In Vivo Antioxidant Activity of Topical Cream of Cassia tora L. Leaves Extract

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Objective. The aim of study was to determine the in vivo antioxidant activity of newly formulated O/W cream of methanolic extract of Cassia tora L. leaves. Methods. Oil in water (O/W) creams (0.05%, 0.1%, and 0.2%), cream base, methanolic extract of C. tora L. leaves (CTM), and standard (0.05% tretinoin cream) were screened for in vivo antioxidant activity. The ultraviolet- (UV-) B-induced rats were treated with different standard, O/W creams, cream base, and methanolic extract of Cassia tora L. leaves (CTM). The parameters like lipid peroxidation (LPO), reduced glutathione (GSH), superoxide dismutase (SOD), and catalase (CAT) levels were assessed. Results. The result of acute dermal toxicity shows that the creams were safe up to 2000 mg/kg (topically). Exposure of ultraviolet light of medium wave (UV-B light) decreased the level of GSH, CAT, and SOD and increased the LPO level. Concluding Remarks. We conclude that topical O/W creams of C. tora L. prevent the oxidative stress induced in rats by exposure to UV-B light by virtue of its in vivo antioxidant property, and these findings help to understand the mechanism of the antipsoriatic activity of O/W creams of C. tora L. in UV-B-induced psoriasis in the rat.

1. Introduction

N. P. Kansara found a potent antipsoriatic activity of O/W creams of Cassia tora L. in ultraviolet- (UV) -B-induced psoriasis in the rat. Aerobic organisms produce reactive oxygen species during normal respiration and inflammatory conditions. Exercise can create an imbalance between oxidant and antioxidant levels, a situation known as oxidative stress. Indeed, oxidative stress resulting from acute exercise in unadapted and adapted subjects has been proposed to damage enzymes, protein receptors, lipid membranes, and DNA [1]. The plant C. tora L. (Caesalpiniaceae) is traditionally claimed to be useful in the treatment of psoriasis and other skin diseases [2, 3]. C. tora L. leaves enriched in glycosides and also containing aloe-emodin may be beneficial for the skin diseases [4]. C. tora L. has been reported to possess a significant antioxidant activity in in vitro activity by using DPPH (2,2′-diphenyl-1-picrylhydrazyl) method [5]. In these antioxidant assays, the volatile oil from C. tora L. demonstrated 2,2′-diphenyl-1-picrylhydrazyl radical-scavenging activity in the concentration range from 20 to 500 μg/mL, with the 50% inhibitory concentration (IC50) value at 137 μg/mL; it also showed a significant inhibitory effect against hydroxyl radicals with an IC50 value of 67 μg/mL, lower than that of quercetin (IC50 = 8.15 μg/mL), but superior to that of 4-terpineol (IC50 = 87.5 μg/mL). We had used UV-B ray as a damage inducer because it causes skin to burn and directly damages DNA by interfering with its replication cycle [6]. And excessive exposure to UV-A radiation has its risks, which may cause premature aging, including wrinkles, sunspots, and loss of skin elasticity [7]. Hence, it was proposed to evaluate the efficacy of O/W creams and methanolic extract of C. tora L. leaves (CTM) to prevent the UV-B-induced oxidative stress.
2. Methodology

Standard (tretinoin: 0.05%) cream was obtained from Ethnor pharma (A Johnson & Johnson Div.), Mumbai, Maharashtra, India. Diethyl ether was obtained from Rankem, India. Other ingredients such as light liquid paraffin (Astron), cetostearyl alcohol (Chemdyes), propylene glycol (Nomex), white soft paraffin (Nomex), butyl hydroxytoluene (Rankem), benzyl alcohol (Chemdyes), disodium EDTA (Rankem), isopropyl myristate (FD, Fine Chemicals), and dibasic potassium phosphate (Rankem) were used to prepare O/W creams. Pyrogallol and hydrogen peroxide were obtained from SD Fine Chemicals Ltd., India. Thiobarbituric acid (TBA), trichloroacetic acid (TCA), 5,5′-dithiobis (2-nitrobenzoic acid) (DTNB), phosphate buffer, and Tris buffer were obtained from Sigma, USA. All reagents used were of analytical grade.

Leaves of C. tora L. (Cesalpinaceae) were collected from Dabhoi, Vadodara, Gujarat [8].

The methanolic extract was prepared by cold maceration method [8].

Formulation development was as follows. The extract was heated up to 70 ± 5°C. Aqueous phase consisting of water (q.s) was heated to the same temperature and then were added disodium EDTA (0.01%), butyl hydroxytoluene (0.001%), and dibasic potassium phosphate (0.2%) in it. Then CTM (0.05%, 0.1%, and 0.2%) was mixed in benzyl alcohol (1%) and added in it. Then, oily phase was added to the aqueous phase with continuous stirring at slow speed for 1 hour and slowly decreased the temperature and meanwhile was added isopropyl myristate (4%) in the mixtures of both phases. Allowed to cool at room temperature. Oily phase was consisted of light liquid paraffin (8%), cetostearyl alcohol (10%), propylene glycol (5%), and white soft paraffin wax (12%). The prepared creams were transferred into wide mouth containers and stored in cool place. Base was also prepared by the same previous method and with same ingredients but without CTM.

Evaluation of O/W creams was as follows.

Physical evaluation was done as per Singhal & Kansara, International Scholarly Research Network [8, 9]. Sensitivity test, irritatation test, and grittiness were performed as per Singhal & Kansara, International Scholarly Research Network [8].

Stability studies were done according to Singhal & Kansara, International Scholarly Research Network [8, 10].

Adult Wistar male rats (weight: approximately 300 g; age: 4–6 months) were used for the experiments. Animals were kept in the Shree Dhanvantary Pharmaceutical Analysis and Research Centre, Kim, Surat. After approval from the Institutional Animal Ethical Committee (Reg. no.1103/abc/07/cpcssea), rats were housed in polypolyethylene cages as 3 animals per cage with rice husk as the bedding material for 12 h light-dark cycle, at temperature of 22 ± 02°C, and humidity 30–70%. Rat pellet feed (Pranav agro Ltd.) and pure drinking water were supplied ad libitum. The animals were acclimatized to the laboratory conditions for a minimum period of seven days prior to commencement of treatment.

The acute dermal toxicity test was performed according to Singhal & Kansara, International Scholarly Research Network [8].

In vivo antioxidant studies were as follows. For the assessment of different formulations and crude extract on UV-B-induced oxidative stress in rats, the animals were treated with respective doses (topically, single dose) of different concentrations of O/W creams (Test 1, 0.05%; Test 2, 0.1%; and Test 3, 0.2%), standard (tretinoin: 0.05%), cream base, and crude extract during whole treatment. The animals were divided into seven groups as follow:

Group 1: Positive control received UV-B light,
Group 2: UV-B + standard,
Group 3: UV-B + Test 1,
Group 4: UV-B + Test 2,
Group 5: UV-B + Test 3,
Group 6: UV-B + cream base,
Group 7: UV-B + extract.

The procedure was as follows. Wistar rats were selected and divided into seven groups. Then hair of the dorsal skin was carefully shaved. Then test creams were applied topically on the dorsal part of the skin whichever was exposed to radiation. An area (1.5–2.5 cm) on one side of the flank was irradiated for 15 min (1.5 J/cm²) at a vertical distance of 20 cm with UV-B lamps. Then following parameters were assessed in the blood which was withdrawn from the retro-orbital plexus at the end of last treatment in the UV-B-induced psoriasis.

To evaluate lipid peroxidation (LPO), 2 mL of 5% suspension of separated RBC in 0.1 M phosphate-buffered saline and 2 mL of 28% trichloroacetic acid were taken in test tube and centrifuged. Then supernatant was separated in another test tube. Then 4 mL of supernatant was taken in test tube and 1 mL of 1% thiobarbituric acid was added in it. Then it was heated in boiling water for 60 minutes and cooled immediately. The absorbance was measured in UV spectrophotometer (Schimadzu 1601, Japan) at 532 nm [11–13].

Superoxide dismutase (SOD) was evaluated as follows. The erythrocyte lysate was prepared from the 5% RBC suspension of the blood. Then 50 μL of the erythrocyte lysate was taken in test tube and then 75 mM of Tris-HCl buffer (pH 8.2), 30 mM EDTA, and 2 mM of pyrogallol were added in it. Then absorbance was recorded at 420 nm for 3 min by UV spectrophotometer (Schimadzu 1601, Japan) [11–13].

Catalase (CAT) activity was determined in erythrocyte lysate using Aebi’s method with some modifications. 50 μL of the lysate was added into a cuvette containing 2 mL of phosphate buffer (pH 7.0) and 1 mL of 30 mM H₂O₂. Catalase activity was measured at 240 nm for 1 min using UV spectrophotometer (Schimadzu 1601, Japan) [11–13].

Reduced glutathione (GSH) was evaluated as follows. Blood glutathione was measured by addition of 0.2 mL of whole blood in 1.8 mL distilled water followed by 3.0 mL of precipitating mixture of 1.67 g metaphosphoric acid, 0.2 g EDTA, and 30 g NaCl to make 100 mL of solution. It was
centrifuged at 5000 \times g for 5 min and 1 mL of the filtrate was added to 1.5 mL of the phosphate solution, followed by the addition of 0.5 mL of DTNB reagent. The optical density was measured at 412 nm using UV spectrophotometer (Shimadzu 1601, Japan) [11–13].

Statistical analysis was as follows. All the experimental results were expressed as mean ± SEM. For statistical comparisons, exploratory probabilities were obtained by the one-way ANOVA followed by Dunnett’s multiple comparison tests using GraphPad Prism 5 (GraphPad software, Inc.). The intergroup difference was considered significant when $P \leq 0.05$.

For the evaluation of creams, three different concentrations of O/W creams (Test 1, 0.05%; Test 2, 0.1%; and Test 3—0.2%) were prepared to evaluate antisoriatic activity. Physical evaluation revealed that creams were having light green colour, characteristic odour, semisolid in nature, and pH ranges from 6.5 to 7. They passed the sensitivity test and irritation test. During stability study, no phase separation and liquefaction were observed.

The acute dermal toxicity test of creams was determined according to the OECD 402 (Organization for Economic Corporation and Development). The creams were safe up to the dose of 2000 mg/kg. There were no changes in fur, eyes, and behavior of treated animals as well as no toxic reactions were determined. And from results suitable doses (250 mg (0.05%), 500 mg (0.1%), and 1000 mg (0.2%)) were chosen for each activity in each cream for further in vivo studies.

3. Dataset Description

The dataset associated with this Dataset Paper consists of 4 items which are described as follows.
**Dataset Item 3 (Table).** Effect of different formulations on blood catalase in UV-B-induced psoriatic rat. In the table, each value represents Mean ± SEM, n = 6; “a” indicates P < 0.05; and “b” indicates P < 0.001, compared to positive control group. There was a marked depletion of catalase level in positive control group. Treatment with standard (tretinoin: 0.25 mg cream) had increased the level up to 145.17%. Treatment with different formulations and extract group were found to be 130.83% (UV-B + Test 2), 76.76% (UV-B + Test 3), and 124.71% (UV-B + extract), respectively, which were statistically significant (Figure 3).

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<th>Column 1: Group Number</th>
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<th>Column 3: Mean ± SEM Increase (%)</th>
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**Dataset Item 4 (Table).** Effect of different formulations on blood GSH in UV-B-induced psoriatic rat. In the table, each value represents Mean ± SEM, n = 6; and “b” indicates P < 0.001, compared to positive control group. There was a marked depletion of GSH level in positive control group. Treatment with standard (tretinoin: 0.25 mg cream) has increased the level up to 94.45%. Treatment with different formulations and extract group were found to be 56.42% (UV-B + Test 1), 89.46% (UV-B + Test 2), 71.77% (UV-B + Test 3), and 64.54% (UV-B + extract), respectively, which were statistically significant (Figure 4).

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**4. Concluding Remarks**

Free radical stress leads to tissue injury and progression of disease such as cancer, aging, ischemia, liver injury, arthritis, and Parkinson’s syndrome. Safer antioxidants suitable for long-term use are needed to prevent or stop the progression of free radical-mediated disorders [14]. Many plants possess antioxidant ingredients that provided efficacy by additive or synergistic activities. Some studies have shown that a number of plant products including polyphenolic substances and herb extracts exert potent antioxidant action. Some traditional natural antioxidants are already exploited commercially either as antioxidant additives or as nutritional supplements [15].

Superoxide dismutase mainly acts by quenching of superoxide, catalase by catalyzing the decomposition of hydrogen peroxide to water and oxygen. Glutathione reductase is a good scavenger of many free radicals like $O_2^-$, $HO_2$, and various lipid hydroperoxides and may help to detoxify many inhaled oxidizing air pollutants like ozone, $NO_2$, and free radicals in cigarette smoke in the respiratory tract [16]. Exposure of UV-B light leads to cellular changes, DNA damage, and also skin damage [17].

The present study has shown that exposure of UV-B light for 15 min not only damaged the skin but also produced oxidative stress in rats. The literature has documented free radical generation and DNA damage occurs during the exposure of UV-B light [17]. UV exposure, particularly UV-B rays, causes the generation of free radicals and related reactive oxygen species, which contribute to carcinogenesis by directly damaging cellular macromolecules, including DNA [18]. The level of the markers of oxidative stress, observed in UV-B-induced rats, substantiates the possibility of extensive generation of free radicals. It is further observed that application of different concentrations of O/W creams and CTM prevented the UV-B-induced oxidative stress parameters, and the effect was comparable to that of standard.
Hence, the observed *in vivo* antioxidant activity of different concentrations of O/W creams and CTM was substantiation of its earlier reported activity in *in vitro* studies [5].

Antioxidants act as free radical scavengers that destroy single oxygen molecules (free radicals) in the body, thereby protecting against oxidative damage of cells. SOD, catalase (CAT), GSH, and so forth are the well-known enzymes present in plasma which act as antioxidants by transforming reactive oxygen species and reactive nitrogen species into the stable compounds and involved in a scavenging of the excessive free radicals. The restoration of blood SOD, CAT, and GSH levels by the treatment with test creams is indicating that the inbuilt protective mechanism is being restored.

**Dataset Availability**

The dataset associated with this Dataset Paper is dedicated to the public domain using the CC0 waiver and is available at http://dx.doi.org/10.7167/2013/243949/dataset.

**References**


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