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RESEARCH ARTICLE

Molecular Mechanism of Crocin Induced Caspase Mediated MCF-7 Cell Death: In Vivo Toxicity Profiling and Ex Vivo Macrophage Activation

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Abstract

Background: Crocus sativus and its major constituent crocin are well established to have anti-cancer properties in breast cancer cells (MCF-7). However the role of C. sativus extract (CSE) and crocin on caspase signaling mediated MCF-7 cell death at molecular level is remains unclear. In this study, we tried to unravel role of CSE and crocin on caspase mediated MCF-7 cells death and their in vivo preclinical toxicity profiling and immune stimulatory effect. Materials and Methods: CSE extract was fractionated by HPLC and crocin was isolated and characterized by NMR, IR, and MS. MCF-7 cells were treated with both CSE and crocin and expression of Bcl-2 and Bax was assessed after 24 and 36 hours. Furthermore, caspase 3, caspase 8 and caspase 9 expression was determined by Western blotting after 24 hours of treatment. DNA fragmentation analysis was performed for genotoxicity of CSE and crocin in MCF-7 cells. The in vivo toxicity profile of CSE (300 mg/kg of b.wt) was investigated in normal Swiss albino mice. In addition, peritoneal macrophages were collected from crocin (1, 1.5 and 2 mg/kg body weight) treated mice and analyzed for ex vivo yeast phagocytosis. Results: Immunoblot analysis revealed that there was time dependent decline in anti-apoptotic Bcl-2 with simultaneous upregulation of Bax in CSE and crocin treated MCF-7 cells. Further CSE and crocin treatment downregulated caspase 8 and 9 and cleaved the caspase 3 after 24 hours. Both CSE and crocin elicited considerable DNA damage in MCF-7 cells at each concentration tested. In vivo toxicity profile by histological studies revealed no observable histopathologic differences in the liver, kidney, spleen, lungs and heart in CSE treated and untreated groups. Crocin treatment elicited significant dose and time dependent ex vivo yeast phagocytosis by peritoneal macrophages. Conclusions: Our study delineated involvement of pro-apoptotic and caspase mediated MCF-7 cell death by CSE and crocin at the molecular level accompanied with extensive DNA damage. Further we found that normal swiss albino mice can tolerate the maximum dose of CSE. Crocin enhanced ex vivo macrophage yeast phagocytic ability.

Keywords: Crocus sativus - crocin - MCF-7 cells - Bcl-2 - caspase - DNA fragmentation - toxicity

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Introduction

Breast cancer is the most common malignancy in women throughout the world, and it accounts for 18% of all female cancers and there are approximately 600,000 annual deaths worldwide (Kumar et al., 2011). Chemotherapy is the principal therapeutic option to treat cancer patients but emerging drug resistance, tumor relapse and post treatment toxicity limits their use in clinic and these hurdles facilitates breast cancer as number one killer in women (Carol et al., 2014). Finding a drug formulation to overcome above mentioned clinical consequences is an active area of cancer drug discovery research. In this context plants are promising sources for chemotherapeutic agents who can abate or reverse cancer development and/or progression. Number of studies suggests that many edible fruits, vegetables, herbs and spices contain chemicals that may reduce the incidence of cancer (Deng et al., 2002).

Crocus sativus L. commonly known as saffron is a dietary herb of the Iridaceae family. Principal components of saffron are safranal, crocin, picrocrocin and crocetin and they are pharmacologically active (Rios et al., 1996). Anticarcinogenic activity of saffron was reported in the beginning of 1990 and research on this subject has increasingly continued during the past decade (Salomi et al., 1991). Anti-cancer and anti-tumor properties of saffron have been studied in several cancer cell lines and animal model (Abdullaev, 2002; Abdullaev and Espinosa-Aguirre, 2004). It is well reported that saffron can induce apoptosis in different cancer cell lines (Tavakkol-Afshari et al., 2008; Mousavi et al., 2009a, 2009b).
Our previous studies shown that saffron can inhibit the growth of different cancer cells such as breast (Bakshi et al., 2012), pancreatic (Bakshi et al., 2010), and lung (Bakshi et al., 2012) and it also shown to be an active tumor remission agent in dalton lymphoma model (Bakshi et al., 2009). Crocin is major active component of saffron and it is reported by our group and elsewhere. It has been reported that crocin has different properties including antiplatelet (Liakopoulou-Kyriakides and Skubas, 1990), neuron-protecting (Abe and Saito, 2000), antiatherogenic (He et al., 2005), antioxidant (Chen et al., 2008), aphrodisiac (Hosseinzadeh et al., 2008), hypolipidemic effect by inhibiting pancreatic lipase, leading to the malabsorption of fat and cholesterol (Sheng et al., 2006), and anxiolytic-like effect in the rat (Pitsikas et al., 2008). Crocin possesses significant anti-proliferation effects on human colorectal cancer cells (Aung et al., 2007). This carotenoid can induce the significant alteration of gene expression profile of T24 (transitional cell carcinoma of bladder) cell. It is suggested that the antitumor effects of crocin are medicated at least in part by regulating the cell cycle controlling gene expression (Lv et al., 2008).

In the current study, we intended to elaborate the molecular mechanism of caspase mediated MCF-7 cell death induced by saffron. In vitro cytotoxicity of saffron on MCF-7 cells is well established by our group and others (Mousavi et al., 2008; Mousavi et al., 2011; Bakshi et al., 2012; Nikouei et al., 2013; Samarghandian and Borji, 2014). Further saffron reported for initiation of apoptosis in MCF-7 cells by flow cytometric analysis (Mousavi et al., 2009; Mousavi et al., 2011). However, the precise molecular mechanism of apoptosis induced by saffron and its active constituent crocin in MCF-7 cells is unclear. Flow cytometric analysis of apoptosis by propidium iodide staining does not indicate the exact mechanism of cell death rather it denotes the proportion of cells reached apoptotic state. Further In vivo preclinical toxicity profile and immunomodulatory effect of saffron is not yet reported. In order to get clear picture of caspase involvement in saffron medicated MCF-7 cell death in this study, we delineate efficacy of crocin on Caspase 3, 8 and 9 at molecular level and DNA fragmentation as genotoxic marker. Further In vivo toxicity and macrophage activation of saffron were investigated in normal Swiss albino mice.

Materials and Methods

Chemicals and reagents

Roswell park memorial institute medium (RPMI) and fetal bovine serum (FBS; heat inactivated) was obtained from life technologies (GIBCO BRL), USA, gentamycin (100µg/ml) and amphotericin B (1µg/ml), were obtained locally. Penicillin/streptomycin 100units, Dimethyl sulfoxide (DMSO), Phosphate buffer saline (PBS), Sodium bicarbonate, Calcium chloride, HEPES Buffer, Sodium chloride and disodium hydrogen phosphate were purchased from Merck, USA. Polyclonal antibody to caspase 8 and monoclonal antibodies to caspases 3 and Bcl-2 were obtained from oncogene, USA and active caspase 9 antibodies, etoposide were obtained from Calbiochem, USA. All reagents were prepared using deionized (Millipore) distilled water. All other fine chemicals were obtained from Sigma, Mo, USA. All HPLC and analytical grade solvents and silica gel required for column purification and TLC were obtained from SISCO Research Laboratories, India. The pre-coated TLC plates mesh size 60-120 was obtained from E-MERCK India.

Plant material and extraction

Saffron stigma was powdered using mortar and pestle. Powdered materials were extracted with ethanol and it is store in -20°C until use for experiment. The active component identified by HPLC (Figure 1).

Isolation and Characterization of Crocin from C. Sativus

Crocin was isolated from saffron as previously described method (Lussignoli et al., 1999). Briefly, 500 mg saffron was washed thrice with 20 ml ethyl ether, and the residual ether was evaporated in air. It was then suspended in 15 ml of 30% methanol (v/v) in distilled water and stirred for 5 minutes at room temperature. The extract was filtered through a 0.45 µm Millipore filter. It was then diluted with 10 mmol/l phosphate buffered saline (PBS, pH=7.4), and the concentration of crocin was adjusted to 25 μmol/l, using the coefficient €443=89,000 M⁻¹ cm⁻¹ reported for crocin in aqueous solution (Jorgensen et al., 1997). The crocin structure (Figure 2) was elucidated on the basis of 1HNMR, 13CNMR, IR, and mass spectral data.

Drug preparation

Different concentration of CSE (25, 50 and 75 µg/ml) was freshly prepared by reconstituting them in cell culture medium. Crocin (1, 1.5 and 2 mg/ml) was prepared in PBS.

Cell lines and Culture Method

MCF-7 cells were purchased from ATCC, USA. Cells were cultured in RPMI 1640 medium with 10% FBS and 1% antibiotics (Penicillin/streptomycin) and maintained in humidified cell incubator at 37°C and 5% CO₂.

Western blot analysis

The whole cell lysate was prepared from CSE (10 µg/ml) and crocin (10 µg/ml) treated MCF-7 cells after 24 and 36 hours as described earlier (Narayan et al., 2000). Then whole cell lysate were resolved in a 10% SDS polyacrylamide gel electrophoretically and electro transferred onto a nitrocellulose membrane. The immunoblots were probed with Bcl-2, Bax and caspase 3, 8, 9 antibodies and were visualized with the NBT/BCIP chromogenic substrate and documented.

DNA fragmentation by agarose gel electrophoresis

Inter-nucleosomal cleavage of DNA was analyzed as described previously (Kim et al., 2001). Cells were seeded in 6-well plates at a concentration of 1x10⁶ cells per ml of medium. Then cells were treated with CSE (25, 50 and 75 µg/ml), Crocin (25, 50 and 75 µg/ml). Cells were harvested after 24 hours of treatment and DNA fragmentation was assessed by gel electrophoresis.

Flow cytometric analysis

MCF-7 cells were harvested after 24 hours of CSE
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and crocin treatment at 10µg/ml. DNA content analysis (Sub-G1) was done by flow cytometry using propidium iodide (PI, sigma) (Nicoletti et al., 1991). Flow cytometry was performed using a FACSscan equipped with cell quest software (Becton Dickinson). The fluorescence emitted PI-DNA complex, was collected through an FL-2 Filter (585 nm).

In Vito Toxicity Assessment

Swiss albino (Mus musculus) male mice (25 - 30 g) were maintained under standardized, environmental conditions (22-28˚C, 60-70% relative humidity, 12 hours dark/light cycle and water ad libitum). Then animals were divided in to two groups (n=16). Group I treated with vehicle (saline) and group II treated with CSE (300 mg/kg of body weight) at weekly intervals of 60 days. All the experiments were conducted under the guidelines of Institutional Animal Ethical Committee.

Histopathological analysis

At the end of experimental period, animal were euthanized by cervical dislocation and liver, kidney, spleen, lung and heart tissues were dissected and tissue sections were stained with hematoxylin and eosin staining as described earlier (Landen et al., 2002). Microscopic evaluation was done blindly by pathologist.

Peritoneal macrophage activation in murine model by purified crocin

Animals were divided in to four groups (n=4). Group I (control), group II (crocin; 1 mg/kg of body weight), group III (crocin; 1.5 mg/kg body weight) and group IV (crocin; 2 mg/kg of body weight).

Macrophage activation was analyzed as described earlier (Bajaj et.al 2001). Mice were orally treated with different doses of crocin regularly for five days. Then peritoneal macrophages were isolated from the treated and control mice. Then macrophage was incubated for 24 hours at 37˚C in 5% CO₂. After harvesting, macrophage culture was co-incubated with heat inactivated yeast (Saccharomyces cerevisiae) at 57˚C for 45 minutes. Then macrophage cells with ingested yeast cell were counted. This was expressed as percentage of phagocytosis or percentage inhibition of yeast digestion which was calculated as:

\[
\text{% inhibition of yeast digestion} = \frac{\text{No. of active macrophages}}{\text{Total no. of macrophages}} \times 100
\]

Statistical evaluation

Data presented as mean ± SD of triplicates of three independent experiments. Experimental data were evaluated by students’ t test and one or two way analysis of variance (ANOVA). Significant difference between each set of data were considered at the confidence level of p<0.05 & p<0.001.

Results

Infrared (IR) spectrum of purified Crocin

IR spectra showed absorption bands at 945cm⁻¹(-CH=CH-trans), 1050cm⁻¹(-C-O-CO, ether), 1160cm⁻¹(-C-O-ester), 1562 and 1618cm⁻¹(-C=C-, conjugated polyene), 2920 (CH₃-aliphatic) and 3420cm⁻¹(-OH) (Table 1; Figure 3)

Mass spectral analysis of purified crocin

Purified crocin contained ions at M/Z 999, [M⁺Na]⁺, and 1015, [M⁺K]⁺ (molecular mass 976) . Elimination of Terminal gentibiosyl groups was observed, generating peaks at m/z 675, [(M⁺Na⁺H)Gnt] and 691, [(M⁺K⁺H) Gnt]⁺ (Figure 4)

¹H and ¹³C NMR Spectral analysis of purified crocin

The ⁱH and ¹³C NMR spectra of compound showed simple sub duplet signals, because crocin has C₂ structural symmetry. The ¹HNMR spectrum of crocin revealed a crocetin moiety (δH 6.49-7.31), anomoric doublets of β-configuration (δH 5.38 d, J=7.8Hz), 4.55 (d, J=6.9Hz), 4.13 (d, J=7.8Hz), 4.15 (d, J=7.8Hz) and an anomeric

Figure 1. HPLC Profile of Crocin (Bakshi et al., 2010)

Figure 2. The Chemical Structure of Crocin (Bakshi et al., 2008)

| Table 1. Infrared spectrum of Crocin |
|-------------------------------------|
| S. No. | Wavelength (cm⁻¹) | Functional group |
|--------|-------------------|-----------------|
| 1      | 945               | -CH=CH-trans    |
| 2      | 1050              | -C-O-CO, ether  |
| 3      | 1160              | -C-O-ester      |
| 4      | 1562              | -C=C-, conjugated polyene |
| 5      | 1618              | -C=C-, conjugated polyene |
| 6      | 2920              | CH₃-aliphatic    |
| 7      | 3420              | -OH             |

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MCF-7 cell proliferation through induction of apoptosis accompanied with irreparable DNA damage; we examined the cell death by DNA fragmentation. Each concentration of CSE and crocin exhibit significant DNA fragmentation in MCF-7 cells (Figure 6).

Role of CSE and crocin on Caspase expression

To investigate the role of CSE and crocin on caspase’s, we analyzed the expression level of caspase 8, caspase 9 and caspase 3 by immunoblot. The western blot analysis of whole cell lysate obtained from MCF-7 cells treated with 10 µg /ml of CSE and crocin revealed an increase

Up regulation of Bax and down regulation of Bcl-2 by CSE and Crocin

To determine the mechanism of apoptotic induction in MCF-7 cells by CSE and crocin treatment, we analyzed the expression level of Bcl-2 and Bax by western blot. The immunoblot analysis showed gradual decline in the expression levels of Bcl-2 with marked increase in the Bax protein expression levels in a time dependent manner (Figure 5). Untreated MCF-7 cells showed basal expression levels of both proteins. This data suggest the involvement of Bcl-2 in the CSE and crocin induced apoptotic process.

Role of CSE and crocin on DNA fragmentation

To elucidate whether the CSE and crocin inhibits doublet of α-configuration (δH 5.06 (d,J=3.6Hz). Two anomeric protons (δH 4.53, 5.06) were shifted to a high magnetic field and two oxymethineo on H-2 (δH 4.39 (dd, J=3.7, 10.1 Hz), 4.83 (d, J= 6.9Hz) were shifted to a low magnetic field. This observation indicated the existence of a free hydroxyl group at an anomeric position on the glycosyl groups. Furthermore, HMBC correlations were observed between the H2 of the α-glycoside and β-glycoside (Glucose C) at δH 5.06 and δH 4.53 and carbonyl carbons on the crocetin moiety at δC 167.92 and δC 167.11 respectively. Based on this chemical and spectra data compound was determined to be an all trans-crocetin β-D-2-deoxy glucopyranos-2-yl ester (Crocin).

Figure 3. IR Spectrum of Crocin

Figure 4. Mass Spectra of Crocin

Figure 5. Western Blot Analysis of Bcl-2 and Bax.

Western blot analysis of anti-apoptotic protein Bcl-2 and proapoptotic Bax done using whole cell lysate of CSE and crocin treated MCF-7 cells at different time points from 0 to 36 h: (1) untreated cells used as corresponding controls, (2) etoposide (50 µM) treated, (3) CSE (10 µg/ml) treated, and (4) crocin (10 µg/ml) treated

Figure 6. DNA Fragmentation Induced by CSE and Crocin.

Agarose gel electrophoresis demonstrating DNA fragmentation in MCF-7 cells: 1×106 MCF-7 cells treated with 25, 50, 75 µg/ml of CSE and purified crocin for 24 hours induced classical DNA fragmentation

Figure 7. Activation of Caspases (3, 8, and 9) by CSE and Crocin.

Activation of caspase 8, 9, and 3. MCF-7 cells were treated with CSE (10 µg/ml) and Crocin (10 µg/ml) for 24 hours. Whole cell lysate was used for western blot analysis with specific antibodies, respectively: (1) control (untreated MCF-7 cells), (2) etoposide (50 µM) treated, (3) CSE (10µg/ml) treated, and (4) Crocin (10 µg/ml) treated
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in caspase 8, and 9 expression and there is cleavage of caspase 3 after 24 hours (Figure 7). This data reveals that the apoptosis induction in MCF-7 cells by CSE and crocin mediated by caspase 3 dependent pathway.

Flow cytometry
To confirm that cell death observed by DNA fragmentation and expression of proapoptotic proteins which corresponds to apoptosis, we analyzed the changes in DNA content by flow cytometry. The profile of PI stained events clearly distinguished nuclei with normal diploid DNA in control cells from the nuclei with hypodiploid DNA found in treated cells. The proportion of cells in the sub-G1 region increased from 3.36% (control) to 51.45% (CSE) and 49.85% (crocin) after 24 hours of treatment (Figure 8). Taken together these results suggest that CSE and crocin were capable of inducing apoptosis in MCF-7 cells with extensive DNA damage.

In vivo pre-clinical toxicity profile
To determine the maximum In vivo tolerance level of CSE, we treated normal swiss albino mice with CSE (300mg/kg body weight) for 60 days in weekly intervals. In order to analyze the toxicity on normal tissues, we examined the liver, kidney, spleen, lung and heart of treated and untreated groups. Evaluation of numerous histological sections of these tissues from animals treated with CSE did not indicate any detectable pathologic abnormalities as examined by H&E staining. There were no observable histopathologic differences in these tissues. The liver showed normal hepatic lobular architecture, intact central vein with trapped red blood cells in a liver section from CSE treated animals. The kidney revealed normal glomeruli, proximal and distal tubules, interstitium, and blood vessels (solid arrow heads) (Figure 9A). The splenic follicles and vascular sinusoids were indistinguishable between the CSE treated and untreated control groups. The lung tissue showed normal alveoli (solid arrow heads) (Figure 9B). The heart muscle showed normal morphology among the two groups (Figure 9C).

Figure 8. Determination of Sub-G1 DNA Content in MCF-7 Cells by Flow Cytometry. Determination of sub-G1 DNA content in MCF-7 cells by flow cytometry. The 1x10^6 MCF-7 cells were treated with CSE (10 µg/ml) and crocin (10 µg/ml) for 24 hours. Cells were harvested and stained with propidium iodide as described. The area labeled M1 is the sub-G1 apoptotic population, M2 is G1, M3 is S and M4 is G2/M-phase: (A) untreated control cells, (B) MCF-7 cells treated with the CSE (10 µg/ml), and (C) MCF-7 cells treated with Crocin (10 µg/ml). Experiment repeated thrice and the profile was similar.

Figure 9. A: Organ Toxicity (Liver and kidney) Profile of CSE treated and untreated Mice. B: Organ Toxicity (spleen and lungs) Profile of CSE treated and untreated Mice. C: Organ Toxicity (Heart) Profile of CSE treated and untreated Mice

Figure 10. Activation of Peritoneal Macrophage by Crocin. (A) Macrophage (control) (B) Crocin untreated macrophage with yeast (C) Crocin treated macrophage with yeast (arrow head indicates the yeast engulfed by macrophage) normal morphology among the two groups (Figure 9C).

Ex vivo immune stimulatory effect of purified crocin
9. Treatment of MCF-7 cells with 10 µg/ml of CSE and issue, we determine expression of pro caspase 3, 8 and Bcl-2 down regulation in MCF-7 cells. To address this consequent signaling cascade of CSE and crocin mediated apoptosis inducing signaling complex and translocated to mitochondria where apoptosis initiated (Shamas-Din et al., 2011). This phenomenon led us to determine apoptotic proteins such as caspase 3, 8 and 9 undergo post-translational modifications that include dephosphorylation and cleavage leading to their activation and formation of death inducing signaling complex and translocated to mitochondria where apoptosis initiated (Shamas-Din et al., 2011). This phenomenon led us to determine consequent signaling cascade of CSE and crocin mediated Bcl-2 down regulation in MCF-7 cells. To address this issue, we determine expression of pro caspase 3, 8 and 9. Treatment of MCF-7 cells with 10 µg/ml of CSE and crocin leads to up regulation of caspase 3, 8 and 9 in time dependent manner (Figure 7). These findings suggest that the CSE and crocin induced apoptosis in MCF-7 cells via the Bcl-2 down regulation and caspase 3 dependent pathways. Biochemical signatures accompanied with apoptosis include chromosomal DNA cleavage into inter-nucleosomal fragments, phosphatidylserine externalization and a number of intracellular substrate cleavages by specific proteolysis (Cohen et al., 1994; Martin and Green, 1995). As apoptosis occurs when DNA damage is irreparable, we tested the efficacy of CSE and crocin on MCF-7 cells DNA fragmentation. Each concentration of CSE and crocin treatment elicits considerable DNA fragmentation after 24 hours (Figure 6). Taken together this data evidenced that CSE and crocin modulating the expression of pro apoptotic markers in MCF-7 cells along with severe DNA damage and thus elicits significant cell death.

Toxicity in many tissues following chemotherapy is a major clinical concern. Therefore, the search for safe, well tolerated regimen has been major goal of clinical cancer research. To determine whether 300mg/kg of CSE treatment for 60 days results in toxicities to normal tissues, we examined the liver, kidney, spleen, lung and heart of treated and untreated groups. Evaluation of numerous histological sections of these tissues from animals treated with CSE did not indicate any detectable pathologic abnormalities as examined by H&E staining. There were no observable histopathologic differences in these tissues.

As a consequence of toxicity, cancer patients are prone for variety of infections due to weakening of their immune system by chemotherapeutic drugs (Vento and Cainelli, 2003). Macrophage playing a pivotal role in neutralizing infections but, deteriorated effect of chemo drugs abrogates its functions in cancer patients. To address this issue and to assess the immune toxicity of crocin, in this study, we tested the efficacy of crocin on macrophage activation (ex vivo). Macrophage collected from crocin treated mice found to be active and its phagocytic ability enhanced in each time intervals of experimental period (Figure 11). Activated macrophages release a wide array of mediators including reactive oxygen and nitrogen species, hydrolytic enzymes, bioactive lipids, and cytokines such as tumor necrosis factor alpha (TNF-α) (Nathan, 1987; Janeway and Medzhitov, 2002; Martini, 2004). Our ex vivo data reveals that peritoneal macrophage collected from crocin treated mice engulf the yeast prominently however, precise molecular mechanism for macrophage activation by crocin is remains unclear. In support our results, previously Escribano et.al reported that non-toxic dose of saffron derived proteoglycan enhance macrophage mediated nitric oxide (NO) release cum phagocytosis (Escribano et al., 1999). Production of NO is an important cytotoxic function of macrophage to resolve infection of protozoan and bacterial parasites (De Groote and Fang, 1995). Intracellular signaling cascades alter nitric oxide synthase expression in macrophage upon activation and facilitate the release of NO by oxidizing terminal guanidine nitrogen of L-arginine (Raddassi et al., 1994; Dong et al., 1995; Park et al., 1996; Jun et al., 1996). We speculate that in the current study crocin could induce the

**Discussion**

We and other reported that *C. Sativus* and its major constituent crocin is an effective *in vitro* anti-proliferative agent of MCF-7 cells (Mousavi et al., 2011; Bakshi et al., 2012; Bizhan Malaekeh-Nikouei et al., 2013; Samarghandian and Borji, 2014). Initiation of apoptosis and proportion of MCF-7 cells attained apoptotic state after treatment with saffron has been well enumerated by flow cytometric analysis (Mousavi et al., 2009; Mousavi et al., 2011). However, the precise role of CSE or crocin on pro-apoptotic, anti-apoptotic, and caspase signaling mediated MCF-7 cell death at molecular level is ambiguous. Bcl-2 family includes key regulators of apoptosis and the molecule is over-expressed in many types of cancer cells (Llambi and Green, 2011). While reduced Bcl-2 expression accompanied with high expression of Bax may promote apoptotic response to anticancer drugs, increased expression of Bcl-2 leads to resistance to chemotherapeutic drugs (Ouyang et al., 2012). In this study, we found that CSE and crocin down regulate the expression of Bcl-2 with simultaneous up regulation of pro apoptotic Bax in MCF-7 cells (Figure 5). Upon receiving death signal in form of ligands (Fas, Death receptor 3, 4, 5, and TRAIL) or drugs, pro-apoptotic proteins such as caspase 3, 8 and 9 undergo post-translational modifications that include dephosphorylation and cleavage leading to their activation and formation of death inducing signaling complex and translocated to mitochondria where apoptosis initiated (Shamas-Din et al., 2011). This phenomenon led us to determine consequent signaling cascade of CSE and crocin mediated Bcl-2 down regulation in MCF-7 cells. To address this issue, we determine expression of pro caspase 3, 8 and 9. Treatment of MCF-7 cells with 10 µg/ml of CSE and crocin leads to up regulation of caspase 3, 8 and 9 in time dependent manner (Figure 7). These findings suggest that the CSE and crocin induced apoptosis in MCF-7 cells via the Bcl-2 down regulation and caspase 3 dependent pathways. Biochemical signatures accompanied with apoptosis include chromosomal DNA cleavage into inter-nucleosomal fragments, phosphatidylserine externalization and a number of intracellular substrate cleavages by specific proteolysis (Cohen et al., 1994; Martin and Green, 1995). As apoptosis occurs when DNA damage is irreparable, we tested the efficacy of CSE and crocin on MCF-7 cells DNA fragmentation. Each concentration of CSE and crocin treatment elicits considerable DNA fragmentation after 24 hours (Figure 6). Taken together this data evidenced that CSE and crocin modulating the expression of pro apoptotic markers in MCF-7 cells along with severe DNA damage and thus elicits significant cell death.

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expression of nitric oxide synthase and subsequent NO production in macrophage and thus enhanced phagocytosis of yeast ex vivo.

In conclusion, our study delineates the involvement of pro apoptotic and caspase mediated MCF-7 cell death by CSE and crocin at molecular level accompanied with extensive DNA damage. Further we found that normal swiss albino mice can tolerate the maximum dose of CSE. Crocin enhance macrophage activation and its phagocytic ability ex vivo. However the detailed molecular mechanism studies are required to translate this formulation in to clinic.

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