Hesperidin, A Citrus Bioflavonoid Attenuates Iron-Induced Biochemical Oxidative Stress in Mouse Liver

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Received: July 31, 2018; Published: August 17, 2018

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Abstract

Iron is an essential element, which is abundant in the red blood cells and plays a key role during oxidative phosphorylation in the mitochondria of all cells. The excess of iron has been indicated in several diseases as it induces oxidative stress. The present study was undertaken to understand the modulation of iron induce oxidative stress in mice by hesperidin. The mice were administered with 250 mg/kg body weight of hesperidin for five days before feeding with 5000, 10,000 and 20,000 ppm FeCl₃ for 30 days. The glutathione, glutathione-s-transferase, catalase, superoxide dismutase, lipid peroxidation, lactate dehydrogenase, aspartic acid transaminase, and alanine aminotransaminase levels were estimated in the liver of iron treated mice after 30 days post-iron treatment. The feeding of different concentrations of iron for 30 days led to an increase in the oxidative stress as indicated by a decline in the glutathione concentration, and glutathione-s-transferase, catalase, and superoxide dismutase activities accompanied by the rise in the lactate dehydrogenase, aspartic acid transaminase, and alanine aminotransaminase levels.

Treatment of hesperidin for 5 days before iron overload elevated the activities of glutathione-s-transferase, catalase, and superoxide dismutase and glutathione concentration, whereas the activities of lactate dehydrogenase, aspartic acid transaminase, and alanine aminotransaminase and lipid peroxidation declined significantly. Our study demonstrates that pre-treatment of hesperidin for five days reduced the iron induced oxidative stress indicated by the reduction in lipid peroxidation, lactate dehydrogenase, aspartic acid transaminase, and alanine aminotransaminase activities and a rise in the glutathione-s-transferase, catalase, and superoxide dismutase and glutathione contents in the mice liver.

Keywords: Mice; Liver; Antioxidants; Lipid Peroxidation; Lactate Dehydrogenase; Aspartic Acid Transaminase; Alanine Aminotransaminase

Introduction

Iron is one of the most abundant common elements on Earth. Iron is a trace element and it is extremely important in carrying out various metabolic processes in all living organisms for their survival. It participates in a wide variety of metabolic processes, including oxygen transport, electron transfer, nitrogen fixation and DNA synthesis. Iron exists in two oxidative states, ferrous (Fe²⁺) and ferric (Fe³⁺), which can donate or accept electrons [1-3]. In addition, the iron can ligand up to six places that reduces it otherwise high reactivity. “Free” iron produces hydroxyl radicals from superoxide and hydrogen peroxide via Fenton chemistry inducing oxidative stress in the cells [4]. These free radicals produced by iron are highly reactive and they interact with proteins, DNA and lipid inflicting severe damage to these biomolecules and subsequently tissue injury [5]. Cell has got inbuilt mechanism to protect itself from oxidative stress-induced by iron. The iron is oxidized to Fe³⁺, which tightly combines with transferrin and is kept in a non-toxic redox-inert state [6].

In addition, two-thirds of serum transferrin exists as apo transferrin and will quickly capture the free iron which is released from the cell whenever required [7]. Iron and iron salts are poorly water soluble however, it is an essential micronutrient, which is required for numerous biological reactions including mammalian cell growth, heme and DNA biosyntheses, oxygen transport, cell proliferation, cytochrome p450 enzyme activity, and hypoxic response reactions [8]. Iron is essential constituent of numerous enzymes, including iron-sulphur and heme proteins of the respiratory chain, as well as ribonucleotide reductase, which catalyses the rate-limiting step in DNA synthesis [9,10]. It is common prosthetic group composed of protoporphyrin IX and a Fe (II) ion. Presence of iron is a double-edged sword, it is extremely indispensable to carry various important metabolic processes at the same time its slight excess induces oxidative stress due to production of free radicals, especially the hydroxyl radical which is highly reactive and provoke critical and even irreversible cell injury. Iron deficiency also exerts negative effects on the cells [8].

The mismanagement of iron in the cells is the principle cause of underlying pathogenic event in oxidative stress that is harbinger of several diseases [11,12]. The presence of excess iron in the body is referred to as iron overload. Iron –induced oxidative stress may be of several types however, the most common being the genetic. Apart from the genetic cause the presence of excess iron in the diet
the body content of iron is regulated primarily by absorption because humans have no physiological mechanism to eliminate excess iron. Accumulation of excess iron in the body may be highly toxic as it may lead to a variety of serious conditions including aging, cardiovascular diseases, gastrointestinal tract disorders, diabetes, cataractogenesis, degenerative retinal damage, autoimmune nephrotic syndromes, heavy metal toxicity, Parkinson’s disease, Alzheimer’s disease, bronchopulmonary dysphasia, ischemia reflow states, kidney damage, macular degeneration, stroke and cancer [17-21]. The tissue damage associated with iron overload is believed to result basically from free radical reactions mediated by iron as iron is an effective catalyst in free-radical production [22]. The natural products may be of great significance in preventing or curing iron-induced toxicity and disorders.

Hesperidin is a bioflavonoid (flavonoids have diphenol structure with the molecular formula C_{27}H_{34}O_{16} and is found in vegetables, herbs, fruits and legumes. Nearly 5000 kinds of flavonoids are known) abundantly present in citrus fruits. Its aglycone form is called hesperitin. The hesperidin derives its name from the word “hesperidium”, the kind of fruit produced by citrus trees. The French chemist Lebreton (1828) was first to isolate hesperidin from the white inner layers of citrus peels. Hesperidin is believed to play a role in plant defence against fungal and bacterial invasions [23]. The sweet oranges (Citrus sinensis) and tangelos contain very high amount of hesperidin. The orange juice that is without pulp [24]. Hesperidin exerts a variety of pharmacological actions including anti-inflammatory, antioxidant, free radical scavenging and antiulcer effects. It also inhibits selected cytochrome p450 enzymes resulting in drug interactions [25,26]. Hesperidin increases the strength of the capillaries (blood vessels) and regulates their permeability.

It assists vitamin C in maintaining the collagen in healthy condition and hesperidin also helps in the proper absorption of vitamin C. It prevents vitamin C from being destroyed in the body by oxidation; beneficial in hypertension; helps haemorrhages and in conditions of ruptured capillaries and connective tissues and builds a protective barrier against infections [27]. Hesperidin has been reported to be analgesic, anti-inflammatory, antiallergenic, anticancerous, antihypotensive, antimicrobial and vasodilators [28-32]. The consumption of 500 ml orange juice for 4 weeks has been reported to modulate the expression of 3422 genes whereas only hesperidin intake led to the modulation of 1819 genes studied by microarrays [33]. Hesperidin has been found to be active against oxidative stress, hypotension, nitric oxide synthase inhibition, neurotoxicity, Herpes simplex virus infection, apoptosis, inflammatory bowel disease, arthritis, atherogenesis platelet and erythocyte aggregation and infection [34-44]. The hesperidin has been reported to scavenge free radicals and accelerate wound healing of irradiated wounds. It was found to be non-toxic up to 2g/kg body weight in mice orally [45,46]. The sub chronic administration of 5% hesperidin for 13 weeks has been found to be non-toxic in mice [47]. The liver is an important organ that metabolizes iron therefore the present study was undertaken to obtain an insight into the effect of hesperidin on the iron-induced oxidative stress in the mouse liver.

**Materials and Methods**

**Chemicals**

Ferric chloride, Hesperidin, di-sodium hydrogen phosphate (Na_{2}HPO_{4}), hydrogen peroxide (H_{2}O_{2}), di-potassium hydrogen phosphate (K_{2}HPO_{4}), potassium di-hydrogen phosphate (KH_{2}PO_{4}) were procured from MERCK, India Ltd., Mumbai, whereas 1-chloro-2,4 dinitrobenzene (CDNB), phenazene methosulphate, nitroblue tetrazolium, sodium pyruvate, cumene hydroperoxide, thio-barbituric acid, 5,5’-dithiobis[2-nitrobenzoic acid, 5-thionitrobenzoic acid, mM tert-butyl-hydroperoxide and nicotinamide adenine dinucleotide (NADH) were supplied by the Sigma Aldrich Chemical Company, Kolkata, India.

**Animal care and Handling**

The animal care and handling were done according to the guidelines issued by the World Health Organization, Geneva, Switzerland and the INSA (Indian National Science Academy, New Delhi, India). Usually, six to eight-week-old inbred Swiss albino mice weighing 25 to 30 g of either sex were selected from the inbred colony maintained under the controlled conditions of temperature (23 ± 2°C), humidity (50 ± 5%) and 12 h of light and dark cycle, respectively. The animals had free access to the standard food and water. Five animals were housed in a polypropylene cage containing wood powder (procured locally) as bedding throughout the experiment. The Animal Ethics Committee of Mizoram University, Aizawl, India had approved the study.

**Preparation of drug and mode of administration**

The required amount of hesperidin was dissolved in 1% CMC in distilled water and the animals were administered with 250 mg/kg body weight of hesperidin (HPD) orally through an oral gavage.

**Experimental**

The effect of hesperidin on iron-induced oxidative stress in the mouse liver was studied by dividing the animals into the following groups:

a) **Iron**: The animals of this group were administered with 0, 5000, 10,000 and 20,000 ppm of ferric chloride in ordinary drinking water daily for 30 days consecutively.

b) **HSP+iron**: Animals of this group were orally administered with 250 mg/body weight of hesperidin intraperitoneally for five days before administering the animals with 0, 5000, 10,000 and 20,000 ppm of ferric chloride in ordinary drinking water daily up to 30 days.

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Thirty days after the administration of Ferric chloride, the animals from both the groups were killed by cervical dislocation. The animals were dissected and perfused with ice cold saline transcardially. The liver from both group of animals was removed and blot dried. A total of 80 mice were used to complete this experiment.

**Preparation of Homogenate**

The liver was weighed, and 10% homogenate was prepared in a phosphate buffer saline using an electrical homogenizer (REMI India). Various biochemical parameters were estimated in the liver homogenate after 30 days of iron treatment.

**Total Proteins**

The total proteins were assayed by the modified method of Lowry [48].

**Glutathione**

The glutathione (GSH) contents were estimated by earlier described method [49] with minor modifications. Briefly, the precipitation of proteins was carried out with 0.5 ml ice cold 10% S-sulfosalicylic acid and kept on ice for 10 min and centrifuged (Sorvall Instruments RC5C, DuPont, Minnesota, USA) at 4 °C for 15 min at 15,000 rpm in cold. The protein free supernatant was collected and immediately mixed with 0.5 ml of NADPH (4 mg of reduced form was dissolved in 100 ml of 0.5% NaHCO₃), 0.5 ml of glutathione reductase (6 units/ml in 0.1 M phosphate buffer, pH 7.0) and 1 ml of 0.6 M DTNB [prepared in 0.2 M phosphate buffer (pH 8)]. The formation of TNB was read against the blank in a UV-Visible double beam spectrophotometer (Shimadzu Corporation, Tokyo, Japan) at 412 nm. A sample lacking GSH was used as a blank. The GSH concentration has been expressed as µmol/mg protein. A standard curve was plotted using different concentrations of GSH.

**Glutathione-S-Transferase**

The protocol described by Habig and Pabst [50] was used to measure the activity of glutathione-S-transferase (GST), where the homogenate was mixed with 0.1 M potassium phosphate buffer, CDNB and 10 mM GSH, and kept in water bath for 10 min at 37 °C. The absorbance was recorded at 1 min intervals against the distilled water blank at 340 nm using a UV-VIS spectrophotometer. The GST activity has been expressed as nmol/mg of protein.

GST activity = Absorbance of sample – Absorbance of blank × 1000/9.6 × Vol of sample.

**Catalase**

The catalytic reduction of hydrogen peroxide was used to measure the catalase activity [51], where, hydrogen peroxide was added to the tissue homogenate and the mixture was incubated at 37 °C. The hydrogen peroxide decomposition was monitored at 0.5s, 10s intervals up to 30s and the absorbance was read against the phosphate buffered blank at 240 nm using a UV-VIS spectrophotometer. The average difference in absorbance in 30 s was calculated.

**Superoxide Dismutase**

The activity of superoxide dismutase (SOD) was determined as described earlier [52]. Briefly, the homogenate was mixed with phenazine methosulphate, nitroblue tetrazolium and NADH and incubated at 30 °C for 90 sec. The reaction was terminated by the addition of acetic acid and n-butanol. Blank was prepared in a similar fashion without the sample and the reaction was stopped by adding acetic acid and n-butanol. The sample absorbance was read against the blank at 560 nm in a UV-VIS spectrophotometer.

**Lipid Peroxidation**

The induction of lipid peroxidation (LOO) was measured as various thiobarbituric acid reactive substances (TBARS) including malondialdehyde, lipid hydroperoxides and aldehydes in the homogenate [53]. The liver homogenate was heated with thiobarbituric acid (0.8%), sodium dodecyl sulphate (0.1%) and acetic acid (20%) in a boiling water bath for 30 min (lipoproteins get precipitated). The resultant mixture was cooled, extracted with n-butanol-pyridine, and the absorbance of the butanol layer was recorded at 532 nm using UV-Visible double beam spectrophotometer. The resultant concentration of TBA reactive substances is expressed as nmol/mg protein obtained from a standard curve of tetraethoxypropane.

**Estimation of Lactate dehydrogenase, Aspartate transaminase and Alanine amino transaminase**

The activities of lactate dehydrogenase, aspartate transaminase and alanine amino transaminase were measured using commercially available Respons kits according to the protocol provided by the manufacturers using a Respons 910 autoanalyzer (Diagnostic Systems GmbH, Holzheim, Germany).

**Statistical Analysis**

The significance between the treatments was determined using the Student’s ‘t’ test and one-way ANOVA with Tukey’s post-hoc test. A p value of <0.05 was considered statistically significant. The Solo 4 statistical package (BMDP Statistical Software Inc, Los Angeles, CA, USA) was used for statistical analyses.

**Results**

The results of all biochemical analyses are represented as mean ± standard error of the mean in Tables 1-8 and (Figures 1-8).

**Table 1**: Alteration in the Iron-induced glutathione concentration (µmol/mg protein) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron          | Hesperidin + Iron   |
|------------|---------------|---------------------|
| 0          | 14.10±0.18    | 11.6±0.22           |
| 5000       | 10.61±0.13**  | 12.02±0.23**        |
| 10000      | 10.42±0.12**  | 10.84±0.16          |
| 20000      | 10.27±0.08**  | 10.60±0.13          |

Level of significance: *P<0.05; **P<0.01; ***P<0.001; ****P<0.0001. When Iron group compared with Hesperidin + Iron group.

*P<0.001; **P<0.0001, Iron group compared to control.
Table 2: Alteration in the Iron-induced glutathione-s-transferase activity (nmol/mg protein) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 5.63±0.41 | 4.82±0.14       |
| 5000       | 3.06±0.06* | 3.7±0.17*       |
| 20000      | 2.94±0.05* | 3.42±0.12*       |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.01; **P<0.001, Iron group compared to control.

Table 3: Alteration in the Iron-induced catalase activity (nmol/mg protein) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 86.36±3.29 | 86.33±1.79       |
| 5000       | 32.23±0.25** | 69.68±3.42c       |
| 20000      | 28.15±1.36** | 59.74±2.09c       |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.001, Iron group compared to control.

Table 4: Alteration in the Iron-induced superoxide dismutase activity (nmol/mg protein) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 54.35±0.44 | 51.04±0.33       |
| 5000       | 27.79±0.93** | 38.48±0.92c       |
| 20000      | 15.18±0.58** | 34.99±0.18c       |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.001, Iron group compared to control.

Table 5: Alteration in the Iron-induced lipid peroxidation (nmol/mg protein) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 9.5±0.12 | 6.9±0.11       |
| 5000       | 21.55±0.66** | 18.27±0.67c       |
| 20000      | 10.0±0.73** | 19.83±0.17*       |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.0001, Iron group compared to control.

Table 6: Alteration in the Iron-induced lactate dehydrogenase activity (U/L) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 132.06±2.06 | 141.3±5.15       |
| 5000       | 176.77±1.81** | 158.88±2.06**     |
| 20000      | 290.26±2.48** | 196.71±2.48b     |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.001, Iron group compared to control.

Table 7: Alteration in the Iron-induced Aspartate transaminase activity (U/L) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 57.34±2.28 | 46.25±0.97c       |
| 5000       | 84.99±2.10* | 65.37±0.7c       |
| 20000      | 111.63±1.86** | 86.41±2.82c       |

Level of significance=**P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.001, Iron group compared to control.

Table 8: Alteration in the Iron-induced Alanine aminotransferase activity (U/L) by 250 mg/kg body weight of hesperidin in mouse liver 30 days after treatment with iron.

| FeCl₃ (PPM) | Iron | Hesperidin + Iron |
|------------|------|------------------|
| 0          | 46.44±1.29 | 44.61±0.97c       |
| 5000       | 69.25±1.58** | 65.58±1.22       |
| 20000      | 75.97±2.13** | 69.25±0.89 youthful |

Level of significance= **P<0.001; P<0.01; P<0.001; P<0.0001 When Iron group compared with Hesperidin + Iron group.
*P<0.001; **P<0.0001, Iron group compared to control.

**Glutathione**

The estimation of glutathione in liver revealed a spontaneous level of 14.10±0.18 µmoles/mg protein in control animals and administration of hesperidin alone did not reduce GSH concentration significantly when compared to non-drug treated control (Table 1). The chronic administration of ferric chloride for four weeks led to a dose dependent and significant attrition in the GSH concentration when compared to untreated control (Table 1) and (Figure 1). The Hesperidin treatment before iron overload elevated the concentration of glutathione significantly (Table 1) and (Figure 1).
Biomedical Journal of Scientific & Technical Research

Volume 8 - Issue 1: 2018

Cite this article: Jagetia GC, T Lalrinpuii. Hesperidin, A Citrus Bioflavonoid Attenuates Iron-Induced Biochemical Oxidative Stress in Mouse Liver. Biomed J Sci&Tech Res 8(1)- 2018. BJSTR MS.ID.001602. DOI: 10.26717/BJSTR.2018.08.001602.

Figure 1: Effect of 250 mg/kg body weight hesperidin on the glutathione concentration in mouse liver subjected to iron overload. Red bars: Iron alone and Blue bars: Hesperidin+Iron.

Figure 2: Effect of 250 mg/kg body weight hesperidin on the glutathione-s-transferase activity in mouse liver subjected to iron overload. Red bars: Iron alone and Blue bars: Hesperidin+Iron.

Figure 3: Effect of 250 mg/kg body weight hesperidin on the catalase activity in mice liver exposed to iron overload. Red Bars: Iron and Blue Bars: Hesperidin+Iron.
Figure 4: Effect of 250 mg/kg body weight hesperidin on the superoxide dismutase activity in mice liver exposed to iron overload. Red Bars: Iron and Blue Bars: Hesperidin+Iron.

Figure 5: Effect of 250 mg/kg body weight hesperidin on the iron induced lipid peroxidation in mouse liver. Red bars: Iron and Blue bars: Hesperidin+Iron.

Figure 6: Effect of 250 mg/kg body weight hesperidin on the lactate dehydrogenase activity in mice liver exposed to iron overload. Red Bars: Iron and Blue Bars: Hesperidin+Iron.
Glutathione-S-Transferase
The spontaneous glutathione-s-transferase activity has been 5.63±0.41 nmol/mg protein and administration of hesperidin alone did not alter this activity significantly when compared to non-drug treated control. The chronic administration of ferric chloride for four weeks reduced the glutathione-s-transferase activity significantly in a dose dependent manner and the maximum decline was observed for 20000 ppm iron overload (Table 2) and (Figure 2). Hesperidin treatment before iron overload elevated the glutathione-s-transferase activity significantly and the maximum elevation in the GST activity was observed for hesperidin administered in mice before feeding of 5000 ppm iron. With increasing iron overload, the increase in GST activity by hesperidin was lesser (Table 2) and (Figure 2).

Catalase
The spontaneous catalase activity is 86.360±3.29 nmol/mg protein and administration of hesperidin alone did not alter this activity significantly when compared to non-drug treated control. The chronic administration of ferric chloride for four weeks reduced the catalase activity significantly in a dose dependent manner and the maximum decline in the catalase activity was observed for 20,000 ppm iron overload (Table 3) and (Figure 3). Hesperidin treatment before iron overload elevated the catalase activity significantly (Table 3) and (Figure 3). This rise in catalase activity was more than two folds for all iron concentrations (Table 3).

Superoxide Dismutase
The spontaneous superoxide dismutase activity has been 54.35±0.44 nmol/mg protein and administration of hesperidin alone did not alter this activity significantly when compared to non-drug treated control. The chronic administration of ferric chloride for four weeks resulted in a drastic decline in the superoxide dismutase activity, which was dose dependent (Table 4). This decline in SOD activity was significant and a maximum decline was
observed for 20,000 ppm iron overload, where it was greater than four folds in comparison to spontaneous SOD activity (Table 4) and (Figure 4). Hesperidin treatment before iron overload significantly elevated the superoxide dismutase activity and it was two folds higher than the iron treatment alone, except for 5000 ppm (Table 4) and (Figure 4).

**Lipid Peroxidation**

The results of present study showed that the levels of spontaneous lipid peroxidation in the liver of mouse has been 6.95 ±0.02 nmol/mg proteins (Table 5) and (Figure 5). The chronic administration of ferric chloride for four weeks raised the lipid peroxidation significantly when compared to untreated control (Table 5). This increase in lipid peroxidation after iron treatment was greater than two folds and dependent on the dose of iron treatment. The administration of hesperidin alone treatment did not increase the level of lipid peroxidation which was almost similar to the spontaneous level (6.9±0.11 nmoles /mg protein). Administration of hesperidin for five days before chronic ferric chloride treatment significantly reduced lipid peroxidation when compared to iron loaded group (Table 5) and (Figure 5).

**Lactate Dehydrogenase (LDH)**

The spontaneous LDH activity is 132.06±3.33 (IU/L) and administration of hesperidin alone marginally alleviated its activity when compared to non-drug treated control. The chronic administration of ferric chloride for four weeks increased the LDH activity significantly in a dose dependent manner and the maximum rise was observed for 20000 ppm iron overload (Table 7) and (Figure 7). This rise in LDH activity was more than two folds for 10,000 and 20,000 ppm iron treatment (Table 7). Hesperidin treatment for 5 consecutive days before iron overload reduced LDH activity significantly (Table 6) and (Figure 6).

**Aspartate Transaminase (AST)**

The spontaneous AST activity is 57.34±2.39 (IU/L) and administration of hesperidin alone significantly reduced its activity when compared to non-drug treated control. The chronic administration of ferric chloride for four weeks increased the AST activity significantly in a dose dependent manner and the maximum rise was observed for 20,000 ppm iron overload (Table 7) and (Figure 7). Hesperidin treatment before iron overload reduced the AST activity significantly (Table 7) and (Figure 7).

**Alanine Aminotransaminase(ALT)**

The spontaneous ALT activity is 48.44±1.29 (IU/L) and administration of hesperidin alone significantly reduced its activity when compared to non-drug treated control (Table 8). The chronic administration of ferric chloride for four weeks increased the ALT activity significantly in a dose dependent manner and the maximum rise was observed for 20,000 ppm iron overload (Table 8) and (Figure 8). Hesperidin treatment before iron overload significantly reduced the ALT activity (Table 8) and (Figure 8).

**Discussion**

The iron plays an important role in cellular metabolism due to its transition ability from Fe²⁺ into Fe³⁺. Despite the fact that iron is essential in several metabolic functions including energy production it does not exist in free form and it is always found in bound form with proteins [22]. It is an essential micronutrient that regulates oxygen transport in haemoglobin and plays a crucial role in energy production [54]. The total human body iron ranges from 3 to 5 g and greater than this amount leads to different disorders including anemia, bronchopulmonary dysphasia, cardiovascular and gastrointestinal tract dysfunction, cataract, diabetes, degenerative retinal damage, autoimmune nephritic syndromes, heavy metal toxicity, Alzheimer’s disease, Parkinson’s disease, ischemia reflow states, skeletal and respiratory muscle dysfunction and cancer [55-65]. The maximum iron is stored in the liver and the accumulation of excess iron induces oxidative stress related disorders, which needs to be alleviated to counter the iron induced oxidative stress. The natural products be of great value in this regard. Therefore, the present study was undertaken to evaluate the ability of hesperidin, a citrus flavanone to reduce the iron-induced oxidative stress in the liver of Swiss albino mice.

Glutathione is a tripeptide, which controls thiol redox reactions and maturation of extra-mitochondrial iron sulphur clusters. Glutathione insulates cytosolic function in iron metabolism by varying its concentration during redox stresses [66]. Glutathione is abundantly present in the cell and it participates in several redox reactions vital to the cell. It protects the cells from a host of endogenous and exogenous toxic insults [67]. The iron concentration is tightly regulated in the cells. However, excess iron triggers the cascade of free radical formation leading to increased oxidative stress [22]. This may be the cause for the dose dependent reduction in the glutathione concentration in the present study by iron overload. Earlier iron overload has been reported to reduce glutathione concentration in vitro [68,69]. Hesperidin treatment arrested the iron induced decline in the GSH concentration in mice liver. The studies regarding the use of hesperidin to protect against the iron-induced glutathione depletion are scanty. However, naringin another citrus flavanone has been reported to reduce iron induced glutathione concentration in vitro [68,69].

Glutathione-s-transferases are a group of detoxifying enzymes, found in all aerobic eukaryotes and they protect cells from the toxic insults by deactivating hydrophobic cytotoxic and genotoxic compounds [70]. The treatment of mice with different doses of iron for one month resulted in a dose dependent decline in the glutathione-s-transferase activity in the liver. A similar effect has been observed in Caucasian patients receiving iron therapy for the treatment of β-thalassemia, where GSTT1 and GSTM1 genes were downregulated [71]. This indicates that iron overload is detrimental to the glutathione-s-transferase activity due to the ability of iron to induce oxidative stress in the cells. The hesperidin administration has reduced the iron-induced decline in the glutathione-s-transferase activity. There are no reports regarding the use of hesperidin to protect against the iron induced attenuation in the glutathione-s-transferase activity in mouse liver. However, it has been reported to arrest iron-induced decline in glutathione-s-transferase activity in rat liver earlier [72].

Catalase or oxidoreductase (EC. 1.11.1.6) is a tetrameric enzyme, which is present in all organisms utilizing oxygen for...
It detoxifies $\text{H}_2\text{O}_2$, a strong oxidizing agent involved in tissue pathogenesis by converting it into water and molecular oxygen [73]. The treatment of mice with different doses of iron reduced the activity of catalase in mice liver. A similar effect has been observed in mice liver fed with iron overload diet for four months [74]. Earlier the iron overload has been reported to reduce catalase activity in rat liver [72]. Similarly, iron overload has been reported to attenuate the catalase activity in vitro [68,69]. The hesperidin treatment arrested this decline in the catalase activity by elevating the activity of catalase in mice liver significantly. A similar increase in catalase activity has been reported in rat liver administered with hesperidin earlier [72]. The naringin, a citrus bioflavonoid has been reported to increase the catalase activity in HepG2 cells and isolated liver mitochondria in vitro [68,69].

Superoxide dismutases are ubiquitously present in all organisms that use oxygen for their energy requirement. The aerobic organisms produce superoxide radical during respiration, which is not highly reactive, but it becomes reactive with transition metal complexes of iron, copper and manganese leading to increased oxidative stress [75,76]. The superoxide dismutase triggers the dismutation of O$_2^-$ into less toxic hydrogen peroxide and acts as first line of defence against oxidative stress [75,76]. It also inactivates nitric oxide reducing oxidative stress. The iron overload has reduced the activity of superoxide dismutase in a dose dependent manner in the present study. A similar effect has been reported in rat and mice liver earlier [72,74]. Hesperidin treatment for five days led to an increase in the superoxide dismutase activity in mice liver in the present study. A similar effect has been observed in iron overloaded rat liver treated with hesperidin [72]. Similarly, naringin treatment has been reported to elevate the superoxide dismutase activity in iron overloaded HepG2 cells and isolated mice liver mitochondria in an earlier study [68,69].

The inactivation of hydroxyl and hydroperoxyl radicals leads to the formation of lipid peroxidation. hydroxyl radicals are formed due to the redox cycling of Fe$^{2+}$ via Fenton reaction, where it reacts with $\text{H}_2\text{O}_2$ to produce hydroxyl radical. The Haber Weiss reaction regenerates Fe$^{2+}$ when superoxide radical reacts with Fe$^{3+}$ [77,78]. Liver being the primary site for iron accumulation, it is at an increased risk of iron overload that induces oxidative stress. The increased iron overload has been indicated in liver cancer [79]. Therefore, it is pertinent to assay lipid peroxidation in the iron overloaded mice.

The iron overload has increased the lipid peroxidation in the mouse liver drastically. These observations are in conformation with our earlier reports where iron overload has been reported to accelerate lipid peroxidation in vitro [68,69]. Similarly, an increase in lipid peroxidation has been observed in mice and rat liver after iron overload [72,74,79]. Hesperidin administration for five consecutive days has retarded lipid peroxidation significantly. A similar effect has been observed in iron overloaded rat liver treated with hesperidin [72]. Likewise, naringin has been reported to reduce lipid peroxidation in the HepG2 cells and isolated mice liver mitochondria in vitro [68,69].

The lactate dehydrogenase is a glycolytic pathway enzyme that catalyzes the conversion of lactate to pyruvic acid in the presence of NAD$^+$ [80]. The over production of lactate dehydrogenase is hallmark of liver toxicity/damage and organ failure [81,82]. Similarly, the augmented activities of aspartic acid transaminase and alanine aminotransaminase are indicative of liver damage. The iron overload for 30 days has elevated the activities of lactate dehydrogenase, aspartic acid transaminase and alanine aminotransaminase drastically indicative of liver injury. Iron overload has been reported to increase lactate dehydrogenase, aspartic acid transaminase and alanine aminotransaminase in human serum, mice and rat liver [72,74,83,84]. Hesperidin treatment for five consecutive days resulted in a significant reduction in lactate dehydrogenase, aspartic acid transaminase and alanine aminotransaminase in the iron overloaded mice liver indicating that hesperidin pretreatment protected mouse liver against the iron-induced insult. A similar effect has been observed in rat liver treated with hesperidin earlier [72]. Diosmin has been reported to reduce lactate dehydrogenase, aspartic acid transaminase and alanine aminotransaminase in iron loaded rat liver [84].

The exact mechanism of action of hesperidin to reduce the iron induced oxidative stress is not known. Hesperidin may have employed multiple mechanisms to reduce the iron-induced oxidative stress in mice liver.

The iron is known to generate free radicals via Fenton and Haber Weiss reactions during different metabolic processes [77,78]. The presence of hesperidin before iron overload would have inhibited the generation of iron-induced free radicals leading to alleviation in iron-induced oxidative stress in the mice liver. Hesperidin has been found to suppress various free radicals [45]. The iron overload has been reported to activate IKK$\beta$ leading to transactivation of NF-kB, and TNF-$\alpha$ in Kuffer cells of liver [85]. In addition, iron has been also reported to activate COX-II and prostaglandins, which triggers inflammatory cascade leading to pathophysiology [86]. Hesperidin has been reported to suppress the transactivation of NF-kB and COX-II [87,88], which may have also contributed at molecular level to alleviate oxidant status in hesperidin pre-treated group. The transcription activation of Nrf2 inhibits iron accumulation in the mouse liver, the major site of iron storage and metabolism. The hesperidin administration in rats has been reported to stimulate the Nrf2 activation in their kidney [89]. A similar effect cannot be ruled out in the present study in mouse liver.

Conclusions

The present study demonstrates that hesperidin treatment reduced iron-induced biochemical toxicity in the liver attenuating, lipid peroxidation, lactate dehydrogenase, aspartic acid transaminase and alanine aminotransaminase in the mouse liver accompanied by elevated glutathione, glutathione-s-transferase, catalase and superoxide dismutase levels. This action of hesperidin seems to be mediated by free radical inhibition and suppression of NF-kB, COX-II, prostaglandins and TNF-$\alpha$ activation. The activation of Nrf2 signaling pathway may have lead to increased antioxidant status thereby reducing the iron-induced oxidative stress. Our
study indicates the utility of hesperidin to reduce iron-induced oxidative stress in clinical situation.

Acknowledgement
This work was supported by a grant no. F4-10/2010(BSR) UGC from the University Grants Commission, New Delhi, India.

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Cite this article: Jagetia GC, T Lalrinpuii. Hesperidin, A Citrus Bioflavonoid Attenuates Iron-Induced Biochemical Oxidative Stress in Mouse Liver. Biomed J Sci & Tech Res 8(1)- 2018. BJSTR MS.ID.001602. DOI: 10.26717/BJSTR.2018.08.001602.