved sharp singlet). ¹³C NMR; ATP (50 MHz, DMSO) ppm: 191.303 (C=O), 53.4 (OCH₃), 29.474 - 38.750 (-CH₂), 15.7 (CH₃); Molecular formulae (Figure 1). Spectroscopic studies showed that the compounds were unsaturated: The UV absorption values above 260 nm obtained for these compounds falls into the range for cyclic compounds. Transition in 270-350 nm is n-δ transition of ketones. IR spectrum displayed characteristic absorptions for signals due to OH, C=O, C=O functional groups. Oxygen function group with signals of aromatic hydrogen in the ¹H NMR was observed typical of aryl hydrogen adjacent to oxygen functions. These data are in complete agreement with those reported for these compounds in Seidl and Jaffe (1967) and Palmen et al. (1995).

Apocynin or acetovanillone is an organic compound found in natural source and has been reported to have pharmacological activities (Palmen et al., 1995). It is however newly reported in H. crepitans and was obtained as a dark greenish solid. Absorption peaks for OH at 3440.21 cm⁻¹ and 1750.14 cm⁻¹ for C=O were prominent in the IR region. The ¹H NMR revealed that it has aromatic group (6.7-7.9 ppm), ether/hydroxy/ketone group (3.2-4.6 ppm) and methylene group (2.6 ppm). The ¹³C NMR revealed the presence of carboxyl function at 206 ppm. Huratoxin was also obtained as a dark greenish solid. It is a piscicidal compound (widely used to catch fish in different part of the world) and had been isolated from the milky sap of H. crepitans. Prominent peaks corresponding to OH (3433.26 cm⁻¹), C-H (2928.72 cm⁻¹), C=O ring (2129.23 cm⁻¹), C=O (1653.97 cm⁻¹) were observed but absorption for C=O was unusually low because C=O bond adjacent to a carbonyl group results in the delocalization of δ electrons in the C=O and C=C bonds resulting in the lowering of the frequencies. The 1H NMR revealed absorption peaks at (ppm) 6.6-7.3 and 0.4-2.8 due to free OH and polymethylene (CH/terminal CH ) respectively; ¹³C NMR showed prominent peaks due to C=O at (ppm) 207.190 and C=O at 56.387-51.921. About fourteen isomers of this compound are
known to exist (Jaffé, 1943; Jaffé and Seidi, 1969; Kawazu, 1972).

Daphnetoxin is a benzoic acid-ortho ester of a diterpenoid and contained carbonyl, aromatic, hydroxy/ether and methylene/alkyl groups. Acid displacement of the cyclopentenone functional group was observed in the IR Vmax cm⁻¹: 3400.14 for O-H stretch, 2907.06 due to C-H (saturated) and 1712.97(C=O). Signals in the 1H NMR at ppm:12.85 due to OH of a carboxylic acid and 2.913 - 2.491 due to proton of a saturated ring system, oxiran; data from 13C NMR (ppm) further confirmed the structure, C=O at 173.146 and C-O at 52.639 - 51.277. Methylpentadecanoate is an ester. Prominent peaks due to C-H at 2914.08 and C=O at 1710.02 were observed in the IR spectrum. The 1H NMR displayed characteristics resonance peaks in the aliphatic region. Protons in three different environments were observed. The presence of methyl proton signal next to an ester group at 2.9-3.6 ppm, methylene protons next to a carbonyl was observed at 2.1-2.7 ppm while the terminal protons of a methyl group and other methylene protons were accounted for by signals observed at 0.8-1.2 ppm. The attached proton test experiment (ATP) revealed the presence of quaternary carbonyl carbon at 191.303 ppm and saturated carbon; methylene and methyl groups without electronegative element at 29.474 ppm and saturated carbon; methylene and methyl groups without electronegative element at 29.474 ppm and saturated carbon attached to an electronegative atom (C-O). The presence of this ester was accounted for as a break down product from the side chain of the daphnane diterpene isolated in the compound. Daphnane diterpenes with a tricycle [9.3.0] tetradeacne ring system have been isolated from the extracts of sap and seeds of Euphorbiaceae plants. Some of these compounds have been shown to exhibit cytotoxic, irritant, and analgesic activity (Barbieri et al., 1983; Fowomola and Akindahunsi, 2007; Oderinde et al., 2009a; Sharma et al., 1991; Palmen et al., 1995; James and Pickering, 1976).

Hepatoprotective activity

Carbon tetrachloride (CCl₄) is a hepatotoxin commonly used for the production of experimental liver toxicity or damage (James and Pickering, 1976; Recknagel et al., 1976; Thabrew et al., 1987; Md. Rajib et al., 2009). Assessment of liver function is done by estimating the activities of CREA and Urea as well as that of serum marker enzymes like ALT, AST, GGT (Oloyede et al., 2011; Recknagel, 1983; Chattopadhyay, 2003; Mankani et al., 2005). When liver cell plasma membrane is damaged, a variety of enzymes normally located in the cytosol are released into the blood stream. Their estimation in the serum is a useful quantitative marker of the extent and type of hepatocellular damage (Wolf et al., 1980; Ward and Daly, 1999; Venukumar and Latha, 2002). In this present study, it was revealed that the ethylacetate extracts induced suppression of increased activities of the enzymes ALT, AST, GGT, as well as Urea and CREA (which are nitrogenous waste products excreted from the body) when compared to the CCl₄ group (P<0.05). Table 1 shows the effect of ethylacetate fractions of leaves and stem bark of H. crepitans on different biochemical parameters in the serum of rats. There was a significant increase in ALT and AST levels (P<0.05) in Group 2 (CCl₄ only) when compared to other groups (control, CCl₄ + leaves extract, CCl₄ + bark extract, leaves extract only, bark extract only, CCl₄ + kolaviron and kolaviron only). The observed increase can be attributed to the release of these enzymes from the liver into the serum indicating hepatotoxin effect of CCl₄. The co-administration of CCl₄ and the extracts as well as the standard kolaviron at 200 mg extract/kg animal body weight in Groups 3, 4 and 7 brought about a decrease in the ALT and AST activity indicating that the extracts reduced the effect of tetrachloride exposure on both biochemical parameters. There was also significant decrease in ALT level (P<0.05) in Group 1 (control) when compared to Group 6 (bark extract), Group 7 (CCl₄ + kolaviron) and group 8 (kolaviron) indicating that there was no kidney impairment. Gamma-glutamyltransferase (GGT) is elevated in all forms of chemical; however, in this study, it was revealed that the ethylacetate fractions o
Table 1. Effect of Ethyl acetate fractions of leaves and stem bark of Hura crepitans on different biochemical parameters in the serum of rats measured in absorbance units (AU).

<table>
<thead>
<tr>
<th>Group</th>
<th>Treatment</th>
<th>Serum parameter</th>
<th>ALT</th>
<th>AST</th>
<th>GGT</th>
<th>Urea</th>
<th>Creatine</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Control (corn oil)</td>
<td></td>
<td>83.3±7.37*</td>
<td>37.3±9.45**</td>
<td>3.09±3.34**</td>
<td>8.85±3.07**</td>
<td>49.71±8.28</td>
<td>10.17±1.04</td>
</tr>
<tr>
<td>2</td>
<td>CCI₄</td>
<td></td>
<td>105.0±0.01*</td>
<td>100.0±15.0*</td>
<td>8.69±7.47*</td>
<td>13.38±2.89*</td>
<td>53.85±7.18</td>
<td>9.87±1.44</td>
</tr>
<tr>
<td>3</td>
<td>CCI₄ + HCₑᵢ (leaves)</td>
<td></td>
<td>78.3±6.56**</td>
<td>45.3±9.87**</td>
<td>8.42±6.55**</td>
<td>3.52±0.81**</td>
<td>52.2±4.39</td>
<td>9.00±0.01</td>
</tr>
<tr>
<td>4</td>
<td>CCI₄ + HCₑᵢ (bark)</td>
<td></td>
<td>70.0±6.56**</td>
<td>60.0±19.08***</td>
<td>8.39±7.47***</td>
<td>5.79±1.39**</td>
<td>65.25±21.97</td>
<td>8.88±0.63</td>
</tr>
<tr>
<td>5</td>
<td>HCₑᵢ (leaves)</td>
<td></td>
<td>69.0±13.23**</td>
<td>37.0±8.89**</td>
<td>6.66±7.64**</td>
<td>6.06±0.48**</td>
<td>54.38±20.53</td>
<td>8.67±0.29</td>
</tr>
<tr>
<td>6</td>
<td>HCₑᵢ (bark)</td>
<td></td>
<td>53.3±2.52***</td>
<td>47.5±0.71**</td>
<td>3.47±1.16**</td>
<td>8.29±0.11**</td>
<td>64.21±9.49</td>
<td>9.17±0.29</td>
</tr>
<tr>
<td>7</td>
<td>CCI₄ + KV</td>
<td></td>
<td>63.7±4.73***</td>
<td>49.0±1.41**</td>
<td>2.32±0.001**</td>
<td>6.48±0.40**</td>
<td>43.50±10.76</td>
<td>9.00±0.50</td>
</tr>
<tr>
<td>8</td>
<td>KV</td>
<td></td>
<td>60.0±6.19***</td>
<td>27.5±9.19**</td>
<td>2.89±0.81***</td>
<td>5.87±2.38**</td>
<td>43.50±0.001</td>
<td>8.00±2.02</td>
</tr>
</tbody>
</table>

* = The mean difference is significant at P<0.05 when compared with the control.
** = The mean difference is significant at P<0.05 when compared with the toxicant (CCl₄).
KV = Kola viron, Et = ethylacetate, CCl₄ = carbon tetrachloride.

(Recknagel and Glende, 1973) (Table 1).

Creatinine and urea which are nitrogenous waste products excreted from the body in the urine via the kidney are used exclusively in the assessment of the kidney function. Elevation of creatinine and urea is indicative of kidney impairment. There was no significant increase in creatinine (P>0.05) when Group 2 (CCl₄) and Group 1 (control) were compared with other groups but there was significant increase in urea level (P<0.05) in Group 2 (CCl₄ only) when compared to other groups. The observed increase can be attributed to the release of urea from the kidney into the blood stream indicating nephrotoxin effect of the toxicant. The effect was reduced by co-administration with the extracts. Thus, administration of ethylacetate fractions of H. crepitans leaves and stems bark gave hepatoprotective activity against the toxic effect of CCl₄, which is also supported by histological studies. Also lack of variation in protein concentration of the various samples as determined by the Biuret method showed that there was no significant increase (P>0.05) in Group 2 (toxicant only) when compared with other groups (Table 1).

Comparative histopathology studies of livers from different groups

A comparative histopathological study of liver from the groups further corroborated the hepatoprotective effect of H. crepitans. Histopathology of liver from normal control group shows prominent central vein, normal arrangement of hepatic cells. Microscopic examination of carbon tetrachloride treated liver section showed various degrees of pathological changes starting from centrilobular necrosis of hepatic cells and central lobular fatty regeneration. Photomicrographs of the liver samples are shown Figures 2-9.

Histopathological studies revealed that group 1 control animals showed normal hepatocytes (Figure 2), group 2 (CCl₄ treated) showed periportal hepatic necrosis/cellular infiltration (mild), bile duct hyperplasia (Figure 3) and also exhibited intense centrilobular necrosis, vacuolization and macrovesicular fatty change, Group 3 (CCl₄ + leaves extract) and group 4 (CCl₄ + bark extract) showed mild periportal hepatic necrosis/cellular infiltration and moderate bile duct hyperplasia (Figures 4 and 5), Group 5 (leaves extract only) and group 6 (bark extract only) showed no visible lesions (Figures 6 and 7), group 7 (CCl₄ + kolaviron) (Figure 8) had normal arrangement...
Figure 3. Liver section from Group 2 (CCl₄) showing periportal hepatic necrosis/cellular infiltration, bile duct hyperplasia and central lobular fatty regeneration (Mag x 400).

Figure 4. Liver section from Group 3 (CCl₄ + ethylacetate fraction of leaves) showing periportal hepatic necrosis/cellular infiltration (mild), bile duct hyperplasia (Mag x 400).

Figure 5. Liver section from Group 4 (CCl₄ + ethylacetate fraction of bark) showing portal congestion (mild), moderate bile duct hyperplasia (Mag x 400).

Figure 6. Liver section from Group 5 (ethylacetate fraction of leaves) showing no visible lesions (Mag x 400).

Figure 7. Liver section from Group 6 (ethylacetate fraction of stem bark) showing portal congestion (mild) and moderate normal arrangement of hepatocytes around the central vein and absence of necrosis (Mag x 400).

Figure 8. Liver section from Group 7 (CCl₄ + kola viron) showing no visible lesions (Mag x 400).
of hepatic cells while Group 8 (kolaviron only) showed very mild ventral venous congestion (Figure 9). Thus, the animals treated with ethylacetate fractions of leaves and stem bark of *H. crepitans* exhibited significant liver protection against the toxicant as evident by the presence of normal hepatic cords, absence of necrosis and lesser fatty infiltration. The present histopathological and biochemical studies of the ethylacetate fraction of *H. crepitans* confirm its medicinal application as hepatoprotective agent in traditional herbal practice. There were also indications that the presence of daphnane diterpenes in *H. crepitans* may have played a major role in the observed activity however the pure compounds must be subjected to hepatotoxicity study.

**Conclusion**

Spectroscopic analysis of the pure compounds obtained from ethyl acetate and butanol fractions of *H. crepitans* leaves and stem bark using UV-Visible, Infrared, $^1$H and $^{13}$C-NMR revealed the presence of carbonyl, hydroxyl, aromatic alkene and methylene functional groups in the pure compounds. These compounds namely daphnetoxin acid, huratoxin, apocynin and methypentadecanoate were reported. The result of the hepatoprotective activity revealed that the ethyl acetate soluble fractions of *H. crepitans* leaves and stem bark showed significant hepatoprotective activity by reducing elevated levels of serum enzymes, ALT, AST, GGT, as well as Urea and CREA levels. A comparative histopathological study of the liver from different group further corroborated the hepatoprotective efficacy of *H. crepitans* as it showed a significant regeneration of hepatocytes. This present study has therefore been able to establish the hepatoprotective influence of the leaves and stem bark of *H. crepitans* and secondary plant metabolites of medicinal importance have been isolated. However, more in vivo assays are also essential to characterize them as biological hepatoprotective agents. Additional work is also necessary to subject the pure isolated compounds from ethylacetate soluble fractions to hepatoprotective activity to know the isolates that are responsible for the observed activity in this promising plant.

**CONFLICT OF INTERESTS**

The authors have not declared any conflict of interests.

**ACKNOWLEDGEMENTS**

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![Figure 9](image_url)