Protective effect of crocin on liver toxicity induced by morphine

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Abstract

Crocin, a bioactive molecule of saffron can be isolated and crystallized form saffron extract as a pure entity (2). Crocin has shown various pharmacological activities such as antioxidant and anticancer, radical scavenging, and genoprotective (4,5). At pharmacological and high doses, crocin did not exhibit marked damages to major organs of body and no mortality was seen by crocin in mice (6). Opium consumption in young people has been increased. Opioids have oxidative properties and therefore increase apoptosis in many cells by producing free radicals (7). Morphine is an opioid analgesic drug which is the main psychoactive chemical in opium. Morphine has a high potential for addiction, tolerance, and psychological dependence (8). Morphine increases apoptosis in neurons, glial, hepatocytes, cells of immune systems, and epithelial cells (9). Morphine can increase free radicals generation by acting as a lipid peroxidation. Lipid peroxidation is increased following blocking of anti-oxidant enzyme. This process leads to free radical formation or reactive oxygen species (ROS). These free radicals or ROS cause damage to the cell membrane and DNA fragmentation (10). Liver is considered as a major site of morphine biotransformation where morphine exerts a number of adverse physiological effects (11).
Morphine is rapidly metabolized in the liver which induces three major adverse effects on the liver: direct or indirect toxic effects, immunological effects, and oncogenic effects (12). Morphine causes liver cell injury and exerts genotoxic effect in rat liver (13). It has been recognized that morphine results in oxidative stress by inducing the generation of ROS which in turn are capable of initiating and promoting oxidative damage in the form of lipid peroxidation (14). Lipid peroxidation is known to cause cellular injury by inactivation of membrane enzymes and receptors (15). A major pathway of morphine metabolism is co-oxidation to cotinine, which is catalyzed by CYP2A6 in liver (16). Therefore, the present study is conducted to analyze protective effect of crocin on damage induced by morphine by measuring liver parameters in male mice.

**MATERIALS AND METHODS**

**Chemicals**

In this experimental study, crocin (digentiobiosyl 8, 8’-diapocarotene-8, 8’-oate; C44H64O24) powder and morphine was supplied by Merk company (Germany). They were dissolved and diluted in saline (0.9%) for administration (17).

**Experimental animals**

In this study, forty eight Balb/c male mice purchased from Tehran Razi Institute weighting 25 ± 2.2 g were used. All the animals were housed in plastic cages in a room at 22 ± 2 °C, under controlled environmental conditions, 12/12 h light/dark cycle having free access to water and food. All experimentation was conducted in conformity with ethical and humane principles of research and approval of Ethics Committee of Kermanshah University of Medical Sciences (18).

**Experimental design and treatments**

Mice were randomly divided into 8 groups of 6 each. Group 1, saline group, received 1 ml normal saline daily. Group 2, morphine group, was treated with morphine while groups 3 to 5 (crocin groups) were given 12.5, 25, 50 mg/kg crocin respectively; Groups 6 to 8, crocin+morphine groups), were treated with fixed dose of morphine plus 12.5, 25 and 50 mg/kg crocin. In group 2, morphine was injected intraperitoneally at 20 mg/kg once a day for the first 5 days, twice a day for the next 5 days and 30 mg /kg twice per day for the last 10 days of the experiment (19). In groups 3 to 5 crocin was injected interaperitoneally once a day for 20 consecutive days (20). Mice in groups 6-8, received crocin daily for 20 consecutive days and morphine in the same manner as conducted in groups 2 (19,20). The same volume of saline was injected in group 1.

**Collection of blood serum and liver weight**

All animals were anaesthetized with ketamine (75 mg/kg). Midline laparotomy was performed. Blood samples were drawn by cardiac puncture. Serum was separated and stored at −80 °C for nitric oxide measurement. Livers were removed and weighed on a microbalance sensitive to 0.001 mg (Precisa 125A, Switzerland) (21).

**Morphometric measurements**

For each hepatocyte, the total cellular area was measured. The outline of each hepatocyte was measured after taking an image with a 40× objective. The longest and shortest axis were measured in the drawing of each hepatocyte in order to estimate the mean diameter (mean axis). At least 50 hepatocytes from each zone (total 100) were measured in each liver. A separate measurement for central hepatic vein was performed, using the same methodology (22).

**Histopatological examinations**

The specimens (lower one centimeter long part of the right lobe of the liver) were fixed in 10% formalin at room temperature for 72 h. After fixing the tissue, it was thoroughly washed under running water and standard dehydration (dehydrated in ascending concentration ethanol, cleared in xylene). Also, paraffin-wax embedding procedures were used. Sections of 5 mm were cut in a microtome (Leica RM 2125, Leica Microsystems Nussloch GmbH, Germany) and
adhered to glass slides with polylysine. Hematoxylin and eosin-stained slides were prepared by standard methods and were evaluated by light microscopy (Olympus BX-51T-32E01 research microscope connected to a DP12 Camera with 3.34-million pixel resolution and Olysia Bio software (Olympus Optical Co. LTD, Tokyo-Japan) (23).

**Griess assay**

Nitric oxide was measured based on Griess colorimetric assay. Accordingly, N-1-naphthylenediamine (NEED) solution sulfonamide solutions, and nitrite standards were purchased from Sigma (USA) and prepared. To measure nitrite concentration in serum, serum samples were defreezed, 100 µl of the serum sample was deproteinized by zinc sulfate and transferred to the wells. 100 µl chloride vanadium, 50 µl sulfonamide, and 50 µl NEED solutions were added afterwards. The samples were incubated at 30 °C in darkness. Samples’ optical density (OD) was measured by enzyme linked immuno-sorbent assay (ELISA) reader (Hyperion, Germany) at 540 nm (24).

**Liver function tests**

Finally liver was minced and homogenized (10 % w/v) in ice-cold 0.1 M sodium phosphate buffer at pH 7.4. The homogenate was centrifuged twice at 10,000 rpm for 15-20 min at 4 °C. The resultant supernatant was used for various biochemical assays. The activities of aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were assayed by the method of Reitman and coworkers (25). Activities of alkaline phosphatase (ALP) were determined according to the protocol described in laboratory practical manual (26).

**Statistical calculations**

Statistical analyses were performed by analysis of variance (ANOVA) followed by LSD post-hoc test to determine statistical significance between different groups using SPSS software package 16.0. Data was presented as mean ± SEM and $P<0.05$ was considered statistically significant.

**RESULTS**

**Liver weight**

In the present study, morphine administration caused a significant decrease in the liver weight of the mice compared to the saline group ($P<0.05$). Moreover, liver weight was significantly increased in treated animals with crocin and crocin plus morphine in all doses in comparison with morphine group ($P<0.05$) (Fig. 1).

![Fig. 1. Effects of crocin, morphine and crocin plus morphine on the liver weight of various groups of Balb/c male mice. *Significant increase in crocin+morphine group (12.5 mg/kg) compared to morphine group ($P<0.05$). **Significant increase in crocin+morphine group (25 mg/kg) compared to morphine group ($P<0.05$). ***Significant increase in crocin+morphine group (50 mg/kg) compared to morphine group ($P<0.05$)](image-url)
**Morphometric measurements**

The mean diameter of hepatocytes and central hepatic vein, were significantly increased in morphine group in comparison with saline group ($P<0.05$). Further, crocin and crocin plus morphine caused a significant decrease in the mean diameter of hepatocytes and central hepatic vein in all treated groups in comparison with group treated with morphine ($P<0.05$) (Figs. 2 and 3).

**Fig. 2.** Effects of crocin, morphine and crocin plus morphine on the mean hepatocytes diameter various groups of Balb/c male mice. *Significant decrease in crocin+morphine group (12.5 mg/kg) compared to morphine group ($P<0.05$). **Significant decrease in crocin+morphine group (25 mg/kg) compared to morphine group ($P<0.05$). ***Significant decrease in crocin+morphine group (50 mg/kg) compared to morphine group ($P<0.05$).

**Fig. 3.** Effects of crocin, morphine and crocin plus morphine on the mean central hepatic vein diameter of various groups of Balb/c male mice. *Significant decrease of the mean central hepatic vein diameter in crocin+morphine group (12.5 mg/kg) compared to morphine group ($P<0.05$). **Significant decrease in crocin+morphine group (25 mg/kg) compared to morphine group ($P<0.05$). ***Significant decrease in crocin+morphine group (50 mg/kg) compared to morphine group ($P<0.05$).
Histopathological observations

Examination of hematoxylin and eosin-stained sections of liver in the saline group showed normal liver structure. After 20 days treatment with morphine, the liver section appeared with variable changes and marked injury. These changes were evidenced by disruption of tissue architecture, more distributed Kupffer cells around central vein and enlargement of central hepatic vein compared to saline stage. After 20 days treatment with crocin (50 mg/kg), liver section indicated normal histology and architecture. After treatment with morphine plus crocin (50 mg/kg) it was recognized that crocin reduced liver injury caused by morphine toxicity and largely suppressed lymphocytic infiltration (Fig. 4).

Nitric oxide

The mean nitric oxide levels in blood serum increased significantly in morphine group compared to saline group ($P<0.05$). Also, serum nitric oxide levels decreased significantly in crocin and crocin+morphine groups compared to morphine group (Fig. 5).

Biochemical analysis

Morphine caused a significant increase in ALT, AST and ALP enzymes compared to saline group ($P<0.05$). In addition, the mean ALT, AST and ALP enzymes decreased significantly in crocin and crocin+morphine groups compared to morphine group ($P<0.05$) (Fig. 6).

Fig. 4. Histological changes of liver hematoxylin-eosin staining (magnification 100×). A: Micrograph of liver section of mice in saline group showing a normal liver structure. B: Micrograph of liver section treated with crocin (50 mg/kg) showing a normal liver structure. C: Micrograph of liver section treated with morphine showing more distributed Kupffer cells around central vein (thin arrow) and enlargement of central hepatic vein (thick arrow). D: Micrograph of liver section treated with crocin plus morphine showing normal liver structure.
Fig. 5. Effects of crocin, morphine and crocin plus morphine on the mean nitric oxide levels of various groups of Balb/c male mice. *Significant decrease of nitric oxide in group of crocin+morphine group (12.5 mg/kg) compared to morphine group ($P<0.05$). **Significant decrease in crocin+morphine group (25 mg/kg) compared to morphine group ($P<0.05$). ***Significant decrease in crocin+morphine group (50 mg/kg) administration compared to morphine group ($P<0.05$).

Fig. 6. Effect of morphine, crocin and crocin plus morphine administration on liver enzymes of various groups of Balb/c male mice. A; AST enzyme, B; ALT enzyme and C; ALD enzyme. *Significant decrease in crocin+morphine group (12.5 mg/kg) compared to morphine groups ($P<0.05$). **Significant decrease in crocin+morphine group (25 mg/kg) compared to morphine groups ($P<0.05$). ***Significant decrease in crocin+morphine group (50 mg/kg) administration compared to morphine groups ($P<0.05$).
DISCUSSION

In the current study, the changes occurred in liver parenchyma following morphine administration and the protective effects of crocin on these alterations were investigated. Liver is the primary site for transformation of morphine and other opioids, and metabolism of morphine in liver reduces hepatic glutathione (27).

Results obtained from analysis of liver enzymes indicated a significant rise in ALT, AST and ALP enzymes in morphine-treated group compared with saline group, which was stabilized to a large extent by crocin treatment. The increasing activity of enzymatic markers of liver function is an indicative of liver cell impairments (28). Necrosis or impairment of the liver cell membrane would release the enzymes into the blood circulation (29). Morphine increases dopamine and xanthine oxidation and, consequently, can increase ROS. Morphine has also been reported to metabolize free radicals and significantly increase lipid peroxidation in morphine consumers (30,31). The antioxidant property of crocin seems to be responsible for its different pharmacological effects. Therefore, it seems that crocin, due to its antioxidant properties, could protect the liver cells from the damage caused by oxidative stress (32,33).

Crocin has various pharmacologic effects including preventing tumor cell proliferation and preserving the nervous system and hepatocytes. The findings of the present study were in line with those of El-Maraghy and colleagues who showed that crocin has protective effects against biochemical changes resulting from the effect of high doses of iron in liver cells (34). According to the findings of the current research, the reduced weight of liver can be linked to the damage of liver cells and dysfunctions in the real metabolism of the mice due to morphine administration (35). Todaka and coworkers showed morphine caused a decrease in the liver weight in human, which was in agreement with the results of present study (36). The increased appetite due to crocin can be considered as a factor that neutralizes the effects of morphine on weight reduction in the current study (37). This is in agreement with the results of the study conducted by Razavi and colleagues, who reported that crocin increased body weight (38). The results of this study indicated the increasing diameter of hepatocytes and the central vein of liver due to morphine administration in the study groups. Apparently, the change in the size of hepatic cells and liver’s central vein can be the result of an increase in the metabolic activity of cells to excrete toxin from the body during detoxification process (39). The free radicals induced by morphine metabolism caused lipid peroxidation, reaction with DNA and membrane proteins and consequently cell damage through various pathways (40).

Antioxidant compounds such as crocin may exert inhibitory effect on cytochrome P450, preventing further morphine metabolism and reducing the production of free radicals, consequently (41). The findings of the present study are in line with the results of the study carried out by Salahshoor and coworkers in which they showed curcumin administration resulted in the increasing diameter of hepatocytes (42). In the present research, the findings obtained from the measurement of serum nitric oxide levels revealed a significant increase between the group receiving morphine and saline group while crocin administration decreased nitric oxide levels.

The results of this study revealed destructive changes of morphine in liver tissue, increase central vein diameter and accumulation of Kupffer cells (macrophages) around the central vein. Macrophages are activated in response to tissue damage and release toxic mediators such as tumor necrosis factor alpha (TNF-α), interleukin-1 and nitric acid. Accumulation of these cells and secretion of toxic mediators in non-necrotic regions play a major role in liver toxicity induction and can induce liver necrosis (43,44). Production of free radicals and oxidative stress can be the most important reasons for liver cells’ death (45). Results of current study were in agreement with findings of Mohajeri and colleagues showing that treatment with crocin significantly reduced serum markers of liver functions (46). Administration of crocin extract to the groups receiving morphine
caused a significant reduction in degenerative changes and necrotic effects, which confirmed biochemical results. The results obtained from crocin administration seem to be associated with antioxidant properties and reduced oxidative stress of this substance (47).

The hydroxyl radicals produced by nitric oxide and superoxidation interfere with pathogenesis process and liver toxicity (48). Morphine causes an increase in nitric oxide production by regulating the intracellular calcium through activation of calcium-calmodulin-dependent nitric oxide synthase and increasing expression of nitric oxide via naloxone-sensitive receptors (49). It seems that antioxidants, including crocin are able to reduce nitric oxide production (50). The results of the study conducted by Jalili and colleagues showed that P. Crispum administration significantly increased serum nitric oxide (24) which is in contrast with the findings obtained in this study.

CONCLUSION

The present study supported contention that crocin can protect some of liver damages in mice. The results also suggest protective potential of crocin against toxic effects of morphine in morphine-treated male mice. Therefore, crocin could protect liver in individuals who are exposed to or consumed morphine. Antioxidant effect of crocin may be a major reason for its positive impact on liver parameters. However, further studies are required for a better understanding of interaction between crocin and morphine mechanism leading to preventing liver damage.

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Evaluation role of Crocin in mouse liver damage


