

The Effect of Balm's Hydro Alcoholic Extract on Activity of Liver Enzymes and Liver Tissue

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Abstract: Balm (*Melissa officinalis*) is a fragrant plant from Lamiaceae family which has many medical properties including relaxing, anti-oxidant and anti-bacterial. This study was conducted to investigate the effect of hydro alcoholic extract of balm on activity of liver enzymes including AST, ALT and ALP and also liver tissues variation in Balb/C mice. Samples were randomly divided into five groups (three treatment groups, placebo and control group) with eight members in each group. Groups were kept under similar conditions. Hydro alcoholic extract was prepared in three doses (50, 100 and 200 mg kg⁻¹) and were IP for 20 days. Normal saline was used for placebo group. After the last injection blood samples were taken and liver tissues were separated. Changes of AST, ALT and ALP enzymes was evaluated using one way variance analysis and SPSS program (p<0.05). Tissue changes of liver were studied using coloring with H&E technique. According to results, ALT and ALP enzyme was increased (p<0.05) in 200 mg kg⁻¹ dose. Hepatocytes number and histology of liver didn't show significant variations. On the whole, hydro alcoholic extract of balm could have conservative effects via reducing ALT and ALP enzymes and no harmful effects on liver tissue. So, it can play an important role in preventing and curing diseases in traditional and also current medicine.

Key words: Balm, liver enzymes, liver tissue, mice, placebo group

INTRODUCTION

Liver tissue is ready to wide range of harmful effects of alcohol drinking, drugs and infections like viral hepatitis, cancer and other disease (Stout, 1999). In primary stages of liver injuries (before immune system responding), the patient does not feel any pain therefore this disease is called silent disease and patient may not have physical symptoms while liver tissue is hurting (Sallie *et al.*, 1991). So, the first step for diagnosing liver injuries is doing experiments for measuring level of liver enzymes.

Balm is a fragrant, vegetative perennial plant from Magnoliopsida class, Lamiales order, Lamiaceae family and *Melissa* genus (Zargari, 1995). Lemon smell is from properties of this plant and it is sometimes called lemon balm too (Lamaison *et al.*, 1991). The most common therapeutic properties of balm can be sedating, antioxidant, antispasmodic, carminative, anti-bacterial, anti-viral and anti-inflammatory properties (Yousefi *et al.*, 2011; Lamaison *et al.*, 1991). Identified compounds of balm are some of monoterpenoid aldehydes, flavonoid, polyphenolic compounds especially rosmarinic acid and monotrepen glycosides (Carnat *et al.*, 1998; Hohmann *et al.*, 1999). The essence of this plant is anti spasm and relaxing (Ghayour *et al.*, 2012). Balm is useful

in treating childhood colic (Weizman *et al.*, 1993) eliminating chronic asthma, influenza and ague (Agata *et al.*, 1993) memory enhancing and improving Alzheimer (Bennett, 2003). Probable effects of hydroalcoholic extract of Balm on liver enzymes and tissue was investigated in this study.

MATERIALS AND METHODS

Forty mice from Balb/C race with 25-30 g weight from the same age range were divided randomly in five groups including control, placebo and three treatment groups (50, 100 and 200 mg kg⁻¹). Groups were kept under similar conditions (22-26°C temperature, 40-60% humidity and usual light cycle) for 14 days with free access to water and food. Plant samples were selected from young healthy plants without disease and deficits symptoms. Aerial parts of plant were separated, washed completely, dried and doses 50, 100 and 200 mg were prepared. Obtained extract was injected for 20 days every other day. Normal saline was used to inject to placebo group and control group was not injected. The 1 day after last injection blood samples were prepared using Guillotine Method and were poured in heparinized tubes. At last, liver tissues were separated and kept in formalin (10%). Blood samples were centrifuged for 20 min (4000 cycle per min)

and plasma was separated. Activity of serum aminotransferases including alanine-aminotransferase, aspartate-aminotransferase and alkaline phosphatase were evaluated using kits. Also, tissue samples were prepared using H&E Method. Obtained data were analyzed using one way variance analysis of SPSS Program and mean comparisons were done using Duncan's multiple range test.

RESULTS AND DISCUSSION

According to results AST was increased non significantly with increase in dose which is obvious in graph 1 but this increase was more in 100 mg group than the other groups and control had the least amount. The amount of ALT and ALP were decrease significantly ($p < 0.05$) in 200 mg kg^{-1} group (Fig. 1).

Microscopic study of liver tissue show that there were no significant differences between studied groups

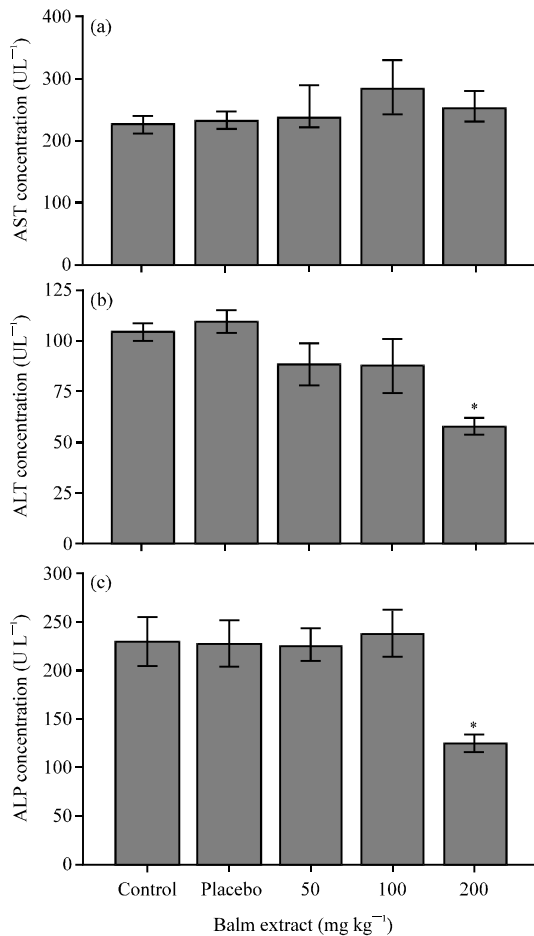


Fig. 1: Mean comparison of: a) AST; b) ALT and c) ALP amount in various experimental groups

and extract didn't cause any tissue destruction (Fig. 2). By injecting 50 mg kg^{-1} of extract lobules boundaries were faded highly, Hepatocytes (H) were became larger and more clear, Central Veins (CV) were obvious completely and hepatocytes of lobules central parts were more clear and bigger than marginal parts.

Also, studies showed that by injecting 100 mg kg^{-1} of extract Central Vhepatocytes (H) around them or hepatocytes of lobules centers have become bigger. By injecting 200 mg kg^{-1} of extract Central Veins (CV) became wider and also were changed in figure so that they have lost their typical global shape and became oval and even irregular. Statistical results showed that the highest number of hepatocytes were obtained min 200 mg kg^{-1} of extract whereas control group had the least number but the difference was not significant.

Results showed that aspartate Aminotransferase (AST) amount was increased non significantly in experimental groups. Also, alanin Amino Tranferase (ALT) amount was decreased in treatment groups especially in last group (200 mg kg^{-1}) which reduction was significant ($p < 0.05$) in this group. Activity of alkaline phosphatase was different in various doses which in 200 mg kg^{-1} it was reduced significantly. Earlier studies show that aminotransferase catalyze the chemical reactions in cells which an amino group is transferred from a donor molecule to a recipient molecule. AST is naturally located in various tissues like liver, heart, muscles, kidney

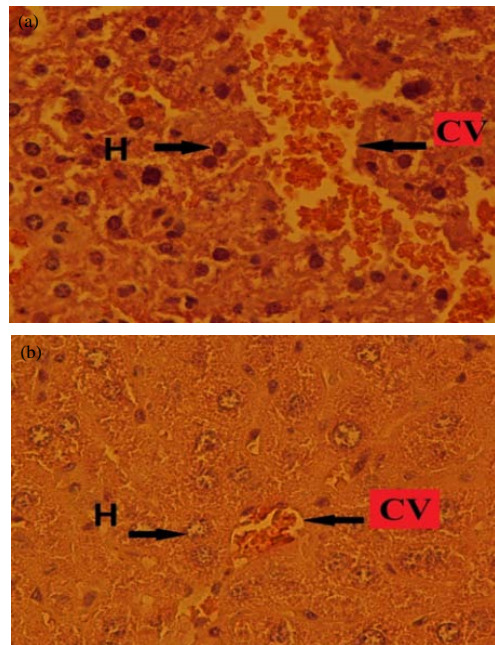


Fig. 2: Cross section of liver tissue in control group: a) x40 and b) 200 mg kg^{-1}

and brain. This enzyme enters to blood at the time of injury to any of these tissues. ALT is also found naturally in liver. Although, researchers cannot say this enzyme is exclusively in liver but this tissue has the highest concentration of this enzyme. ALT is released into the bloodstream as a result of liver damage and then it is a specific indicator of liver status (Aniya *et al.*, 2005). Considering the results of this study, hydroalcoholic extract of balm caused reduction of this enzyme in 200 mg kg⁻¹ treatment. In another study, mixture of balm, cinnamon and nettle extracts caused reduction in ALT amount (Malekirad *et al.*, 2012). It can be because of antioxidants existence because anti oxidants can play an important role in protect internal organs especially heart and liver which cause aminotransferase reduction (Aniya *et al.*, 2005; Ulicna *et al.*, 2003).

Activity of alkaline phosphatase was different in various doses which in 200 mg kg⁻¹ it was reduced significantly. In a study on grape seed and oak fruit alkaline phosphatase amount was decreased also (Mirzaei *et al.*, 2011). Thioacetamide, carbon tetrachloride, ethanol and acetaminophen are matters which are metabolized by cytochrome P450 detoxification system enzymes after entering to body (Ledda-Columbano *et al.*, 1991). Free radicals are active atoms or molecules which have a strong combination tendency with their surrounding molecules because of their last atomic layer status (Chen *et al.*, 2002). These radicals can induce lipid peroxides by alkalizing protein groups and other macro molecules and also attack to non saturated fatty acids which this can lead to liver damage. Liver necrosis leads to increase in serum level of indicator enzymes which are released from liver to blood. Increase in amount of ALT, AST and ALP enzymes and bilirubine show liver damages conventionally (Buettner, 1993). There was not considerable difference between hepatocytes number of groups of study. Also, liver tissue was also not changed considerably from histological and morphologic aspects. But Moradkhani *et al.* (2010) in a study showed that balm reduced total cholestrole and lipid of liver tissue and also increased glutation level in liver tissue and had protecting effects on liver of hyperlipidemic rats. Bolkent *et al.* (2005) announced also this protecting effect. Therefore, considering results of this study about very small histological and morphological changes of liver tissue and non significant changes of hepatocytes numbers, researchers can confirm protecting effect of balm extract too.

CONCLUSION

Finally, considering the existence of flavonoids and anti oxidants in balm extract and precautionary role of

anti-oxidants in radicals formation and also according to previous studies about balm effects on preventing and curing diseases like reducing extra fats of liver, being anti-tumor and anti-cancer and researchers can say that this plant have protective effects on liver tissue and researchers emphasize on proper use of herbal drugs again.

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