Effect of Clausena excavata Burm. f. (Rutaceae) leaf extract on wound healing and antioxidant activity in rats

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Abstract: Clausena excavata is a well-known plant used in folkloric medicine for the treatment of different ailments. This study aimed to determine the in vitro cytotoxicity of its leaf solvent extracts as well as the in vivo wound healing and antioxidant activities of the methanolic extracts of C. excavata (MECE). HaCaT (keratocyte) and Vero cell lines were used for evaluation of the in vitro cytotoxic effects, while the in vivo wound healing and antioxidant activities were determined in skin wounds inflicted on rats. Twenty adult male Sprague-Dawley rats were divided into five groups of four animals each. Approximately 3.14 cm² excisional wound was inflicted on the nape of each rat following anesthesia. The treatment groups received topical application of MECE at 50 mg/mL (MECE-LD [low dose]), 100 mg/mL (MECE-MD [medium dose]), and 200 mg/mL (MECE-HD [high dose]), while the negative control group was treated with gum acacia in normal saline and the positive control group with intrasite gel. Wound contraction was evaluated on days 5, 10, and 15 after wound infliction, and tissue from wound area was collected at day 15 post-wound infliction for antioxidant enzyme evaluation and histopathological analyses. Generally, Vero cells were more resistant to the cytotoxic effects of the solvent extracts as compared with HaCaT cells. Chloroform (CH) and ethyl acetate (EA) extracts of C. excavata were toxic to HaCaT cells at 200 and 400 µg/mL, but the same concentrations showed higher (P<0.05) viability in Vero cells. There was significantly (P<0.01) greater wound contraction at days 10 and 15 post-wound infliction in all the treatment groups than in the control groups. Histopathologically, the MECE-HD-treated wound showed significantly (P<0.05) lesser inflammatory cell proliferation, degeneration, and distribution of granulation tissue than other groups. Similarly, the degree of collagen maturation, angiogenesis, and collagen distribution were significantly (P<0.05) lower in MECE-HD than in other groups. The MECE-HD, MECE-MD, and intrasite treatment groups showed a significantly (P<0.05) higher number of VEGF-positive and TGF-β1-positive cells in the skin wound than the control groups. The activities of superoxide dismutase and catalase were significantly (P<0.01) higher in the MECE-HD and intrasite treatment groups than in the other groups. Lipid peroxidase activity of the treated groups was significantly (P<0.01) lower than that in the control group. The study showed that MECE is a potent wound healing agent through anti-inflammatory and antioxidant effects that enhanced the rate of wound contraction, re-epithelialization, and collagen deposition. The effect of MECE is suggested to be due to its high polyphenolic compound content.

Keywords: Clausena excavata, cytotoxicity, wound healing, antioxidant, histopathology, VEGF, TGF-β1

Introduction

Clausena excavata Burm. f. is one of the potherb species with high antioxidant properties. Its leaves and stem are used in folk medicine for treatment disorders such
as colic, cough, headache, rhinitis, sores, wounds, fever, and
detoxification. This plant has been reported to possess various
biological properties including anti-inflammatory, antimicro-
bial, antinociceptive and immunomodulatory, antifungal,
antioxidant, and analgesic properties.1–5 The acetone extract of
its leaves has been shown to contain several furanocoumarins
with in vitro antiproliferative activities on cancer cells.6

Wound healing is a complex process involving both bio-
chemical and physiological changes. A wound provides an
environment for the growth of microorganisms, and thus an
infected wound shows delayed or prolonged healing time.6

During the inflammation phase, various inflammat-
ory cells like neutrophils, macrophages, fibroblasts, and
endothelial cells produce a sudden burst of reactive oxygen
species (ROS) in the site of the wound, providing signaling
and defense against invading microorganisms. ROS, which
include hydroxyl radical, superoxide anion, hydrogen pero-
oxide, and singlet oxygen, have destructive action on both
DNA and proteins. Overproduction of ROS may also hinder
the rate of wound healing by causing damage to surround-
ning cells. Thus, as expected, plants possessing antioxidant
properties can prevent oxidative damage to cells induced by
wound injury, thereby promoting healing.7,8

One of the many mechanisms of wound healing is the
promotion of angiogenesis in the injured tissue. Vascular
endothelial growth factor (VEGF) and transforming growth
factor β1 (TGF-β1) are among the effectors of angiogenesis,
granulation tissue formation, collagen synthesis, and extracel-
ular matrix deposition.9,10 Increased expressions of these fac-
tors are associated with acceleration of wound healing.11

C. excavata possesses strong antioxidant properties and
is a rich source of polyphenol compounds such as coumarins,
carbazole alkaloid, and flavonoid glycosides.5,12,13 However,
there is no clear evidence that C. excavata, with its potent
antioxidant activity, can enhance wound healing in the animal
model. This study was thus undertaken to determine the
in vitro effect of C. excavata extracts on HaCaT (keratino-
cyte) and Vero cells, and the in vivo effect on experimentally
induced skin wound in rats.

Materials and methods
Collection, identification, processing, and
extraction of plant material
C. excavata leaves were collected, identified by Dr Shamsul
Khamis (Resident Botanist) at the Biodiversity unit, Institute
of Bioscience, Universiti Putra Malaysia, processed, and
extracted as previously described by Albaayit et al.14

Briefly, extraction was done successively at room tem-
perature using petroleum ether (PT) followed by chloroform
(CH), ethyl acetate (EA), and methanol (MOH). The extrac-
tion at a 1:5 dried plant weight to volume ratio began with
PT for 3 days. The filtrate was collected, and the residues
were subjected to further extraction with CH, EA, and MOH,
respectively, and the filtrates were collected after each solvent
extraction. All filtrates were evaporated until dryness using a
rotary evaporator at 45°C–50°C to obtain crude extracts and
stored at 4°C until use. Among the extracts, methanolic extract Clausena excavata (MECE) exhibited the most potent
antioxidant activity, and hence it was selected for wound
healing potential in this study.

In vitro screening of different solvent
crude extracts
Cell maintenance
The HaCaT and Vero cell lines (American Type Culture
Collection [ATCC], Manassas, VA, USA) were cultured in
75 cm² T-flasks using RPMI and DMEM media, respectively,
fortified with 10% fetal bovine serum, and incubated at 37°C
in a 5% CO₂ humidified incubator. Cells reaching 80%
confluency were detached using Trypsin–EDTA (Thermo
Fisher Scientific, Waltham, MA, USA), centrifuged at
1,710×g, and counted using trypan blue exclusion assay.

Evaluation of crude extract fractions on cell viability
using (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenylte-
zolium bromide [MTT]) colorimetric assay
The MTT assay is one of the most commonly used colorimetric
indices to determine cell viability/cytotoxicity.15 The HaCaT
and Vero cells were seeded at a density of 6.0×10⁴ cells/well
in a 96-well plate. After incubation for 24 hours, the cells
were washed with phosphate buffered saline (PBS). The drug
was prepared using corresponding growth media for each cell
line. The vehicle for initial stock of drug was 0.1% dimethyl
sulfoxide (DMSO). Approximately 12.5, 25, 50, 100, 200, and
400 µg/mL of the methanolic MOH, EA, CH, and PT crude
extracts were added to the cells and incubated at 37°C and
5% CO₂ for 24 hours. Twenty microliters of MTT reagent was
then added to each well and incubated for 3 hours. The purple
formazan formed was solubilized with 150 µL of DMSO in
the dark at room temperature for approximately 30 minutes.
The absorbance was recorded at 570 nm, with reference at
630 nm, in a microplate reader (Tecan, Salzburg, Austria).
Each experiment was repeated three times with triplicate wells
for each concentration.

Experimental animal
Twenty adult male Sprague-Dawley rats (200–250 g) were
used in the study. The rats were acclimatized for 2 weeks
in a well-ventilated room, and standard pellet feed and tap water were provided ad libitum throughout the experiment period. The project proposal was approved by the Institutional Animal Care and Use Committee, University of Malaya (ISB/22/007/2013/1111/SFA).

**Wound infliction**

The rats were divided into five groups of four animals each, and all animals were anesthetized with an intramuscular injection of ketamine/xylazine at the ratio of 3:1 (100 mg/mL and 30 mg/mL, respectively), prior to infliction of skin wounds. Skin from the dorsal nape of the neck was shaved with an electrical clipper, disinfected with 70% alcohol, and injected with 0.1 mL lignocaine hydrochloride (2%, 100 mg/5 mL). A 2 cm diameter circular area (~3.14 cm²) of the skin was made using a circular stamp, and full thickness of the marked skin was then cut carefully by sterile scissors and forceps (Figure 1). Wound excision was done carefully to avoid muscle incision. Group 1 (negative control group) was treated with vehicle (gum acacia in normal saline 20 mg/mL); Group 2 was treated with 0.2 mL intrasite gel (positive control group); Groups 3, 4, and 5 were treated with 0.2 mL *C. excavata* MOH extract (MECE) at concentrations of 50 (MECE-LD [low dose]), 100 (MECE-MD [medium dose]), and 200 (MECE-HD [high dose]) mg/mL. All treatments were by topical application twice daily for 14 days starting from day 0 of wound infliction.

**Wound contraction**

After the surgical process, the wound area was measured every 5 days for 15 days by placing transparent paper and defining the borders of the wound. The transparent paper was immediately traced out and placed on a 1 mm² graph paper. The rate of change in wound contraction was expressed as a percentage of the initial wound size as given in the formula below. Differences in wound healing of the treated groups are derived by comparing with healed wound area of the control group on the respective days.¹⁶

\[
\text{Wound contraction} = \frac{\text{Original wound area} - \text{present wound area}}{\text{Total wound area}} \times 100
\]

**Histopathological examinations**

**H&E- and Masson trichome-stained sections**

On day 15 of the experiment, skin specimens from all the rats were collected in 10% buffered formalin, processed, sectioned, and stained with hematoxylin and eosin (H&E) and Masson’s trichrome (MT). Six microscopic fields were examined from three slides per group at 400× magnification and the lesions scored based on a five-point scale: 0 – normal, 0%; 1 – occasional evidence, 25%; 2 – light scattering, 30%–50%; 3 – abundant evidence, 55%–75%; and 4 – confluent cell, 80%–100%. The H&E-stained sections were examined for epithelialization (based on complete or incomplete epithelial proliferation in the epidermal layer), degeneration (based on the distribution of epidermal cells showing vacuolation or vacuolar degeneration), granulation tissue (based on the distribution of necrotic and inflammatory cells in the microscopic fields), and leucocytic infiltration (based on the distribution of polymorphonuclear and mononuclear cells in the microscopic fields) under light microscopy (Olympus BX60; Olympus, Tokyo, Japan) and the images captured (Olympus XC10 camera, Olympus).

The MT sections were similarly evaluated for color intensity (based on a subjective assessment of blue color intensities: no coloration [0], low [1], dull [2], moderate [3], and bright colorations [4]); fibroblast distribution (based on the distribution of fibroblast cells in the microscopic fields) under light microscopy (Olympus BX60; Olympus, Tokyo, Japan) and the images captured (Olympus XC10 camera, Olympus).

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field); collagen maturity (based on the level of maturation of collagen fibers: no collagen [0], thin fibers [2], semithick fibers [3], thick fibers [4]); angiogenesis rate (based on the distribution of new blood capillaries in the microscopic field); and collagen distribution (based on the distribution of collagen fibers in the microscopic fields) using the five-point scale modified as previously described. 9

Staining for VEGF and TGF-β1
Tissue sections were routinely prepared as described earlier for histopathology. The tissue sections were deparaffinized and immersed in graded concentrations of alcohol (100%, 80%, 70%, and 50%) for 5 minutes before being hydrated in PBS twice for 5 minutes. The sections were boiled in 10 mM Tris buffer (pH 9.0) for 15 minutes to retrieve the antigen, and the slides were cooled, blocked by endogenous peroxidase, and incubated for 10 minutes with 3% H2O2. The slides were then washed twice in PBS and incubated with polyclonal primary antibodies (anti-VEGF or anti-TGF-β1), Tween-20 rabbit polyclonal anti-VEGF or anti-TGF-β1 antibody (Abcam, Cambridge, UK) at a dilution of 1:500 and 1:100, respectively, and the tissue was incubated in a humidified chamber at 4°C for 1 hour (VEGF) or 18 hours (TGF-β1) overnight. The tissues were then stained with 3,3′-diaminobenzidine tetrahydrochloride (DAB) and substrate (Dako, Carpintaria, CA, USA) for 5 minutes. All washings were done with 0.1% Tween-20 in PBS, except for washings after rehydration and after chromogen incubation (DAB), in which PBS without Tween-20 was used. The slides were counterstained with hematoxylin, and subsequently dehydrated in increasing concentrations of alcohol (50%, 70%, 80%, and 100%) before being cleared in xylene. Tissue sections were air-dried and mounted onto slides for examination under light microscopy. Stained sections were evaluated for VEGF-positive cells in six fields each from three cut sections per group at 400× magnification. Positive cells stained light to dark brown were counted in each microscopic field. The extracellular matrix (ECM) distribution was evaluated from TGF-β1-stained sections by using a four-point scale: 0 – no ECM, 0%; 1 – little ECM, 25%–40%; 2 – moderate ECM, 50%–65%; and 3 – confluent ECM, 70%–100%, modified as previously described. 9

Antioxidant activity
Tissues samples were collected from all the animals at day 15 post-wound infliction for the determination of tissue superoxide dismutase (SOD), catalase (CAT), glutathione peroxidase (GPx), and lipid peroxidase activities.

Superoxide dismutase activity
The tissues were homogenized (Omni TH; Omni, Kennesaw, GA, USA) in 20 mM HEPES buffer (1 mM EGTA, 210 mM mannitol, and 70 mM sucrose/g tissue), pH 7.2. The homogenate was then centrifuged at 1,500×g for 5 minutes at 4°C, and the supernatant assayed for SOD activity using the Cayman assay kit (Cayman Chemical, Ann Arbor, MI, USA; www.caymanchem.com/pdfs/706002.pdf).

Catalase activity
Tissue samples were homogenized (Omni TH, Omni) in cold potassium phosphate buffer, pH 7.0, containing 1 mM EDTA. The mixture was then centrifuged at 10,000×g for 15 minutes at 4°C, and the supernatant assayed for CAT activity using the Cayman assay kit (Cayman Chemical; www.caymanchem.com/pdfs/707002.pdf).

Glutathione peroxidase activity
Tissue samples were homogenized (Omni TH, Omni) in 50 mM Tris-HCl, pH 7.5, 5 mM EDTA, and 1 mM dithiothreitol. The homogenate was then centrifuged at 10,000×g for 15 minutes at 4°C. The supernatant was assayed for GPx activity using the Cayman assay kit (Cayman Chemical; www.caymanchem.com/pdfs/703102.pdf).

Lipid peroxidation activity
The lipid peroxidation (LPO) assays (Cayman Chemical) were carried out according to manufacturer’s instructions. Tissue samples were homogenized (Omni TH, Omni) in PBS, CH, and extraction buffer provided with the kit to extract lipid hydroperoxides. Real-time chromogenic reaction was allowed to occur for 5 minutes. The absorbance was read at 500 nm (www.caymanchem.com/pdfs/705003.pdf).

Statistical analysis
Data obtained from the experiments were expressed as mean ± SE (standard error of mean), and subjected to one-way analysis of variance using SPSS (Version 19.0; IBM Corporation, Armonk, NY, USA) with significance at P<0.001 and P<0.05. The analyzed data were then presented in tables and bar graphs as shown.

Results
Cytotoxicity assay
The MTT cytotoxicity assay showed that MOH and PT extracts were less toxic to HaCaT cells at 400 µg/mL, where cell viability of more than 50% was observed. EA and CH extracts of C. excavata, on the other hand, were more toxic to
HaCaT cells at 200 and 400 µg/mL (Figure 1A). The MOH and EA extracts of *C. excavata* were slightly toxic to Vero cells at 400 µg/mL only, while CH and PT *C. excavata* extracts were innocuous to these cells at the same dose (Figure 1B). However, proliferation of the HaCaT and Vero cells increased following exposure to the extracts at low doses.

**Gross wound closure**

The gross appearances of wounds at days 5, 10, and 15 post-wound infliction are presented in Figure 2. The groups treated with various concentrations of MECE and intrasite gel showed minimal scar formation when compared with the gum acacia-treated control group, which showed incomplete wound closure. The skin wound areas in MECE-treated rats are presented in Table 1. The gross appearance of wound sites was similar among groups at day 5 post-wound infliction. The difference between the negative control group and the MECE-treated group on day 5 post-wounding was not significant.

However, at days 10 and 15, the rate of wound contraction was significantly different (*P*=0.0005; *P*<0.0001) in the MECE-treated group than in the negative control group. Although the rate of wound contraction appears to be highest on day 15 (Figure 2C–E), there was no significant (*P*>0.05) difference in the rate of healing among the treatment groups.

**Histopathological assessment of wound healing**

**H&E-stained sections**

The histopathological changes observed in the tissue sections are summarized in Table 2. There was significantly (*P*<0.0001) greater epithelialization in the epidermis of the treatment groups than in that of the control group. In general, more inflammatory cells (*P*=0.0039), degeneration (*P*=0.0011), and granulation tissue (*P*=0.0320) were observed in the skin of the control group than in that of the treatment groups, suggesting enhanced healing response in the MECE-treated groups after wound infliction (Figure 3A).

**MT-stained sections**

The changes in Masson trichome-stained skin tissues are presented in Table 3. The color intensity of the stained tissues was significantly (*P*<0.05) greater in the MECE-HD group than in the other groups. There was significantly (*P*=0.0003) lesser distribution of fibroblasts in the control group than in the other treatment groups. Collagen maturity was higher (*P*=0.0009) in the MECE-HD and intrasite groups, while the rates of angiogenesis (*P*=0.0121) and collagen distribution (*P*=0.0041) were significantly (*P*<0.05) higher in the MECE-MD and MECE-HD compared with the other groups (Figure 3B).

**VEGF and TGF-β1 in skin section**

The number of VEGF-positive cells was significantly (*P*=0.0002) lower in the control and MECE-LD groups than in the MECE-HD, MECE-MD, or intrasite groups (Figure 4). Extracellular matrix deposition as determined by the number of TGF-β1-positive cells was significantly (*P*<0.0001) higher in tissues treated with MECE and intrasite gel than in the control (Figure 5).

**Antioxidant enzyme assay**

The SOD activity was significantly higher in the MECE-HD (4.57±0.23; *P*=0.007), MECE-MD (5.39±1.01; *P*=0.001),
and intrasite gel groups (4.91±0.89; P=0.003) than that in the control group (1.83±0.27). There was no statistical difference between the treated groups (Figure 6A). The CAT activity was significantly higher in the MECE-HD (28.55±4.1; P=0.002) and intrasite gel groups (26.90±1.79; P=0.008) than that in the control group (17.82±1.07) (Figure 6B). In addition, there was an increase in the MECE-MD (23.77±1.40) and MECE-LD (20.91±0.90) groups but it was not significant. The GPx activity was significantly (P<0.001) higher in the intrasite and MECE-MD groups (32.9±1.87, 29.33±1.52, respectively) than in the control (16.4±0.35), MECE-HD, (23.0±2.5) or MECE-LD (21.8±2.02) groups (Figure 6C). The LPO activity was significantly higher in the control group (9.12±1.76) than in the intrasite (2.81±0.94; P=0.001), MECE-HD (4.17±1.10; P=0.006), MECE-MD (4.85±0.75; P=0.013), or MECE-LD (4.09±0.76; P=0.001) groups (Figure 6D). These results collectively show that wounds dressed with MECE may protect it from LPO during the healing process and thus contribute remarkably to acceleration of wound healing.

**Discussion**

In the past couple of decades, ethnopharmacology has become a focus area for researchers in the search for natural materials with potent antioxidant and healing properties. Among the properties of pure natural compounds and extracts investigated was wound healing, and the results were remarkable. In this study, the *C. excavata* MECE extract was examined for its wound healing properties. *C. excavata* contains coumarins, flavonoids, and glycosides with a wide array of biological properties.

Various solvents with increasing polarity have been frequently used for the extraction of polyphenols and flavonoids from plants, and since the biological activity of extracts has a strong relationship with the solvents employed on the basis of their polarities, we used different extracts from *C. excavata* leaves to determine their effect on in vitro cell lines. We observed that the MOH extract had cell viability of 71% and 47% at 400 µg/mL to the HaCaT and Vero cells, respectively. It was also observed that at low doses, the MECE and PT extracts of *C. excavata* enhanced the proliferation of these cells. The MECE contains queretin and myricetin that has been shown to enhance the growth of the normal human skin fibroblast (FS5) cell line by conferring protection on these cells against hydrogen peroxide-induced damage.

However, the nonpolar solvent (EA and CH) extracts showed a significant reduction in the percentage of cell viability in the HaCaT cells with 19% and 27% at 400 µg/mL, respectively. This may indicate that nonpolar active principles are responsible for the antiproliferative activity of this plant as previously reported in other plants. High

**Table 1** Effect of *Clausena excavata* on percentage of wound healing in experimental rats

<table>
<thead>
<tr>
<th>Days post-wound infliction</th>
<th>Wound healing postsurgery (%) in different groups</th>
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<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Day 5</td>
<td>21.7±5.1</td>
</tr>
<tr>
<td>Day 10</td>
<td>67.2±4.3</td>
</tr>
<tr>
<td>Day 15</td>
<td>78.9±2.5</td>
</tr>
</tbody>
</table>

**Notes:** All values were expressed as mean ± SE (standard error of mean). Mean with different superscripts within the same row are significantly different from each other (P<0.05), while those with the same superscripts within a row are not significantly different (P>0.05).

**Abbreviations:** MECE, methanol extract *Clausena excavata*; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).

**Table 2** Histopathological evaluation of the epidermis and dermis of H&E-stained skin sections of *Clausena excavata* methanol extract-treated rats

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Lesion score (unit)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Epidermis</td>
<td></td>
</tr>
<tr>
<td>Epithelialization</td>
<td>0.5±0.2</td>
</tr>
<tr>
<td>Inflammatory cells</td>
<td>2.5±0.6</td>
</tr>
<tr>
<td>Degeneration</td>
<td>2.3±0.5</td>
</tr>
<tr>
<td>Dermis</td>
<td></td>
</tr>
<tr>
<td>Granulation tissue</td>
<td>2.7±0.3</td>
</tr>
<tr>
<td>Inflammatory cells</td>
<td>2.5±0.2</td>
</tr>
</tbody>
</table>

**Notes:** All values were expressed as mean ± standard error mean; Mean with different superscripts within the same row are significantly different from each other (P<0.05), while those with the same superscripts within a row are not significantly different (P>0.05); lesions were scored based on a five-point scale: 0 – normal, 0%; 1 – occasional evidence, 25%; 2 – light scattering, 30%–50%; 3 – abundant evidence, 55%–75%; and 4 – confluent cell, 80%–100%.

**Abbreviations:** MECE, methanol extract *Clausena excavata*; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).
Figure 3 Histopathological evaluation of skin sections.

Notes: (A) Skin sections showing (Control) incomplete epidermal epithelialization (EE) with necrosis (N) and abundant inflammatory cell response (IC) in the dermis; (Intrasite) complete EE with cellular vacuolation (CV) and light scattering of inflammatory cells (IC) in the dermis; (MECE-LD) incomplete EE and a focus of hemorrhage (H) in the dermis; (MECE-MD) complete EE with occasional evidence of IC in the dermis; and (MECE-HD) complete EE with occasional evidence of IC in the dermis (H&E, 200×). (B) Skin sections showing (Control) occasional evidence to light scattering of collagen (C) with occasional evidence of blood vessels (BV) in the dermis; (Intrasite) light scattering to abundant evidence of angiogenesis (AG) and collagen distribution (CD), but moderate collagen color intensity; (MECE-LD) occasional evidence to light scattering of collagen (C) with abundant evidence of blood vessels (BV) in the dermis; (MECE-MD) abundant evidence of angiogenesis (AG) and collagen distribution (CD) with moderate collagen color intensity; (MECE-HD) abundant evidence of angiogenesis (AG), confluent distribution of collagen (CD) with moderate collagen color intensity.

Abbreviations: MECE, methanolic extract Clausena excavata; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).
In our previous study, we showed that MECE possesses high radical scavenging activities. However, it was uncertain whether this activity was similar in vivo and whether it would enhance wound healing. Antioxidants enhance the healing process by reducing the damage to tissue and cell structures including membrane lipids, proteins, enzymes, and nucleic acids caused by the production of ROS during the process of tissue injury. The antioxidant enzymes (SOD, CAT, and GPx) are known to overcome radicals and thus prevent the damage to cells caused by them. This was shown in a previous study using Plagiochasma appendiculatum, where enhanced wound healing rate was attributed to decreased activity of LPO and increased activities of SOD and CAT. Similarly, we observed increased SOD and CAT, flavonoids content has been reported to be responsible for antiproliferative activity in cranberry. Our previous study on the phytochemical screening of this plant showed the presence of high flavonoids in the CH extracts, which is responsible for the low proliferation observed here. The percentage cell viability observed in Vero cells was similar to that of HaCaT cells, but Vero cells were more resistant (51% and 78% at 400 µg/mL) than HaCaT cells following exposure to EA and CH. This selectivity could be due to the sensitivity of the cell line to the active compounds in the extract.

The wound healing properties of MECE were determined in a rat model by application of the extract to skin-infl ncted wounds. The extract increased wound contraction in all the treated groups by days 10 and 15. Histopathologically, the treated skin showed decreased inflammatory response, degeneration, granulation tissue, angiogenesis rate, collagen intensity, maturation, and distribution in the MECE-HD group. The findings are similar to that of previous studies that used different compounds and/or plants such as 3-(2-chlorophenyl)-1-phenyl-propenomein, ethanol extract of Bacopa monniera, Elaeag us angustofolia, Centaurea sadleriana janka, Blechnum orientale Linn, and Camellia sinensis. Compounds containing polyphenols have immune health benefits in view of their antioxidant, anti-inflammatory, and antimicrobial effects. These effects are responsible for the acceleration of wound healing that is facilitated by rapid wound closure and increased rate of epidermal regeneration, and these effects are attributed to the high phenolic content of MECE.

VEGF mediates angiogenesis and granulation tissue formation during wound healing. This cytokine, expressed in leukocytes stimulates reactive oxygen species (ROS) production. In fact, increased VEGF expression has been reported to be associated with accelerated wound healing in model animals, while decreased expression occurred in poor wound healing. We observed that MECE-MD and MECE-HD treatments increased the number of VEGF-positive cells in the wound tissue sections, which is suggested to be associated with increase in the growth factor expression. However, the pathway by which C. excavata extract induced this effect is unclear. It is possible that the anti-inflammatory properties of C. excavata extract are attributable to fucanocoumarins such as angelicin, which has been used as a medicinal agent in the treatment of psoriasis. This compound regulates inflammation via inhibition of the MAPK/NF-κB pathways. Our findings also showed that MECE-MD and MECE-HD treatments increased TGF-β1 expression; a cytokine enhances wound contraction, deposition of extracellular matrix, and collagen formation in the healing wounds. This suggests that one of the effects of MECE is to increase deposition of ECM in the skin wounds, facilitating healing.

Table 3 Evaluation of connective tissue and vascular proliferation in Masson’s trichrome-stained skin sections of Clausena excavata methanol extract-treated rats

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Lesion score (unit)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Intensity of coloration</td>
<td>1.6±0.2ab</td>
</tr>
<tr>
<td>Fibroblast distribution</td>
<td>1.5±0.2a</td>
</tr>
<tr>
<td>Collagen maturity</td>
<td>1.3±0.2a</td>
</tr>
<tr>
<td>Angiogenesis rate</td>
<td>2.0±0.4a</td>
</tr>
<tr>
<td>Collagen distribution</td>
<td>2.1±0.6a</td>
</tr>
</tbody>
</table>

Notes: All values were expressed as mean ± standard error of mean. *Mean with different superscripts within the same row are significantly different from each other (P<0.05), while those with the same superscripts within a row are not significantly different (P>0.05); fibroblast distribution, collagen distribution, and angiogenesis rate were scored on a five-point scale: 0 – normal, 0%; 1 – occasional evidence, 25%; 2 – light scattering, 30%–50%; 3 – abundant evidence, 55%–75%; 4 – confluent cell, 80%–100%. Collagen intensity was assessed based on subjective evaluation of blue color intensities as follows: no coloration (0), low (1), dull (2), moderate (3), and bright colorations (4). Collagen maturity was assessed based on level of maturation of collagen fibers: no collagen (0), semithin fibers (1), thin fibers (2), semithick fibers (3), thick fibers (4).

Abbreviations: MECE, methanol extract Clausena excavata; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).
**Figure 4** Quantitative estimation of VEGF reaction.

**Notes:** (A) Skin sections showing epidermis (E) and distribution of light to dark brown VEGF-positive cells (VP) in the dermis (D) in the various experimental groups: (Control), (MECE-LD), (MECE-MD), (MECE-HD), and (Intrasite). (Immunoperoxidase, 400x). (B) Bar graph showing distribution of VEGF-positive cells in the Control, Intrasite, and MECE-treated rat skin wound. *Indicates significance differences at $P<0.05$.

**Abbreviations:** MECE, methanolic extract Clausena excavata; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).
Figure 5 Evaluation of TGFβ1 distribution.

Notes: (A) Skin sections stained for TGF-β1 showing (Control) little deposition of extracellular matrix (ECM), (Intrasite) moderate ECM deposition, (MECE-LD) little to moderate ECM deposition, (MECE-MD) moderate ECM deposition, (MECE-HD) confluent ECM deposition. (Immunoperoxidase, 200×). (B) Bar graph showing distribution of extracellular matrix (ECM) in TGF-β1-stained skin sections of control, intrasite gel, and MECE-treated rat skin wound. *Indicates significance differences at P<0.05. The extracellular matrix distribution (ECM) was evaluated by using a four-point scale; 0 (no ECM); 0%, 1 (little ECM); 30%, 2 (moderate ECM); 65% and 3 (confluent ECM); 70%–100%, modified as previously described by Asba et al.7

Abbreviations: MECE, Methanolic extract Clausena excavata; LD, low dose (50 mg/mL); MD, medium dose (100 mg/mL); HD, high dose (200 mg/mL).
and decreased LPO activities in the MECE-HD group. These enzymes are very important mediators of oxidative stress as they catalyze the breakdown of ROS such as superoxide and hydrogen ions. The inhibition of ROS production and LPO promotes fibroblast proliferation, neovascularization, and the healing process of wounds. Earlier reports have shown that several compounds isolated from the roots of *C. excavata* exhibited profound antimicrobial properties. Thus, the healing properties exhibited by MECE is attributable to its antioxidant and antimicrobial activities, which in turn result from high phenolic content.

The results of this study showed that topical applications of MECE enhanced wound healing by improving different components of wound repair, including reduced inflammation, angiogenesis, increased collagen synthesis, and enhanced wound contraction, as well as decreased activities of lipid peroxidase and increased SOD, CAT, and GPx activities. VEGF and TGF-β expression in the MECE-treated group was dramatically increased and was higher than in the control group. However, an investigation of the major active compounds in MECE is still required to fully understand its mechanism of wound healing.

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**Author contributions**

All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflict of interest in this work.

**References**


