Composition of The Essential Oil From Danggui-zhiqiao Herb-Pair and Its Analgesic Activity and Effect on Hemorheology in Rats With Blood Stasis Syndrome

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ABSTRACT
Background: Angelica sinensis and Aurantii fructu used in a pair, named Danggui-Zhiqiao herb-pair (DZHP), which was rich in essential oil and has been adopted to promote blood circulation, dispel blood stasis, and relieve pain in traditional Chinese medicine (TCM). Objective: To analyze the composition and pharmacological effects of essential oil from DZHP. Materials and Methods: The composition of the essential oil from DZHP was analyzed by gas chromatography/mass spectrometry (GC/MS). Its analgesic activity was evaluated by acetic acid-induced writhing test and hot plate test. The hemorheology test was carried out to evaluate the effect on hemorheology in rats with blood stasis syndrome. Results: Twenty-eight components were identified and the main components were α-pinene (3.07%), β-pinene (2.0%), β-myrcene (3.71%), D-limonene (49.28%), γ-terpinene (9.53%), α-terpinolene (1.80%), α-terpinene (2.02%), β-bisabolene (1.13%), butyldenealthalide (1.43%), and Z-ligustilide (16.08%). The pharmacology test showed that the essential oil significantly inhibited the number of writhes induced by acetic acid with inhibition rate of 44.64% and significantly increased hot-plate latency compared with control group from 30 min to 90 min after oral administration of drugs in mice. It could significantly decrease plasma viscosity, whole blood relative index at high and low shear rate, whole blood reduced viscosity at high and low shear rate, and erythrocyte rigidity index in hemorheology test. Conclusion: The composition of the essential oil of DZHP was determined successfully and it had analgesic and promoting blood circulation activities. Key words: Danggui-Zhiqiao herb-pair, essential oil, composition, analgesic activity, hemorheology

SUMMARY
• Angelica sinensis and Aurantii fructu used in a pair, named Danggui-Zhiqiao herb-pair (DZHP), which was rich in essential oil and has been adopted to promote blood circulation, dispel blood stasis and relieve pain in traditional Chinese medicine (TCM).
• Twenty-eight components were identified and the main components were α-pinene (3.07%), β-pinene (2.0%), β-myrcene (3.71%), D-limonene (49.28%), γ-terpinene (9.53%), α-terpinolene (1.80%), α-terpinene (2.02%), β-bisabolene (1.13%), butyldenealthalide (1.43%), and Z-ligustilide (16.08%). The pharmacology test showed that the essential oil significantly inhibited the number of writhes induced by acetic acid with inhibition rate of 44.64% and significantly increased hot-plate latency compared with control group from 30 min to 90 min after oral administration of drugs in mice. It could significantly decrease plasma viscosity, whole blood relative index at high and low shear rate, whole blood reduced viscosity at high and low shear rate, and erythrocyte rigidity index in hemorheology test.
• The essential oil significantly inhibited the number of writhes induced by acetic acid and increased hot-plate latency compared with control group from 30 min to 90 min after oral administration of drugs in mice.
• The essential oil could significantly decrease plasma viscosity (PV), whole blood relative index (WBR), and erythrocyte rigidity index (ERI) in hemorheology test.

INTRODUCTION
Chinese herbal formulas, referring to the combination of two or more herbs, are the main form of clinical application in traditional Chinese medicine (TCM) and have been adopted to treat and prevent disease for thousands of years in China. More and more people from all over the world have recognized and accepted TCM formulas due to their remarkable efficacies and less adverse reactions. Herb pairs, mixture of two herbs, as the basic and the simplest composition units of Chinese herbal formulas, have special clinical significance without altering their basic therapeutic features in TCM.¹ ² So herb pairs were interested by many research workers and the composition, pharmacological effects, pharmacokinetics of herb pairs were widely studied in recent years.¹ ³ ⁴ Danggui-Zhiqiao herb-pair (DZHP) is comprised of Radix Angelica sinensis and Fructus Auranti (named as Danggui and Zhiqiao in Chinese, respectively). Angelica sinensis has been widely used to treat female

Abbreviations used: DZHP: Danggui-Zhiqiao herb-pair; TCM: traditional Chinese medicine; GC/MS: gas chromatography/mass spectrometry; PV: plasma viscosity; WBR: whole blood relative index; ERI: erythrocyte rigidity index

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menstrual disorders via nourishing the blood and activating blood circulation in TCM.[5] *Fructus Aurantii* is one of well known traditional herbal medicines to treat cardiovascular symptoms in China.[6] DZHP exists in many famous Chinese herbal formulas, such as Wu Ji san, Xuefu-Zhuyu decoction, Shujin Huxue decoction, Gexia Zhuyu decoction, Danggui-Zhiqiao decoction, which are all used to promote blood circulation, remove blood stasis, and relieve pain for long history.[7] Clinically, pain is one of the characteristics of blood stasis.[8] Essential oil is the DZHP’s active ingredient. For example, *Z-ligustilide*, presented in the essential oil of *Angelica sinensis*, can inhibit platelet aggregation, resist oxidation, reduce ischemic brain injury, and blood pressure, and so on.[9-11] *d-Limonene*, an active component of essential oil of *Fructus Aurantii*, has been reported possessing anti-inflammatory, anti-oxidative, lipid-lowering, and decreasing blood pressure activities.[12-14] Previous studies mostly have been focused separately on the essential oil from *Angelica sinensis* or *Fructus Aurantii*,[15,16] respectively. So far, to the best of our knowledge, none of the study on the composition and pharmacological effects of essential oil from DZHP has been reported.

Therefore, this article aimed to analyze the composition of the essential oil from DZHP and evaluate its analgesic activity and effect on promoting blood circulation. The analgesic activity was determined by two mice experimental models: acetic acid-induced writhing response and hot-plate latent pain response test.[17,18] The effect on activating blood circulation was evaluated by determining hemorheological parameters in rats with blood stasis syndrome.[19,20] Our result might provide scientific information to further understand the main components of the essential oil from DZHP, as well as its mechanism in relieving pain and activating blood circulation to dissipate blood stasis. It would be helpful for the clinical application of the essential oil from DZHP on blood stasis syndrome.

### MATERIALS AND METHODS

#### Materials and reagents

*Radix Angelica sinensis* was purchased from Dong Tang Pharmacy (Lot number 101218, Changsha, China), *Fructus Aurantii* was provided by Hunan Hansen Pharmaceutical Co. Ltd. (Lot number 101102, Yiyang, China). The two herbs were authenticated by Professor Ta-si Liu from the Hunan University of Chinese Medicine. Adrenaline hydrochloride injection was obtained from Wuhan Grand Pharmaceutical Group Co. Ltd. (Lot number 110307, Wuhan, China); TW-80 was supplied by Hunan Er Kang Pharmaceutical Co. Ltd. (Lot number 100608, Changsha, China). Acetic acid (analytical grade) was obtained from Changsha Reagent Company (Changsha, China).

#### Instrument and analytical conditions

The gas chromatography/mass spectrometry (GC/MS) analyses were performed on QP 2010 GC/MS instrument (Shimadzu, Japan) equipped with a flame ionization detector (FID), mass selective detector (MSD), and a Rtx-5ms capillary column (30 m × 0.25 mm, 0.25 μm). The initial oven temperature was held at 60°C for 2 min, and raised from 60 to 140°C at 10°C/min, and then heated to 200°C at 4°C/min, and held for 8 min; injector temperature, 250°C; FID temperature, 300°C; ion-source temperature, 200°C; transfer line temperature was scanned over 40-500 amu with an ionizing voltage of 70 eV and ionization current 150 mA. Samples (1.0 μL) were automatically injected into the GC/MS and Helium was used as the carrier gas (1.0 mL/min) with split mode (1:50).

#### Compound identification

Most constituents were identified by comparing their mass spectra with those in either a computer library or with the authentic compounds and were confirmed by comparing the Kovats retention indices with those of authentic samples or from the literature.[16, 22]

### Preparation of the essential oil sample

The essential oil from DZHP was obtained by hydrodistillation. A 480 g amount of herb pair, that is, *Radix Angelica sinensis* and *Fructus Aurantii* in the weight ratio of 1 : 2, was crushed to coarse powder. The mixture was then steam distilled for 6 h and the yield of essential oil was 2.27% (w/w). The essential oil samples administrated to animals were prepared by dissolving the essential oil in 1% (w/v) Polysorbate 80 (TW-80), and the essential oil concentration reached to 0.035 and 0.092 g/mL for mice and rats, respectively.

### Animals

Sprague–Dawley rats (220 ± 20 g) and Imprinting Control Region (ICR) mice (20 ± 2 g) were purchased from the Animal Center of Hunan University of Chinese Medicine [Certificate No. SCXK (Xiang) 2009-0004]. They were individually housed in plastic bottom cages with food and water provided ad libitum, and maintained at temperature 24 ± 2°C with 55 ± 5% relative humidity and a 12 h light/dark cycle. The animal experiments were approved by Animal Ethics Committee of Hunan University of Chinese Medicine.

### Acetic acid-induced writhing response

The acetic acid-induced writhing test was carried out according to the report.[17,18] 30 ICR mice, half male and half female, were divided into three groups of 10 mice each: control group (vehicle, 1% TW-80), positive group (aspirin, 0.12 g/kg), and the test drug group (essential oil of DZHP, 0.35 g/kg). They were administered vehicle or drugs for five days. Until half hour after the final administration, 0.6% acetic acid (0.1 mL/10 g) was intraperitoneally injected to them. The number of writhes was counted for 15 min after the acetic acid administration. The inhibition rate of writhes was calculated as follows: inhibition rate = [(control mean–test mean)/control mean] × 100.

### Hot plate test

The hot plate test was performed according to the method previously reported.[19, 20] The female ICR mice were divided into three groups (n = 10): control group (vehicle, 1% TW-80), positive group (aspirin, 0.12 g/kg), and the test drug group (essential oil of DZHP, 0.35 g/kg). Each female mouse was placed on the hot plate with a fixed temperature of 55 ± 0.5°C to observe the pain responses (hind-paw-licking or jumping). Before test each mouse was habituated twice to the hot plate. The response latency was defined as the recorded time in seconds between the platform and reaction. The mice were excluded when they exhibited latency time greater than 30 s or less than 5 s. They received continuous dosing for five days. After the final administration, the latency time was determined at 30, 60, and 90 min. A cut-off time of 60 s was fixed to prevent damage to the mice.

### Experiment on hemorheology in rats with blood stasis syndrome

The Sprague–Dawley rats, composed of male and female in half, were randomly divided into four groups with eight rats per group, control group, model group, positive medicine group (aspirin, 0.1 g/kg), and essential oil group (0.92 g/kg). The animals received continuous dosing for six days. Until 1 h after the final administration, except for the control group, the acute blood stasis models in the other three groups were established according to the report:[21] animals were subcutaneously injected at the intraperitoneal area with the adrenaline hydrochloride injection (0.8 mg/kg). The injection was administered twice, each separated by 4 h. Two hours before the second injection, animals were soaked in ice water for 5 min with their heads above the water. Rats were fasted overnight and administration continued after...
performed the model. Blood samples were collected 0.5 h after the last administration at the second day. The blood collection was carried out by carotid artery intubation and was collected into tubes at the ratio of 1:9 with 3.8% sodium citrate. The hematological parameters including whole blood relative index (WBRI) at high and low shear rate, whole blood reduced viscosity (WBRV) at high and low shear rate, and erythrocyte rigidity index (ERI) were detected by SA6000 automatic hemorheology meter. The plasma was separated from blood by centrifugation at 4000 rpm for 10 min and detected for plasma viscosity.

Statistical analysis
The results were analyzed by nonparametric Mann–Whitney U-test or one-way analysis of variance based on Dunnett’s multiple comparisons test. The differences were considered significant at \( P \) value less than 0.05.

RESULTS AND DISCUSSION
Composition of essential oil
The essential oil was yellow and fragrant liquid. The GC/MS chromatogram of the essential oil from DZHP is shown in Figure 1 and the main components in the mixtures are presented in Table 1. The presence of 28 compounds, accounting for 96.22% of the total oil composition, was detected by GC/MS, a main analytical technique adopted to quantification and quality control for the essential active substances of Chinese herb plants.

The constituents were identified by comparing their mass spectra with those in either a computer library or with the authentic compounds and were confirmed by comparing the Kovats retention indices with those of authentic samples or from the literature. As shown in Table 1, the main components were \( \alpha \)-pineine (3.07%), \( \beta \)-pineine (2.09%), \( \beta \)-myrcene (3.71%), D-limonene (49.28%), \( \gamma \)-terpinen (9.53%), \( \alpha \)-terpinolene (1.80%), \( \alpha \)-terpineol (2.02%), \( \beta \)-bisabolene (1.13%), butylidenehydralide (1.43%), and Z-ligustilide (16.08%). The oil was dominated by D-limonene, \( \gamma \)-terpinen, and Z-ligustilide, which made up 74.89% of the identified oil composition.

To our knowledge, there are reports on the chemical composition of the essential oil of Angelica sinensis and Fructus Aurantii, respectively. The main compounds in Angelica sinensis were ligustilide, \( \beta \)-pineine, and butylidenehydralide and so on.[22] Limonene, \( \alpha \)-pineine, \( \gamma \)-terpinen, carveol, terpineol, and linalool were the main compounds in Fructus Aurantii and the percentage of limonene was over 50%. In previous reports, little is known about the essential oil of the herb pair, which was obtained by extracting the two herbs commonly. In our present study, the essential oil of the herb pair was dominated by the presence of D-limonene at 49.28% and Z-ligustilide at 16.08%. It was consistent with the results of the previous study that the major constituents of the essential oil from Danggui-Zhiqiao herb-pair.

Table 1: Chemical composition of essential oil from Danggui-Zhiqiao herb-pair

| Compounds                 | RI*  | Percentage (%) | Identificationb |
|---------------------------|------|---------------|-----------------|
| \( \alpha \)-Thujene       | 902  | 0.61          | RI, MS          |
| \( \alpha \)-Pinene        | 943  | 3.07          | RI, MS, Co-I    |
| \( \beta \)-Pinene         | 976  | 2.09          | RI, MS, Co-I    |
| \( \beta \)-Myrcene        | 990  | 3.71          | RI, MS, Co-I    |
| D-Limonene                | 1018 | 49.28         | RI, MS, Co-I    |
| \( \gamma \)-Terpinen      | 1046 | 9.53          | RI, MS, Co-I    |
| \( \alpha \)-Terpinolene   | 1052 | 1.8           | RI, MS          |
| \( \beta \)-Linalool       | 1082 | 0.47          | RI, MS          |
| 3-Butyl-4-vinyl-1-cycloptene | 1092 | 0.32          | RI, MS          |
| 4-Carvomenthenol          | 1137 | 0.81          | RI, MS          |
| \( \alpha \)-Terpinolene   | 1143 | 2.02          | RI, MS, Co-I    |
| Carvone                   | 1190 | 0.08          | RI, MS          |
| Nerol acetate             | 1352 | 0.31          | RI, MS          |
| \( \beta \)-Elemen         | 1393 | 0.06          | RI, MS          |
| \( \beta \)-Cedrene        | 1398 | 0.15          | RI, MS          |
| Caryophyllene             | 1494 | 0.51          | RI, MS          |
| \( \beta \)-Bisabolene     | 1500 | 1.13          | RI, MS          |
| Caryophyllene oxide       | 1507 | 0.23          | RI, MS          |
| Chamigren                 | 1507 | 0.22          | RI, MS          |
| Germacrene                | 1515 | 0.74          | RI, MS          |
| Spathulenol               | 1536 | 0.59          | RI, MS          |
| \( \alpha \)-Cadinol       | 1580 | 0.25          | RI, MS          |
| Butylidenehydralide       | 1655 | 1.43          | RI, MS          |
| Z-Ligustilide             | 1741 | 16.08         | RI, MS, Co-I    |
| 1,4-(1'-hydroxycyclopentyl)-1,3-butadiyne | 1834 | 0.28      | RI, MS          |
| Hexadecanoic acid, methyl ester | 1878 | 0.14      | RI, MS          |
| n-Hexadecanoic acid, methyl ester | 1968 | 0.17  | RI, MS          |
| 8,11-Octadecadienoic acid, methyl ester | 2093 | 0.14   | RI, MS          |
| Total                     | 1027 | 96.22         |                 |

a) RI: Retention indices. b) Identification: MS, comparison of mass spectra with those listed in the NIST08 and mass spectral libraries; RI, comparison of the Kovats retention indices with those of authentic samples or from the literature.

Table 2: Effect of the essential oil from Danggui-Zhiqiao herb-pair on acetic acid-induced writhing response in mice

| Groups                   | Dose (g/kg, p.o.) | Numbers of writhing response* | \( P \)  | Rate of inhibiting (%)b |
|--------------------------|-------------------|-------------------------------|--------|------------------------|
| Control                  | -                 | 28.00 (23.25-35.75)           | -      | -                      |
| Aspirin                  | 0.12              | 13.00 (11.25-18.00)           | 0.001  | 53.57                  |
| Essential Oil            | 0.35              | 15.50 (11.00-21.50)           | 0.019  | 44.64                  |

a) Data are expressed as median (interquartile range) \( n = 10 \). b) Rate of inhibiting = \( [(control \text{ mean}-test \text{ mean})/control \text{ mean}] \times 100 \).
the two herbs were Z-ligustilide and D-limonene, respectively. The
γ-terpinene, β-pinene, α-pinene, β-myrcene, α-terpinolene, α-terpinol, β-bisabolene, and butylidenephthalide were the other important
products in the essential oil.

Acetic acid-induced writhing response
In analgesic study mice are widely used because of short reproduction
period, cheaper to purchase and maintain. Moreover, mice models
showed advantages of smaller quantity of agents required to induce
and/or treat disease. Therefore, analgesic studies were carried out
in mice in our research. The acetic acid-induced writhing test model
is a highly sensitive and useful test for screening analgesic drug and
has been widely regarded as a classical peripheral analgesic animal
model.[24] The results of the acetic acid-induced writhing response are
shown in Table 2. The essential oil, at the dose of 0.35 g/kg, showed
remarkable inhibition on the number of writhes induced by acetic
acid with inhibition rate of 44.64%. The positive control aspirin (0.12
g/kg) also very significantly inhibited the writhing response with
inhibition rate of 53.57%. Compared with the blank control group, the
results of the positive control aspirin and the tested essential oil were
statistically significant (P < 0.01 and P < 0.05). Several studies suggested
that acetic acid could result in an increase in peritoneal fluid levels of
prostaglandins (PGE₂ and PGF₂α) involving in part peritoneal receptors
and inflammatory pain by inducing capillary permeability.[25] The effect
of the essential oil of DZHP might inhibit the synthesis and/or the
release of the prostaglandins and may be mediated through inhibition
of cyclooxygenases and/or lipoxygenases.[25,26]

Hot plate latent pain response test in mice
The hot plate test, using thermal stimulus to induce pain, was proposed
by Eddy and Leimbach[27] and is commonly applied to evaluate its
centrally mediated antinociceptive activity. Significant analgesic efficacy
of the essential oil of DZHP was observed by the hot plate test and is
shown in Figure 2. Aspirin and the essential oil of DZHP significantly
increased pain threshold of mice compared with preadministration
(0 min) and a significant increase in hot plate latency occurred from
30 to 90 min after oral administration of drugs in mice compared with
control group (P < 0.01). In our study, the essential oil of DZHP could
significantly increase hot plate latency and this good analgesic activity
may be mediated through inhibiting central pain receptors.

Effect on hemorheology
According to report, [2] rats were selected to take part in hemorheology
test. Effect on hemorheology was observed and the results are presented
in Table 3. Compared with the control group, the model group had
conspicuous plasma viscosity, WBRI at high and low shear rate, WBRV
at high shear rate, and ERI, which indicated that the blood stasis model
was built successfully. After administration, these hemorheological
parameters significantly decreased in aspirin group and essential oil
group compared with the model group. Obviously, the essential oil from
DZHP can activate blood circulation to remove blood stasis.

Hemorheology is the study on the relation of blood flow to pressure,
flow volume, and resistance of blood vessels.[2] Once blood stasis
develops, hemorheology and microcirculation will be further affected.
It will lead to the elevation of blood viscosity, deterioration of
erthrocyte deformability, acceleration of erythrocyte aggregation, and
platelet aggregation, as well as microcirculatory dysfunction.[27] The
hemorheological parameters will display abnormal when blood stasis
occurs. In our study, hemorheological indexes were determined to
evaluate whether the essential oil of DZHP had the effect on promoting
blood circulation. The results demonstrated that the essential oil
of DZHP could significantly decrease hemorheological parameters
including plasma viscosity, WBRI, WBRV, and ERI compared with the
model group and had strong effect on activating blood circulation.

CONCLUSION
The composition of the essential oil of DZHP and its analgesic, promoting
blood circulation activity are reported for the first time. The essential
oil of DZHP was analyzed successfully by GC/MS, and D-limonene
(49.28%), Z-ligustilide (16.08%), and γ-terpinene (9.53%) were rich in
the oil. It had good analgesic and promoting blood circulation activities,
which may give evidences contributing to explain clinical application of
essential oil of DZHP on blood stasis syndrome and provide foundation
for clarifying the nature of the effect in the further study.

Table 3: Effect of essential oil from Danggui-Zhiqiao herb-pair on the hemorheology in rats with acute blood stasis syndrome

| Groups     | Dose (g/kg) | PVa (mPa.s) | WBRib | WBRVc (mPa.s) | ERId |
|------------|-------------|-------------|-------|---------------|------|
|            |             | High shear rate | Low shear rate | High shear rate | Low shear rate | Low shear rate | Low shear rate |      |
| Normal     | -           | 2.60 ± 0.16 | 3.68 ± 0.19 | 17.86 ± 1.16 | 13.20 ± 1.11 | 81.31 ± 4.07 | 5.22 ± 0.49 |
| Model      | -           | 2.88 ± 0.21* | 4.11 ± 0.49* | 19.56 ± 1.60* | 15.03 ± 1.60* | 85.96 ± 5.15 | 6.21 ± 0.98* |
| Aspirin    | 0.10        | 2.35 ± 0.26** | 2.72 ± 0.43** | 15.60 ± 1.47** | 8.69 ± 1.80** | 74.42 ± 4.43** | 3.78 ± 1.01** |
| Essential Oil | 0.92       | 2.51 ± 0.18** | 3.59 ± 0.53* | 17.38 ± 1.36** | 12.46 ± 2.47* | 78.72 ± 5.08** | 5.01 ± 1.17* |

* PV, Plasma viscosity; ** WBRI, Whole blood relative index; WBRV, α Whole blood reduced viscosity; α ERI, Erythrocyte rigidity index. Data are expressed as mean ± S.D. (n = 8); # P < 0.05 versus control group; * P < 0.05 versus model group; ** < 0.01 versus model group.
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Conflict of interest
There are no conflicts of interest.

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