Abstract

Jojoba [Simmondsia chinensis (Link 1822) Schneider 1907] is an arid perennial shrub grown in several American and African countries. Jojoba seeds, which are rich in liquid wax, were used in folk medicine for diverse ailments. In the current study, the potential anti-inflammatory activity of jojoba liquid wax (JLW) was evaluated in a number of experimental models. Results showed that JLW caused reduction of carrageenin-induced rat paw oedema in addition to diminishing prostaglandin E2 (PGE2) level in the inflammatory exudates. In a test for anti-inflammatory potential utilizing the chick’s embryo chorioallantoic membrane (CAM), JLW also caused significant lowering of granulation tissue formation. Topical application of JLW reduced ear oedema induced by croton oil in rats. In the same animal model, JLW also reduced neutrophil infiltration, as indicated by decreased myeloperoxidase (MPO) activity. In addition, JLW ameliorated histopathological changes affected by croton oil application. In the lipopolysaccharide (LPS)-induced inflammation in air pouch in rats, JLW reduced nitric oxide (NO) level and tumor necrosis factor-alpha (TNF-α) release. In conclusion, this study demonstrates the effectiveness of JLW in combating inflammation in several experimental models. Further investigations are needed to identify the active constituents responsible for the anti-inflammatory property of JLW.

Keywords: Jojoba liquid wax; Anti-inflammatory; Prostaglandin E2; Myeloperoxidase

1. Introduction

Jojoba [Simmondsia chinensis (Link 1822) Schneider 1907] is an arid perennial shrub indigenous to Arizona, California and Northwestern Mexico [1]. It is also grown in Australia [2], Brazil, Argentina and some Middle East countries [3]. Jojoba seed is rich in liquid wax, commonly mistaken for “jojoba oil” [4]. As much as 97% of jojoba liquid wax (JLW) consists of a mixture of esters of long chain fatty alcohols and long chain fatty acids. More than 60% of this mixture of esters contains cis-11-eicosenoic (jojobenoic) acid (C20) [5]. JLW contains a natural antioxidant postulated to be an allylic derivative of hydroxytoluene [6]. JLW was used in folk remedies for renal colic, sunburn, chaffed skin, hair loss, headache, wounds and sore throat [7]. Animal studies demonstrate that JLW can be classified as a non-toxic substance [8]. JLW has shown moderate digestibility that increases when the wax is mixed with other oils [9]. In addition, JLW possesses moderate absorption when applied topically [10]. In human studies, sulfurized JLW was effective in the treatment of acne while the unmodified wax was used for treatment of psoriasis [11]. Furthermore, dermatological research suggests that JLW may help to reduce inflammation [12].

Inflammation is an essential protective process preserving the integrity of organisms against physical, chemical and infective insults [13]. However, it is frequent that the inflammatory response to several insults erroneously leads to the damaging of normal tissues [14]. Prostaglandin E2 (PGE2) is generated from arachidonic acid by the enzyme cyclo-oxygenase (COX) at sites of inflammation in substantial amounts and can mediate many of the pathologic features of inflammation [15]. One of the early cellular events in inflammation is the margination of leukocytes, primarily neutrophils. This response can be measured by using the neutrophil specific enzyme myeloperoxidase (MPO), an indicator of neutrophil accumulation [16]. In addition, nitric oxide (NO) plays an important role in inflammation and NO synthase (NOS) inhibitors can reverse several classic inflammatory symptoms [17]. Tumor necrosis factor-alpha (TNF-α), a cytokine, plays an important role in inflammation. TNF-α stimulates neutrophils to transcribe and release cytokines,
and chemokines biosynthesis [18]. The conventionally used therapies for inflammation; non-steroidal anti-inflammatory drugs (NSAID’s), have very important role in managing pain and inflammatory conditions [19], though with rather discouraging profile of side effects [20]. Even the newer anti-inflammatory drugs, cyclo-oxygenase 2 (COX-2) inhibitors, are not devoid of adverse effects [21]. Meanwhile research has revealed that oxidative mechanisms are at the origin of inflammation, and has suggested the use of antioxidant substances [22]. This demonstrates the need for new and safe anti-inflammatory drugs. In this regard, natural products have long gained wide acceptance among the public and scientific community [23]. Therefore, the present study was designed to explore the potential anti-inflammatory effects of JLW.

2. Materials and methods

2.1. Chemicals

Croton oil, dimethylbenzidine, hexadecyltrimethylammonium bromide, o-dianisidine, indomethacin, lipopolysaccharide from E. coli serotype 0111:B4 (LPS), N-1-naphthylethyamine dihydrochloride (Gries reagent II), sodium nitrite and sulfanilamide (Griess reagent I) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Acetone, ethanol, ether and pyridine were purchased from SDS (Peypin, France). All other chemicals were of the highest available commercial grade.

2.2. Jojoba liquid wax

Before any processing, the plant (S. chinensis) was authenticated by Department of Pharmacognosy, Ain Shams University, Cairo, Egypt. Jojoba seeds were harvested in August by El-Salam Collaborative Agricultural Society, Mansyph, Ismailiya, Egypt. Fresh JLW was then prepared by the Extraction and Authentication Unit (EAPRU), Faculty of Pharmacy, Ain Shams University, Cairo, Egypt. Physical properties of the oil were as follows (Table 1).

Table 1

<table>
<thead>
<tr>
<th>Physical properties of JLW</th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Freezing point</td>
<td>9 °C</td>
</tr>
<tr>
<td>Boiling point</td>
<td>396 °C</td>
</tr>
<tr>
<td>Smoke point</td>
<td>195 °C</td>
</tr>
<tr>
<td>Flash point</td>
<td>295 °C</td>
</tr>
<tr>
<td>Refractive index at 25 °C</td>
<td>1.46</td>
</tr>
<tr>
<td>Specific gravity at 25 °C</td>
<td>0.87</td>
</tr>
<tr>
<td>Viscosity (25 °C)</td>
<td>50 cP</td>
</tr>
<tr>
<td>Iodine value</td>
<td>81</td>
</tr>
<tr>
<td>Saponification value</td>
<td>93</td>
</tr>
<tr>
<td>Acid value</td>
<td>2</td>
</tr>
<tr>
<td>Acetyl value</td>
<td>2</td>
</tr>
<tr>
<td>Unsaponifiable matter</td>
<td>51%</td>
</tr>
<tr>
<td>Total acids</td>
<td>52%</td>
</tr>
</tbody>
</table>

a) Determined according to official method, CC6a-48, of the American Oil Chemists, Society [24].

2.3. Animals

Throughout the experiments, adult male Sprague-Dawley rats weighing 150–175 g were used. Animals were housed at a temperature of 23 ± 2 °C with free access to water and standard food pellets (El-Nasr Co., Abo-Zaabal, Egypt). Rats were acclimatized in our animal facility for at least 1 week prior to any experiment. Protocol of the present work was approved by Experiments and Advanced Pharmaceutical Research Unit (EAPRU), Faculty of Pharmacy, Ain Shams University, Cairo, Egypt.

2.4. Measurement of paw volume and PGE2 in carrageenin-induced rat oedema model

Thirty rats were equally divided into five groups assigned Latin numbers I–V. Animals were fasted, with free access to water, 16 h before the experiment. Groups I and II were given saline by intragastric tube, while groups III and IV received JLW in two doses; 5 ml kg−1 (≈4.35 g) and 10 ml kg−1 (≈8.7 g), respectively. Animals in group V were orally treated with indomethacin as standard anti-inflammatory drug (10 mg kg−1). Dosing volume was kept constant (10 ml kg−1) and was completed with saline when required. The choice of the used doses and time of measurement and sampling was based on pilot studies in our laboratory. Thirty minutes after oral treatment, group I received 0.05 ml saline, while groups II–V received 0.05 ml carrageenin (1% solution in saline) sc on the plantar surface of the right hind paw. The right hind paw volume was measured immediately after carrageenin injection by water displacement using UGO-BASILE 7140 plethysmometer (Comerio, Italy) [25]. The volume was measured again 3 h after carrageenin injection and immediately before decapitation.

After decapitation, right hind paws were removed. A volume of 0.1 ml saline containing 10 µM indomethacin was injected to aid removal of the eicosanoid-containing fluid and to stop further production of PGE2. Paws were incised with a scalpel and suspended off the bottom of polypropylene tubes with Lipendorf pipette tips to facilitate drainage of the inflammatory exudates. For the purpose of the removal of the inflammatory exudates, paws were centrifuged at 18,000 g for 15 min [26]. PGE2 was quantified in the collected exudates using a quantitative binding PGE2 enzyme immunoassay kit. The kit uses a monoclonal antibody to bind, in a competitive manner, the PGE2 in the sample as well as alkaline phosphatase-labelled PGE2 provided in the kit. The enzyme bound, through PGE2 molecule, to the monoclonal antibodies processes the specific substrate to produce a colour that is measured spectrophotometrically [27].
2.5. Chick’s embryo chorioallantoic membrane (CAM) test

The chick’s embryo CAM test was carried out according to the procedures described by D’Arcy and Howard [28]. Fertile hen’s eggs of a local strain (Fayoumi chickens) were purchased from the chicken farm of The Faculty of Agriculture, Cairo University (Giza, Egypt) and Whatmann No. 1 filter paper was purchased from Fisher Scientific Co. (Pittsburgh, PA, USA). Eggs were collected 24 h after being laid. Twenty four fertile eggs, arranged in four groups, were used. Eggs were immediately incubated at a temperature of 37 °C at the time of receipt. Relative humidity was adjusted at 80%. Eggs were rotated four times daily. Eight days later, eggs were implanted with filter paper discs measuring 10 mm in diameter and weighing 7 ± 0.04 mg. The discs were prepared by carefully punching them out from No. 1 Whatmann filter paper using a pneumatic cork borer. After autoclaving, discs were impregnated with sterile drug solutions under aseptic conditions. Each disc was loaded with 14.5 μg drug solution. Indomethacin was dissolved in water, while JLW was dissolved in ether and thoroughly dried prior to application to the CAM. JLW was applied at the dose of 3.5 and 7 μg per disc (approximately 3.05 and 6.1 mg JLW per disc, respectively). Indomethacin was used as a positive control in concentration of 2.5 g per disc, respectively. Group IV received indomethacin-loaded discs. Only the unmedicated discs while groups II and III were implanted with discs loaded with JLW low and high doses, respectively. Group IV received indomethacin-loaded discs. Using aseptic techniques, eggshells were opened using a dental drill; loaded discs were placed on the surface of CAM, the removed piece of shell was placed again and the opening was sealed using molten paraffin wax. Eggs were re-incubated under the same conditions of temperature and humidity without rotation. Eggs were opened, 4 days later, by cutting the shells circumferentially along the longer perimeter. CAM membranes were eased out of the shell and the discs along with the adhering, and sometimes infiltrating, granulation tissue were cut with fine scissors. The discs carrying the granulation tissue were dried overnight at 55 °C and weighed individually.

2.6. Assessment of ear oedema, tissue myeloperoxidase activity, and histopathology in croton oil-induced ear oedema model in rats

The experiment was performed using a slight modification of the procedure described by Tonelli et al. [29]. An irritant solution was prepared by dissolving 4 parts croton oil (the irritant) in a solvent mixture of 10 parts ethanol, 20 parts pyridine, and 66 parts ethyl ether. JLW and indomethacin, serving as positive control, were dissolved in the same vehicle of the irritant. Two different concentrations of JLW were employed; namely 30 and 50% (v/v). Indomethacin was used in dose of 12.5% (w/v) [30]. Thirty rats were equally arranged in five groups numbered I–V. Irritant, JLW and indomethacin solutions were applied in volume of 20 μl topically on both sides of the right ears. The left ear was kept untreated to serve as a control. Group I served as negative control hence received only the irritant-free solvent mixture. Third and fourth group received JLW solutions and group V received indomethacin solution. One hour later, groups II–V received croton oil solution and group I was re-administered the croton oil-free solvent mixture again. After 4 h, animals were decapitated. An 8-mm cork borer was used to punch out discs from both the treated as well as the control ears. The two punches were weighed immediately after decapitation and the difference in weight was used to assess the inflammatory response.

The entire tissue of the right ear was homogenized for 10 min in ice bath (10%, w/v) in 50 mM phosphate buffer (pH 6.0) containing 0.5% hexadecyltrimethylammonium bromide with a Glas-Col® homogenizer. Tissue suspensions were centrifuged at 40 000 × g for 15 min. An aliquot of 0.1 ml of the supernatant was added to 2.9 ml of 50 mM phosphate buffer (pH 6.0) containing 0.167 mg ml⁻¹ o-dianisidine dihydrochloride, serving as MPO substrate, and 0.0005% hydrogen peroxide. The change in absorbance was measured at 460 nm with a SHIMADZU® UV-1601 spectrophotometer at 25 °C. MPO activity was quantified kinetically; change in absorbance was measured over a period of 2 min, sampled at intervals of 15 s. The maximal change in absorbency per minute was used to calculate the units of myeloperoxidase activity based on the molar absorbancy index of oxidized o-dianisidine dihydrochloride which equals 1.13 × 10⁷ M⁻¹ cm⁻¹. One unit of myeloperoxidase is defined as that degrading one micromole of peroxide per minute at 25 °C [32]. Results were expressed as units of activity per milligram protein. Protein content was determined according to Lowery et al. [33]. A representative ear tissue from each group was fixed in 10% formalin solution and kept until the time of preparation of sections. Ear tissue was embedded in paraffin; sections were cut using the microtome at a thickness of 5 μm. Sections were stained using hematoxylin and eosin. Then the specimens were observed under a light microscope and photographed. Specimens were examined regarding leukocytic infiltration, oedema as indicated by collagen fiber dispersion and extravasations.

2.7. Measurement of NO and TNF-α in rat air pouch model

Thirty rats uniformly divided in five groups were used in the study. Air pouches were formed by subcutaneous injection of 20 ml sterile air in the suprascapular area of the back of the animal. Three days later, the pouches were re-inflated with 10 ml sterile air. Three days later, 100 μg ml⁻¹ solution of LPS in physiological saline (1 ml kg⁻¹) was administered intrapouch to groups II–V. Group I received only saline. Thirty minutes later, drugs were injected also intrapouch. JLW dose was 5 and 10 mg kg⁻¹, animal body weight. Indomethacin concentration used was 10 mg kg⁻¹. The volume of administered solutions was kept constant at three
Groups I and II received only saline, groups III and IV received JLW low and high doses, respectively and group V received indomethacin. The doses of JLW in this model were based on a pilot trial in our laboratory. Eight hours later animals were sacrificed [34]. Each pouch was lavaged using 1 ml of sterile physiological saline. The lavage fluid was then centrifuged at 3000 × g for 5 min. Supernatant was used immediately for analysis of NO and TNF-α.

NO was assayed by measuring nitrite accumulation utilizing Griess reaction [35]. Briefly 100 μl of the air pouch exudates was combined with 100 μl of a 1:1 mixture of Griess reagent I (1% sulfanilamide in 5% phosphoric acid) and Griess reagent II (0.1% N-1-naphthylethylenediamine dihydrochloride in water) in a flat-bottomed microtiter plate. The reaction mixture was incubated for 40 min at 37°C in a shaking water bath. In the presence of nitrite, azo-dye reaction proceeds in an acid medium producing scarlet color, which can be measured spectrophotometrically at 540 nm in an ELISA reader. The concentrations of nitrite samples were calculated from sodium nitrite standard curve run in the same plate. Results were expressed as μM nitrite. TNF-α was assayed using rat TNF-α enzyme immunometric assay kit. Rat TNF-α was immobilized on polyclonal antibody bound to microtitre plate. Excess sample was washed. A monoclonal antibody specific to rat TNF-α, coupled to horseradish peroxidase, was added. The monoclonal antibody binds specifically to the immobilized rat TNF-α. Excess monoclonal antibody was washed and the substrate, tetramethyl benzidine, was added. After incubation period, the developed colour was measured spectrophotometrically at 450 nm [27].

3. Statistical analysis

Results are reported as mean ± S.E.M. Statistical analysis was performed using one-way analysis of variance (ANOVA). If the overall F-value was found statistically significant (P < 0.05), further comparisons among groups were made according to post hoc Tukey’s test. All statistical analyses were performed using SPSS statistical software package (SPSS® Inc., USA) version 8. Graphs were sketched using GraphPad Prism (ISI® software, USA) version 2 software.

4. Results

4.1. Effect of JLW on paw volume and PGE2 in carrageenin-induced rat oedema

Intraplantar injection of carrageenin to rats resulted in severe discernible inflammation and significant increase in the mean volume of the challenged paw compared to that of the untreated paws (162.3% of the untreated paws) (Table 2). Pretreatment of rats with JLW in doses of 5 and 10 ml kg⁻¹ significantly inhibited the carrageenin-induced increase in the oedema volume of the paws by 26.4 and 34%, respectively. Similarly, indomethacin-treated group showed significant anti-oedema effect (43.4% of the induced paws).

Carrageenin challenge resulted in more than five-fold increase in PGE2 concentration in inflammatory exudates in group II compared to unchallenged animals in group I (Fig. 1). Animals receiving JLW showed significant reduction of the PGE2 concentration in exudates that was dose-related in nature (reduction by 58.15 and 77.4% of the carrageenin-treated animals, respectively). The higher dose of JLW as well as indomethacin could lower PGE2 level in carrageenin-challenged animals approaching normal levels.

![Fig. 1. Effect of JLW on PGE2 production in exudates from carrageenin-treated rats.](image-url)}
Table 3

Effect of JLW on weight of granulation tissue in chick’s embryo CAM experiment

<table>
<thead>
<tr>
<th>Group</th>
<th>Weight of granulation tissue (mg)</th>
<th>Inhibition of the induced discs (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Induced</td>
<td>12.92 ± 0.06</td>
<td>0</td>
</tr>
<tr>
<td>JLW (3.05 µg per disc)</td>
<td>10.87 ± 1.08</td>
<td>15.86</td>
</tr>
<tr>
<td>JLW (6.1 µg per disc)</td>
<td>8 ± 0.23</td>
<td>38.08</td>
</tr>
<tr>
<td>Indomethacin (2.5 µg per disc)</td>
<td>4.3 ± 0.40</td>
<td>66.7</td>
</tr>
</tbody>
</table>

* Statistically significant difference from the induced discs at P < 0.05.

4.2. Effect of JLW on granulation tissue formation in the chick’s embryo CAM experiment

The data in Table 3 show that introduction of filter paper discs to CAM resulted in accumulation of granulation tissue on discs that mounted to 12.9 mg per disc. Preloading of the filter paper discs with JLW in concentration of 30% (v/v) and 50% (v/v) resulted in statistically significant reduction of granulation tissue weight by 15.8 and 38%, respectively compared to plain discs. Similarly, indomethacin produced significant lowering (66.7%) of the granulation tissue weight per disc compared to untreated discs.

Table 4

Effect of JLW on weight of croton oil-induced ear oedema in rats

<table>
<thead>
<tr>
<th>Group</th>
<th>Ear-punch weight (g)</th>
<th>Un-induced ear-punch weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.018 ± 0.0004</td>
<td>100</td>
</tr>
<tr>
<td>Croton oil-induced</td>
<td>0.039 ± 0.0014</td>
<td>216.7</td>
</tr>
<tr>
<td>JLW (30%, v/v)</td>
<td>0.028 ± 0.0006</td>
<td>155</td>
</tr>
<tr>
<td>JLW (50%, v/v)</td>
<td>0.023 ± 0.0005</td>
<td>122</td>
</tr>
<tr>
<td>Indomethacin (12.5%, w/v)</td>
<td>0.029 ± 0.0004</td>
<td>161</td>
</tr>
</tbody>
</table>

* Statistically significant from the control group at P < 0.05.

4.3. Effect on croton oil-induced ear oedema, MPO tissue activity and histopathological changes

Application of croton oil to rat ears caused massive increase in the weight of the ear punch (216%) compared to unchallenged ears. Pretreatment of rat ears with the two dose levels of JLW, 30% (v/v) and 50% (v/v) significantly reduced the increase of punch weight by 28 and 43.6%, respectively. In addition, indomethacin produced significant reduction of oedema by 25.6% (Table 4).

Histopathological examination of the ear tissue confirmed those results obtained by assessing MPO activity. Fig. 3, Plate A shows normal histological characteristics of the epidermal, dermal as well as subcutaneous layers with no obvious neutrophil infiltration. The croton oil-treated rats showed massive neutrophil infiltration with extravasations of RBC’s as well as oedema in the dermal layer (Fig. 3, Plate B). Fig. 3, Plate C, corresponding to JLW 30% (v/v), shows less neutrophil infiltration and less hyperemia. Fig. 3, Plates D and E, which represent the ear tissue treated with 50%v(v) JLW solution and indomethacin, respectively showed almost normal dermal tissue with mild degradation of discrete cells of sebaceous glands.

4.4. Effect of JLW on NO and TNF-α in air pouch model

Injection of LPS into the air pouch caused about 60-fold increase of NO production compared to that untreated animals (group I). JLW injection in doses of 5 and 10 ml kg⁻¹ showed significant reduction of NO level by 31.4 and 32.8%, respectively compared to group II (Fig. 4). However, the...
Fig. 3. Effect of JLW on histopathological changes in croton oil-induced ear oedema experiment. Plate A shows normal architecture of the covering dermal and epidermal layers as well as subcutaneous tissue of the skin. Plate B shows ear tissue from croton oil-alone treated rats exhibiting massive neutrophil infiltration with extravasations of red blood cells and oedema in the dermal layer. Plates C–E, corresponding to JLW 30% (v/v), 50% (v/v) and indomethacin 12.5% (w/v), respectively, show less neutrophil infiltration, collagen fiber dispersion and oedema (40×).
Fig. 3. (Continued)
effect was not dose related. Similarly, indomethacin treatment significantly lowered NO level by 36.6%.

In addition, LPS injection caused about 8-fold increase in TNF-α level in the pouch exudates compared to untreated animals. JLW treatment in doses of 5 g kg⁻¹ and 10 ml kg⁻¹ lowered TNF-α release by 62.2 and 75.8%, respectively (Fig. 5). Indomethacin, serving as positive control caused a reduction mounting to 43.4% of TNF-α level in rats of group II.

5. Discussion

Jojoba [Simmondsia chinensis (Link 1822) Schneider 1907] has been long used as a folk medicine for a variety of conditions, including inflammatory diseases, by the ancient natives of the American continents [7]. Inflammation is the most common aspect of tissue pathology and has always taken a central role in medical practice. The aim of the present work was to evaluate the potential anti-inflammatory effect of JLW as well as the underlying possible mechanism.
The results of carrageenin-induced paw oedema [25] showed that JLW reduces carrageenin-induced paw oedema. This finding could be interpreted as a potential anti-inflammatory activity of JLW. Such conclusion was further substantiated by assessing PGE2 level in the inflammatory exudates of rat paws in the same model. Prostaglandins have been long recognized as a major mediator of inflammation. They are arachidonic acid metabolites synthesized by COX-1 and COX-2 isozymes [36]. Our results indicate that JLW caused statistically significant reduction of PGE2 content. This finding clearly underlines the anti-inflammatory effect of JLW. Such activity may be explained on the basis of JLW content of long chain fatty acid esters. These esters include erucyl jojobenoate and jojobenyl jojobenoate, which constitute 61% of the total esters present. Jojobenoic acid; cis-11-eicosenoic acid, is produced by hydrolysis of these esters by lipases [37] and constitutes 71% of the total fatty acids [5]. The hydrolysis of these esters by lipases results in liberation of free fatty acids which are anticipated to compete with arachidonic acid for the binding site on COX isozymes [38]. It may be suggested that this competition results in decreased synthesis of PGE2.

Chick’s embryo CAM test is an accurate method for the screening of anti-inflammatory effect of an agent directly on the developing CAM [28]. JLW demonstrated an anti-inflammatory effect as indicated by significant reduction of the weight of the granulation tissue.

The results of croton oil-induced ear oedema support this finding [29]. A reduction in ear punch weight was shown after application of the two dose levels of JLW compared to the croton oil-alone condition. This activity could be justified by the remarkable cutaneous absorption of JLW. Sebum consists of several lipids, the major constituents of which are wax, free fatty acids and other lipid esters [39]. Owing to the similarity of JLW composition to that of sebum, JLW is efficiently absorbed which may account for its powerful anti-inflammatory activity.

The findings of the same experiment demonstrated that JLW reduced MPO activity in challenged ear tissue. This could be explained on the basis of inhibition of neutrophil infiltration which was confirmed visually by microscopic examination of the ear tissue. JLW reduced neutrophil infiltration, red cell extravasation, oedema and hence collagen fiber dispersion. Cutaneous inflammation is characterized by the infiltration of polymorphonuclear leukocytes such as neutrophils [40]. MPO, a heme protein, is a major component of azurophilic granules of neutrophils. Oxygen-dependent microbicidal activity depends on MPO as the critical enzyme for the generation of hypochlorous acid and other toxic oxygen products [41]. The primary function of neutrophils is the release of MPO into the phagosome containing the ingested microorganism leading to a rapid microbicidal effect. MPO can be released to the outside of the cell inducing damage to adjacent tissue and, thus, contribute to the pathogenesis of inflammation [42].

NO is a highly fat soluble, relatively stable free radical, having numerous promiscuous roles [43]. NO synthesis is greatly amplified by LPS [44]. Several studies have demonstrated that inflammation correlates with level of NO [45]. In this study JLW have shown remarkable, though not dose related, reduction of nitrite accumulation in the air pouch model. This finding may be attributed to the anti-oxidant property of JLW reported to protect the oil itself against rancidity [6]. Furthermore, lowering of NO level after JLW administration may account for the reduction in PGE2 level since many reports have shown that NO rapidly and strongly stimulates COX enzymes [46]. In addition, NO synthase inhibitors reduce PGE2 biosynthesis [47].

TNF-α is a pleiotropic cytokine which plays a critical role in both acute and chronic inflammation [48]. TNF-α promotes an acute phase reaction [49]. Several inflammagens have the ability of inducing the synthesis of TNF-α. The formation of a number of small molecular mediators of inflammation is linked with TNF-α and thus contributes to the range of mediators that critically control inflammation [50]. TNF-α facilitates inflammatory cell infiltration by promoting the adhesion of neutrophils and lymphocytes to endothelial cells [51]. When TNF-α effect is specifically blocked, the severity of inflammation is reduced [52]. More importantly, TNF-α induces the synthesis of PGE2 [49]. JLW, in particular the higher dose, strongly inhibited TNF-α release. The reduction of TNF-α release by JLW is in agreement with the PGE2 finding obtained in the current study. An alternative explanation of the resultant reduction of PGE2 level in inflammatory exudates may be through interference with TNF-α release.

In conclusion, JLW exerts anti-inflammatory activity in several animal models. This is evidenced by decreased paw oedema, PGE2 content in exudates, CAM granulation tissue formation, ear oedema, MPO activity, NO generation and TNF-α formation as well as amelioration of inflammatory histopathological changes. These results lend support to the effectiveness of JLW in combating inflammation via multilevel regulation of inflammatory mediators. Further investigations are needed to identify the active constituents responsible for the anti-inflammatory property of JLW.

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References

