MITRAGYNA SPECIOSA-INDUCED HEPATOTOXICITY-TREATED EFFECTIVELY BY PIPER BETLE: SCOPE AS A FUTURE ANTIDOTE

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ABSTRACT

Objective: Consumption of Mitragyna speciosa (MS) leads to various toxicities including hepatotoxicity. Piper betle (PB) is a herb that possesses various therapeutic properties. The aim of the present study was to examine the protective effect of PB methanol extract (PBME) on MS-induced hepatotoxicity which could pave the way for any future antidote.

Methods: Twenty-four male Sprague–Dawley rats were randomized into control and experimental groups. The control group was further divided into the negative (G-T80) and positive (G-PB) control groups. The G-T80 group (n=6) received oral gavage of the vehicle, 15% Tween 80. The G-PB group (n=6) received PBME 200 mg/kg/day, orally. The experimental group was divided into two groups, i.e., The G-MS and G-MS (PB) groups. The G-MS group (n=6) received only MS methanol extract (MSME) 500 mg/kg/day, while the G-MS (PB) group (n=6) received MSME with concomitant treatment with PBME.

Results: Histopathology examination of the G-T80 and G-PB groups showed normal histology of the liver. The G-MS group showed liver injury features such as microvesicular steatosis, ballooning degeneration, acidophilic bodies, scattered focal necrosis, fibrous portal expansion, bridging fibrosis, sinusoidal congestion, and dilatation. These features were fewer in the G-MS (PB) group which received concomitant treatment with PBME.

Conclusion: Administration of PBME exerted a protective effect against MS-induced hepatotoxicity. Future clinical trials using PB as an antidote may help in combating MS-induced hepatotoxicity.

Keywords: Mitragyna speciosa, Piper betle, Liver injury, Methanol extract, Microvesicular steatosis, Fibrosis.

INTRODUCTION

Mitragyna speciosa (MS) is a native plant of South-East Asia and it is also known as “ketum” in Malaysia or “kratom” in Thailand. It is a tropical, evergreen plant which belongs to the Rubiaceae family. Its height may reach up to 25 feet with a diameter of 2–3 feet [1]. Conventionally, it has been used in this region to treat various ailments including pain, diarrhea, fever, worm infestation, cough, and increase sexual performance [2,3].

Recent evidence shows an increase in toxicity associated with MS usage [2,3]. MS has been misused by locals in many countries. Liver being the main organ in detoxification process is the most vulnerable organ to any toxic injury [4]. Several case reports and experimental studies reported liver injury associated with MS consumption. Intrahepatic cholestasis was diagnosed in a 25-year-old man following consumption of MS with rapid dosage increment [5]. Cholestatic hepatitis was reported to develop in a 58-year-old man following 3 months’ consumption of MS [6]. Numerous experimental studies reported the development of liver injury following administration of MS extract in acute and subchronic duration [7-10]. Hence, there is a need to look for a potential antidote which can inhibit hepatotoxicity caused by MS.

Piper betle (PB) is herbal plant belongs to Piperaceae family. It is a perennial climber which is commonly found in many Asian countries. Numerous scientific studies reported therapeutic properties of PB [11,12]. PB was reported to exhibit hepatoprotective effect against various toxic insults including cadmium, carbon tetrachloride, ethanol, and D-galactosamine [13-16]. Hence, in the present study, we aimed to observe the effect of PB in hepatotoxicity caused by MS.

MATERIALS AND METHODS

Ethical clearance

The animal studies were performed after receiving approval from the Institutional Animal Care and Use Committee in Universiti Kebangsaan Malaysia.

Plant materials

M. speciosa and PB were bought from Hilleaf resources, Sungai Buloh, Selangor. The leaves were identified by Herbarium, Faculty of Science and Technology, Universiti Kebangsaan Malaysia (UKM), Bangi, Malaysia, and conserved there with specimen voucher numbers for MS and PB as UKMB 30028 and UKMB 40238, respectively.

Methanol crude extracts preparation

M. speciosa leaves were washed thoroughly, and twigs were removed to ensure that only the leaves were left out. The leaves were dried under sunlight. The dried leaves were grounded into finer pieces. A total of 250 g of grounded leaves were exhaustively Soxhlet extracted with 2.5 L methanol, producing dark-brown extract solution. The methanol was evaporated from the extract solution using rotary evaporator (HeiDolph, Germany) and vacuum pump (Buchi, Switzerland) producing 33 g of black paste of crude MS methanol extract (MSME). PB leaves underwent similar process as per description for MS leaves. Thus, 250 g of PB dried leaves yielded 30 g of dark-green paste of crude PB methanol extract (PBME).

Animals

Twenty-four (n=24) male Sprague–Dawley rats (190±10 g) were bought from the Laboratory Animal Resource Unit, UKM. Each rat...
was housed in individual plastic cage and was maintained under standard environmental condition (12 h light/dark cycle) with a room temperature of 27°C ± 2°C. The rats were fed with standard rat pellets and water *ad libitum* and were acclimatized for 1 week before being subjected to the experimental procedures.

**Study design**

The rats (n=24) were randomly divided into two groups, i.e., control and experimental groups. Control group was further subdivided into two groups: (i) Negative (G-T80) and (ii) positive (G-PB) control groups with each group consisting of six rats. The G-T80 group received the vehicle, 15% Tween 80 administered through oral gavage for 28 days. The G-PB group received 200 mg/kg/day PBME dissolved in 15% Tween 80 administered through oral gavage for 28 days. The experimental group was subdivided into two, i.e., G-MS and G-MS (PB) groups with each group consisting of six rats. The experimental groups received 500 mg/kg/day MSME dissolved in 15% Tween 80 only for 28 days. The G-MS group received only MSME, and no treatment was given. The G-MS (PB) group received MSME and concomitant treatment with PBME 200 mg/kg for 28 days. The animal experiments were performed as per earlier published study [9].

**Histopathological examination**

At the end of experiment, the rats were sacrificed and the liver was harvested. The liver was washed with 0.9% normal saline to remove excess blood. The liver tissue was fixed in 10% formalin at room temperature. The fixed tissues underwent dehydration with a series of increasing concentration of alcohol solutions, followed by clearing with xylene. The tissues were embedded in paraffin wax, and 5μm tissues were section and stained with hematoxylin and eosin, reticulin, and Masson’s trichrome. The slide was examined and the image was captured using image analyzer DMRXA2 (Leica, German).

**RESULTS**

**G-T80 group**

The G-T80 group showed normal histology of the liver (Fig. 1). Liver parenchyma mainly comprised hepatocytes that were polyhedral in shaped with euchromatic nucleus. Hepatocytes formed the hepatic plates which converged toward the central vein (CV). Hepatic plates were mostly formed by single layer of hepatocytes. Under reticulin staining, the converging pattern of hepatic plates could be clearly appreciated (Fig. 2). Liver sinusoids were located in between the anastomosing hepatic plates. Portal tract (PT) consisted of portal veins tributaries, branches of hepatic artery, and bile duct. These structures were bounded together by connective tissue. Under Masson’s trichrome staining, collagen around the PT could be clearly appreciated and the collagen was stained blue (Fig. 3).

**G-PB group**

Histopathology assessment of the G-PB group showed that the liver tissue was comparable to the G-T80 group which indicated normal histology of the liver (Fig. 1). Liver lobules had CV in the center with portal triad at its periphery (Fig. 2). Hepatocytes were polyhedral and formed anastomosing hepatic plates. No fibrotic change was observed in the G-PB group (Fig. 3).

**G-MS group**

Liver tissue of the G-MS (PB) group showed liver injury features. There was severe sinusoidal congestion and dilation with disrupted CV (Fig. 1). There was scattered focal necrosis with inflammatory cell infiltration. Under reticulin stain, there were areas of “drop out” lesion which were devoid of parenchyma (Fig. 2). Acidophilic bodies were observed and were characterized by shrunken cells with eosinophilic cytoplasm that was irregularly shaped and contained pyknotic nucleus (Fig. 4). Few hepatocytes developed ballooning degeneration, characterized by swollen and paler hepatocytes (Fig. 4). Microvesicular steatosis characterized by several tiny lipid droplets was present in some hepatocytes (Fig. 4). It appeared as if there was discontinuation of reticulum fiber when viewed under reticulin stain (Fig. 2). Masson’s trichrome staining showed few areas of fibrous portal expansion and bridging fibrosis (Fig. 3). The liver tissue showed features suggestive of hepatotoxicity.

**G-MS (PB) group**

Normal liver architecture was preserved in liver tissue of G-MS (PB) group best viewed with reticulin stain (Fig. 2). Compared to the G-MS group, there was a reduction in ballooning degeneration, microvesicular steatosis, sinusoidal congestion, and dilation. There were very minimal focal hepatic necrotic and acidophilic bodies (Fig. 1). No bridging fibrosis was found, with only few portal triads showing fibrous portal expansion (Fig. 3). Compared to the G-MS group, the G-MS (PB) group showed less features of liver injury.
necrosis found in our study compared to earlier studies which observed and scattered focal necrosis. There were differences in the pattern of microvesicular steatosis, G-MS group which received MSME 500 mg/kg/day, the liver showed reported to be safe with its median lethal dose (LD50) more than administration of PB ethanol extract for 28 days did not exhibit liver comparable features with the G-T80 group. Earlier study reported Liver of the G-PB group, which received 200 mg/kg/day of PBME, had was portal-central bridging fibrosis (black arrow) between the central vein (CV) and PT. (d) The G-MS PB group showed normal distribution of collagen in the PT and CV (Masson’s trichrome stain, ×10) hepatocytes showed ballooning degeneration (black arrow), microvesicular steatosis (red arrow), acidophilic body (blue arrow), and scattered focal necrosis with infiltration of inflammatory cells (yellow arrow). (Hematoxylin and eosin stain, ×20)

**DISCUSSION**

Histopathology assessment is the main method for assessing liver pathology in studies looking at the safety of the new drugs [17]. Histopathology assessment of toxicology studies in experimental animals is needed in predicting toxicity before it can safely proceed to human clinical trials [17]. An earlier histological study was performed using Khat with clove or green tea also showed a protective effect on the liver [18]. Hence, histopathology assessment was used in various toxicological studies in assessing toxicity of new drug or natural product. We also relied on histopathological evidence.

Liver of the G-PB group, which received 200 mg/kg/day of PBME, had comparable features with the G-T80 group. Earlier study reported administration of PB ethanol extract for 28 days did not exhibit liver injury [16]. Oral administration of methanol extract of PB leaves was reported to be safe with its median lethal dose (LD50) more than 5000 mg/kg [19].

Acute liver injury can be expressed as cytotoxic injury [4]. Cytotoxic injury may present as steatosis, degeneration, or necrosis [4]. In the G-MS group which received MSME 500 mg/kg/day, the liver showed microvesicular steatosis, ballooning degeneration, acidophilic bodies (eosinophilic degeneration), and scattered focal necrosis. There were differences in the pattern of necrosis found in our study compared to earlier studies which observed fatty changes with centrilobular necrosis following a single dose of MS extract [7,8]. In the present study, the liver showed scattered focal necrosis with microvesicular steatosis. Liver injury caused by toxic substance had variable histological presentation [4]. Scattered focal necrosis which was classified under non-zonal necrosis was reported to be the most common necrotic pattern of drug-induced liver injury [4]. Focal and non-zonal necrosis was presumed to be caused by apoptosis and non-zonal confluent necrosis may have resulted from severe apoptosis [4]. Acidophilic bodies suggestive of degenerative changes that preceded apoptosis were found scattered in the liver of the G-MS group, which may explain the scattered focal necrosis pattern found in this group.

Sinusoidal congestion was present in the group treated with MSME which was the similar to findings found in the earlier studies which observed the effect of MS extract in acute and subacute administration [7–9]. This feature was also found in liver biopsy of an individual who abused MS and had liver injury [5].

In general, fibrosis is considered as an irreversible consequence of liver damage [20]. Under Masson trichrome stain, areas of fibrous portal expansion and bridging fibrosis were observed in the G-MS group. This suggests that subchronic exposure of MSME may lead to fibrosis formation and had potential to cause irreversible liver damage.

Following concomitant administration of PBME in the G-MS (PB) group, the liver showed less hepatotoxicity features. There was fewer microvesicular steatosis, ballooning degeneration, acidophilic bodies, focal hepatic necrotic, sinusoidal congestion, and dilatation with very minimal fibrous portal expansion. PB was reported to possess hepatoprotective action. Studies examined the effect of PB on various toxic compounds including cadmium, carbon-tetrachloride, D-galactosamine, and ethanol which showed PB extract significantly reduced liver injury caused by the toxic compounds [13–16]. PB extract exhibited high antioxidant activities [21]. Active compound isolated from PB including allylpyrocatechol and chavibetol increases radicals scavenging activities [20]. Several mechanisms were postulated to be involved in the development of hepatotoxicity which included increase oxidative stress [22]. Hence, high level of antioxidant in PB may increase radical scavenging activities and this may lead to a reduction of oxidative stress. Thus, administration of PBME may reduce the toxic effect of MS and reduce any injury to the liver.

**CONCLUSION**

In the present study, subchronic administration of MSME 500 mg/kg/day caused liver injury with fibrosis formation. Concomitant treatment with PBME for 28 days significantly reduced the liver injury induced by MSME. Hence, the present study opens the opportunity for future research to observe the mechanism involved in the hepatoprotective effect of PBME in inhibiting liver injury in abusers with MS ingestion. Future clinical trials are advised to try PB extract as an effective antidote for treating hepatotoxicity.

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**AUTHOR’S CONTRIBUTION**

All the authors contributed equally to planning, conductance of study, interpretation of results and writing.

**CONFLICTS OF INTERESTS**

None of the authors have any conflicts of interests to declare.

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