Identification of compounds with antipyretic effects and anti-endotoxin activity in different species of *Lonicera japonica* using spectrum-effect correlation

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**Abstract.** Liquid chromatography (LC) is a common and straightforward approach used in the evaluation of the quality of Traditional Chinese Medicines (TCMs). Quality control is a critical step when systematically assessing the efficacy of TCMs. In the present study, the spectrum-effect correlation method was used to identify pharmacologically active substances. The aim of the present study was to investigate the underlying correlations between common chemical compounds with antipyretic effects and the anti-endotoxin activity of *Lonicera japonica*. The common chemical constituents of *Lonicera japonica* were analyzed using LC, and the antipyretic effects and anti-endotoxin activity were determined using ELISAs. Combining the results of bivariate and principal component analysis methods, eight active constituents were qualitatively and quantitatively analyzed. The results of these analyses indicated that neochlorogenic acid, chlorogenic acid, cryptochlorogenic acid and isochlorogenic acids A, B and C served a synergistic role with respect to antipyretic effects and anti-endotoxin activity. The present study lays a foundation for the future clinical use of *Lonicera japonica*.

**Introduction**

Traditional Chinese Medicine (TCM) has become increasingly popular worldwide, due to its reported therapeutic effects and low toxicity. As a result, it is vital to develop novel types of TCMs and to understand the active compounds used during their clinical application (1-3). Fever is a pathological phenomenon that causes the set point of the thermoregulatory center to increase, resulting in the body producing more heat than it dissipates, thus raising the body temperature (4). Heat sources can be divided into endogenous and exogenous. Endogenous heat sources are also known as leukocyte heat sources (such as interleukins and interferons), which can act on the thermoregulatory center through the blood-brain barrier, causing the thermoregulatory point to increase. Exogenous heat sources, causing the body temperature to increase (5,6). The animal body is extremely sensitive to lipopolysaccharide (LPS). Even small amounts of endotoxin (~5 ng/kg) can cause the body temperature to rise and fever to be maintained for >4 h (7,8). LPS is a component of the outer membrane of the cell wall of Gram-negative bacteria. It is released after the death of bacteria and has a wide range of biological activities. It can act on monocytes and macrophages to produce interleukins, tumor necrosis factor and other cytokines. These cytokines act on the thermoregulatory center in the hypothalamus, thereby increasing body temperature and causing inflammation, as well as other pathological changes (9-11).

The diversity of TCM arises from its multi-component and multi-target pharmacodynamic activity and treatment characteristics. Early research has focused on the potential antipyretic effect of compounds used in TCM and their...
underlying mechanism of action (12-18). Clarifying the efficacy and function of TCM is crucial to ensuring drug safety and quality control. The spectrum-effect correlation is studied to discern the correlation between fingerprint (chemical, biological and metabolic) with pharmacodynamic efficacy. This allows the identification of active components in TCM, and the formulation of standards to assess their internal quality (19). The spectrum-effect relationship has also been applied in multi-field research, such as single herb treatment, classic couplet medicines, compound compatibility, processing mechanism and technology optimization (20-24). In addition, it can be used to identify key substances, optimize the ratio of ingredients, improve preparation technology and track various characteristics, providing insight into novel approaches for the development of novel TCM drugs (19,25-31).

*Lonicera japonica* Flos is prepared from a dried flower bud or flower collected from the first harvest of *Lonicera japonica* blooms. Its chemical components include organic acids, flavonoids, triterpenoid saponins, iridoid terpenes, volatile oils and trace elements (32). The principal active ingredients are chlorogenic acid and luteolin, which may be quantified to ensure quality control of the preparation (33). Previous studies have reported that *Lonicerae japonica* Flos has numerous pharmacological effects, such as broad-spectrum antibacterial, antiviral, immunemychenying, antioxidant, antipyretic and anti-inflammatory effects, as well as liver protection, hypoglycemia and anti-tumor activity (32,34-36). *Lonicera japonica* Thunb. (LJT), *Lonicera fulvotomentosa* Hsu et S. C. Cheng (LFH) and *Lonicera macranthoides* Hand.-Mazz. (LMH) represent different species of *Lonicerae japonica* Flos. Although their efficacy is the same, their chemical compositions differ (37).

The quality of TCM drugs has a direct impact on their efficacy (31). In the present study, LPS endotoxin was injected intraperitoneally to successfully establish a rat model of pyrotoxemia. Physiological and biochemical indexes, such as cAMP and arginine vasopressin (AVP) levels in the hypothalamus and serum endotoxin (ET) levels were measured, as previously described (38-40). The antipyretic effect and the anti-endotoxin activity of different species of *Lonicera japonica* were also examined. The present study provided a theoretical basis for the molecular mechanism underlying the antipyretic effects and anti-endotoxin activity of different species of *Lonicera japonica* and the development of safer and more effective antipyretic TCM drugs.

**Materials and methods**

**Instruments, reagents and animals.** An Agilent 1260 HPLC system (Agilent Technologies, Inc.) with a diode array detector (DAD) was used for analytical chemistry experiments and data processing. Milli-Q ultrapure water (Merck KGaA) was used for the preparation of test samples of standards. Additional instruments included the LGJ-12 Freeze dryer (Beijing Songyuan Huaxing Technology Development Co., Ltd.), the 011 Electronic thermometer (Henan Muxiang Veterinary Pharmaceutical Co. Ltd.), the 1510 microplate reader (Thermo Fisher Scientific, Inc.) and the 5415D tabletop high-speed centrifuge (Eppendorf).

LJT, LFH and LMH were collected from different villages in Guizhou (China) and identified by Professor Wei Shenghua (Guizhou University of Traditional Chinese Medicine). These samples were: i) LJT (sample no. 20170610; Guizhou, China); ii) LFH (sample no. 20170612; Qingzhou, China); and iii) LMH (sample no. 20170611; Daozhen, China). The 12 different villages of LJT were as follows: 1a-Xingtai, 1b-Xingtai, 2-Xinxiang, 3-Huihua, 4a-Suiyang, 4b-Suiyang, 4c-Suiyang, 4d-Suiyang, 5a-Linyi, 5b-Linyi, 5c-Linyi and 6-Xiushan. The 30 different villages of LFH were as follows: 1-Haiyan, 2-Baling, 3-Maoying, 4a-Yingpan, 4b-Yingpan, 4c-Yingpan, 5a-Yingtawon, 5b-Yingtawon, 11-Gantian, 12-Zhengchang, 13-Huangyangtai, 14-Yongle, 15-Yuxi, 16-Dagan, 17-Hekou, 18-Daba, 19-Zhouzhu, 20-Shichao, 21-Dejiang, 22-Sinan, 23-Dushan, 24-Tianzhu, 25a-Songtao, 25b-Songtao and 26-Xiaoguan. The 12 different villages of LMH were as follows: 1a-Xingren, 1b-Xingren, 2-Huilong, 3-Mamaya, 4-Xingyi, 5-Pengzuo, 6-Dewo, 7-Waina, 8-Hongni, 9-Zhenfeng, 10-Qingzhen and 11-Anlong.

The standards used were as follows: i) Neochlorogenic acid (NCA; cat. no. PS0601-0025); ii) chlorogenic acid (CA; cat. no. PS0131-0025); iii) cryptochlorogenic acid (CCA; cat. no. PS0775-0025); iv) secologanin (ca. no. PS2210-0025); v) secologanic acid (SL; cat. no. PS2215-0020); vi) isochlorogenic acid B (ICAB; cat. no. PS0607-0025); vii) isochlorogenic acid A (ICAA; cat. no. PS0066-0025); and viii) isochlorogenic acid C (ICAC; cat. no. PS0068-0025). All standards were obtained from Chengdu Push Bio-technology Co., Ltd.). The purity of each standard was ≥98. Acetonitrile, methanol and formic acid were of chromatographic grade, whereas the water used was ultrapure. LPS was purchased from Hefei Domei Biology Co., Ltd. (cat. no. L2880; Sigma-Aldrich; Merck KGaA). The cAMP (cat. no. LE-H3962), AVP (cat. no. LE-H1112) and ET (cat. no. LE-H8973) ELISA kits (rat) were obtained from Heihei Laier Biological Technology Co., Ltd.

Specific pathogen free Sprague-Dawley rats (age, 2 months, n=96), weighing an average of 245±15 g (48 males and 48 females), were purchased from Chengdu Animal Health Supervision Institute. The rats were housed in a temperature-controlled room (22±2˚C) with 60±10% humidity on a 12-h light/dark cycle and unrestricted access to food and water.

**Preparation of the sample solution.** LJT, LFH and LMH extracts were prepared using distilled water as the solvent in a 1:20 mass to volume ratio. The extracts were incubated three times at 100˚C for 90 min each time, filtered twice, then dried at 60˚C after concentration.

With a volume of 20 ml/kg, LJT, LFH and LMH solution samples were prepared in distilled water for oral administration. The high dose was 20 g/kg body weight (equivalent to 4 times the clinical dose), the medium dose was 10 g/kg body weight (twice the clinical dose) and the low dose was 5 g/kg body weight (clinical dose). With a volume of 20 ml/kg and the crude drug dosage of 6.75 g/kg, *banlangen granules (Radixisatidis extract)* (Guanzhou Baiyunshan Pharmaceutical Holdings Co., Ltd.) was prepared for oral administration as a positive control. The blank and model groups were given equal volumes of distilled water.
LC conditions. An Agilent Eclipse XDB-C18 column (3.9x150 mm; 5 µm; Agilent Technologies, Inc.) was used to analyze the samples. Data analysis was performed on an Agilent 1260 HPLC system (Agilent Technologies, Inc.) equipped with a diode array detector. The mobile phase consisted of acetonitrile (A) and 0.4% (v/v) acetic acid aqueous solution (B), and was pumped at a flow rate of 0.8 ml/min. The column temperature was maintained at 35°C. The injection volume of each sample was 10 µl. The chromatograms were monitored at 254 nm. The gas was N2, and its flow rate was 1.60 SLM with the DAD.

The gradient elution program of LJT was as follows: 5-6% A, 0-10 min; 6% A, 10-12 min; 6-7% A, 12-20 min; 7-8% A, 20-25 min; 8-10% A, 25-30 min; 10-12% A, 30-35 min; 12-14% A, 35-40 min; 14% A, 40-45 min; 14-16% A, 45-50 min; 16% A, 50-55 min; 16-18% A, 55-60 min; 18-20% A, 60-65 min; 20-30% A, 65-70 min; 30-40% A, 70-75 min; 40-60% A, 75-80 min; 60-5% A, 80-85 min; and 5% A, 85-90 min.

The gradient elution program of LMH was as follows: 5-10% A, 0-10 min; 10-15% A, 10-20 min; 15% A, 20-25 min; 15-20% A, 25-30 min; 20-25% A, 30-40 min; 25-30% A, 40-50 min; 30-40% A, 50-60 min; 40% A, 60-70 min; 40-5% A, 70-75 min; and 5% A, 75-80 min.

The gradient elution program of LFH was as follows: 5-10% A, 0-10 min; 10-15% A, 10-20 min; 15% A, 20-25 min; 15-20% A, 25-30 min; 20-25% A, 30-40 min; 25-30% A, 40-50 min; 30-40% A, 50-60 min; 40% A, 60-70 min; 40-5% A, 70-75 min; and 5% A, 75-80 min.

For quantitative analysis, the mobile phase was pumped at a flow rate of 1.0 ml/min. The column temperature was maintained at 30°C. In addition, the gradient elution program was the same as LMH. All other conditions were kept the same.

Fingerprint evaluation. LC fingerprints of the samples collected from different species were established and matched automatically using the Similarity Evaluation System for Chromatographic Fingerprint of TCM (version 2012; China Pharmacopoeia Committee). Furthermore, cluster analysis using SPSS (version 22; IBM Corp.) was applied to evaluate the quality of the samples from different species.

Principal component analysis (PCA). PCA is a multivariate statistical method that can retain sufficient information from data acquisition. In the present study, characteristic peaks from LC chromatograms were screened using PCA in order to identify active constituents based on spectrum-effect relationships. PCA was conducted using SPSS software (version 22.0; IBM Corp.).

Establishment of rat models of LPS toxemia. Animal experimentation was initiated following 3 days of acclimatization. The rats were divided into 12 groups (8 per group, with equal numbers of males and females). Groups 1-9 received high, medium and low doses of water extracts from LJT, LFH and LMH. The high dose was 27 g/kg body weight (20 times the clinical dose), the medium dose was 13.5 g/kg (10 times the clinical dose) and the low dose was 6.75 g/kg (5 times the clinical dose). Group 10 was the fever model group, which was administrated water extract by gavage daily for 6 days. Group 12 (blank) received no treatment and group 11 (positive control) received low dosage normal saline and 6.75 g/kg Banlangen Granules, which were administered by gavage daily for 6 days. The rats were fasted overnight with free access to water prior to administration of the test solutions. On day 7, the model was considered to be established successfully when after 1 h, 100 µg/kg LPS endotoxin was injected into the abdominal cavity of rats in groups 1-11. The protocol was approved by the Animal Ethical Committee of Guizhou University. All animals were treated according to the guidelines of the National Institutes of Health (41). All procedures were performed under sodium pentobarbital anesthesia (intraperitoneal injection at a dose of 50 mg/kg).

Tissue lysate ELISA. Following anesthesia the rats were sacrificed by decapitation 6 h after the peak of fever. The whole brain and the hypothalamus tissue were removed quickly and placed in an ice bath. Then, 0.5 ml cell lysis buffer (cat. no. 87792; Thermo Fisher Scientific, Inc.) was added to 50 mg fresh hypothalamus tissue. After lysis at room temperature for 30 min, sonication was continued for 1 min. The samples were centrifuged (3,662 x g; 15 min; 4°C), and the supernatants were separated for testing. In each standard well, 50 µl standard of different concentrations were added. In each sample well, 10 µl testing sample and 40 µl sample diluent were added. In blank wells, nothing was added. In all wells except the blanks, 100 µl horseradish peroxidase (HRP) labeled detection antibody was added. cAMP (cat. no. LE-H3962), AVP (cat. no. LE-H1112) and ET (cat. no. LE-H8973) ELISA kits (rat) were obtained from Heifei Lyle Biotech Co., Ltd. The protocol was in accordance with the manufacturer’s instructions. The plates were sealed with sealing film, then incubated at 37°C for 1 h. After discarding the excess liquid, each well was filled with detergent five times for 1 min each time. In each well, 50 µl substrate A and 50 µl substrate B were added to each well, and the plates were incubated at 37°C in the dark for 15 min. After incubation, 50 µl termination solution was added to each well. After 15 min at 37°C, the optical density (OD) value of each well was measured using a 1510 microplate reader at 450 nm. A standard curve was drawn using the OD values obtained from the standards. According to this curve equation, the concentration of each sample was calculated.

Serum ELISA. Rats were anesthetized with sodium pentobarbital 6 h after the peak of fever, and 5 ml blood was obtained from the abdominal aorta. The samples were centrifuged (845 x g; 30 min; 4°C) in order to obtain serum for testing. The concentrations of ET were determined using a double antibody sandwich ELISA, for which the determination method was the same as that of cAMP (cat. no. LE-H3962) and AVP (cat. no. LE-H1112). All kits were obtained from Heifei Lyle Biotech Co., Ltd.
Statistical analysis. Two-tailed paired t-tests and one-way ANOVA followed by Tukey's post hoc test were used to identify statistically significant between groups. P<0.05 (95% confidence interval) was considered to indicate a statistically significant difference. Pearson's correlation coefficient was used to calculate the relative peak areas, which corresponded with antipyretic effect and anti-endotoxin activity of the samples. Correlation coefficients >0.3 were considered significant (P<0.05), whereas correlation coefficients >0.5 were considered highly significant (P<0.01). PCA was used to evaluate characteristic peak areas in the chromatograms of the samples from different species. Bivariate analysis was used to assess the correlation of peak areas with their antipyretic effect and anti-endotoxin activity. Statistical analysis was conducted using SPSS software (version 22.0; IBM Corp.). Data are presented as the mean ± SD and 6 experimental repeats of each test were performed.

Results

Establishment and evaluation of fingerprint. The relative standard deviation of the retention time and peak area of characteristic peaks were 0.041 and 0.082% for precision, 0.390 and 0.173% for reproducibility, and 0.098 and 0.103% for stability. The similarities of the chromatograms were all >0.9. These findings indicated that the present LC method of fingerprint establishment was reliable.

Cluster analysis. The results of cluster analysis are presented in Fig. 1. The 12 different batches of LJT were divided into two categories. The first category consisted of 4b, 6, 5c, 2, 5a, 4c, 1b and 1a. The second category consisted of 5b, 3, 4d and 4a. The 30 different batches of LFH were divided into two categories. The first category consisted of 18, 25, 15, 22, 8, 11, 6, 13, 4, 20, 7, 3, 19 and 16. The second category comprised 9b, 2, 14, 24, 12, 17, 9a, 23, 9c, 1, 21, 5, 26, 25b, 10b and 10a. The 12 different batches of LMH were divided into two categories as follows: i) a; and ii) 11, 10, 6, 9, 4, 1b, 8, 7, 2, 5 and 3. Although they were clustered into two categories, the differences were not significant, which indicated that there was no marked difference between different species of LJT, LFH and LMH.

Qualitative and quantitative analysis of the eight main chemical constituents. Under quantitative analysis conditions, the linear regression equations for NCA, CA, CCA, SLA, SL, ICAB, ICA and ICAC were as follows: i) NCA, \(Y=1104.2X+26.119\) (r=1) in the concentration range of 1.50-13.50 µg/ml; ii) CA, \(Y=1105.4X+15.676\) (r=1) in the concentration range...
of 2.59-17.28 µg/ml; iii) CCA, Y=973.46X+5.8348 (r=1) in the concentration range of 0.72-12.96 µg/ml; iv) SLA, Y=598.22X+28.842 (r=0.9998) in the concentration range of 0.76-12.96 µg/ml; v) SL, Y=509.85X-2.865 (r=1) in the concentration range of 1.04-9.36 µg/ml; vi) ICAB, Y=1277.9X+2.5354 (r=1) in the concentration range of 2.14-19.26 µg/ml; vii) ICAA, Y=1169.4X-1.0839 (r=1) in the concentration range of 0.84-15.12 µg/ml; and viii) ICAC, Y=1588.8X+8.6825 (r=1) in the concentration range of 2.24-21.78 µg/ml. These results all indicated a linear relationship (Table I). The structures of eight main chemical constituents identified in different species of LJT, LFH and LMH are presented in Fig. 2.

Determination of the antipyretic effect and anti-endotoxin activity. There was a highly significant difference in body temperature between the normal groups and the model control group (Table II), indicating that the rat endotoxin model was successfully established. Compared with the model group, there was significant difference in antipyretic effect in each experimental group. The extract significantly inhibited the increase in cAMP and its secretion, thus reducing the heat output of the body. Compared with the model group, there were highly significant differences in the experimental groups 2, 3, 5, 6, 7, 8 and 9 (Table II). This indicated that the extract promoted an increase in the levels of the anti-diuretic hormone AVP to different degrees. Compared with the model group, there was a highly significant difference in all 9 experimental groups. Thus, the extract had an anti-endotoxin effect in vivo.

PCA. As presented in Table III, the eigenvalues of the first two principal components were >1, and the contribution rate of the first principal component was 67.447%, indicating that the first principal component was able to explain 67.447% of the antipyretic effect of different species of LJT, LFH and LMH. The second principal component could explain 32.553%, and the cumulative contribution rate of the first two principal components was 100%. Therefore, the first two principal components were further evaluated. Table III presents the eigen values and factor loadings of these two principal components. The first principal component exhibited significant load based on the eigenvalues of NCA, CA, CCA, ICAB, ICAA and ICAC (>0.8). Consequently, these six constituents were selected as representative variables for the antipyretic effect.

As presented in Table IV, the eigen values of the first two principal components were >1, and the contribution rate of the first principal component was 82.246%. The results indicated that the first principal component could explain 82.246% of the anti-endotoxin activity of different species of LJT, LFH and LMH. The second principal component could explain 17.754%, and the cumulative contribution rate of the first two principal components was 100%. Therefore, the first two principal components were evaluated further. Table IV indicates their eigen values and factor loadings. The first principal component exhibited significant load based on the eigenvalues of the six constituents (>0.8). Consequently, NCA, CA, CCA, ICAB, ICAA and ICAC, were selected as representative variables for anti-endotoxin activity.

Bivariate analysis (BA). The antipyretic effect and ant-endotoxin activity of these extracts are indicated in Figs. 3 and 4. Based on these results, a total of eight constituents significantly correlated with the concentration of cAMP (P<0.05), whereas seven constituents highly significantly correlated with the concentration of cAMP (P<0.01); a total of seven constituents significantly correlated with AVP levels (P<0.05), whereas four constituents highly significantly correlated with AVP levels (P<0.01). The constituents that significantly correlated with the concentrations of cAMP and AVP were NCA, CA, CCA, SL, ICAB, ICAA and ICAC (Fig. 3). These seven constituents exerted antipyretic effects. A total of eight constituents highly significantly correlated with ET levels (P<0.01). The constituents that significantly correlated with ET content were NCA, CA, CCA, SL, ICAB, ICAA and ICAC (Fig. 4). Therefore, these eight constituents were demonstrated to exert anti-endotoxin activity. The comprehensive findings of the present study verified that different species of LJT, LFH and LMH exhibited antipyretic effects and anti-endotoxin activities.

Discussion

To obtain a better chromatogram map, methanol-water, acetonitrile-water, methanol-0.15% formic acid, acetonitrile-0.15% formic acid, acetonitrile-0.4% acetic acid, acetonitrile: Methanol (1:1)-0.4% acetic acid and acetonitrile-0.4% phosphoric acid were investigated and screened as mobile phases. The combination of acetic acid (0.4%, v/v)-acetonitrile was the best mobile phase for separation and analysis. In the range of 190-400 nm full wavelength scanning, 254 nm was selected as optimal. The column temperature was set at 25, 30 and 35°C, respectively. A temperature of 35°C was selected. Flow rates of 0.8, 1.0 and 1.2 ml/min were also tested, and peak resolution of the samples was improved when the flow rate was 0.8 ml/min. Lastly, injection volumes of 10, 15 and 20 µl were also compared, and peak resolution was optimized when the injection volume was 10 µl.

TCM has the characteristics of multi-component, multi-effect and collaborative integration. The composition of its components and the interactions between them is

| Compound | LJT (mg/g) | LFH (mg/g) | LMH (mg/g) |
|----------|------------|------------|------------|
| NCA      | 19.33±3.35 | 11.05±0.34 | 33.62±0.91 |
| CA       | 48.48±8.05 | 19.81±0.60 | 64.78±1.86 |
| CCA      | 26.05±4.27 | 18.40±0.56 | 47.49±1.53 |
| SL       | 29.42±5.97 | 29.94±0.95 | 0.00       |
| ICAB     | 12.92±2.32 | 39.90±0.88 | 27.50±1.21 |
| ICAA     | 13.85±2.28 | 30.89±0.84 | 17.22±0.32 |
| ICAC     | 24.58±4.22 | 52.55±1.44 | 25.60±0.54 |

LJT, *Lonicera japonica* Thunb.; LFH, *Loniceranum przewalskii* Hsiao et S. C. Cheng; LMH, *Loniceranum przewalskii* Hsiao et S. C. Cheng; NCA, neo-chlorogenic acid; CA, chlorogenic acid; CCA, cryptochlorogenic acid; SL, secologanic acid; SL, secoxyloganin; ICAB, isochlorogenic acid B; ICAA, isochlorogenic acid A; ICAC, isochlorogenic acid C.
complex. It is important to identify a suitable research method to determine the efficacy of TCM and its mechanism of action. At present, the most common research method is to determine pharmacodynamics and assess the correlation of its ‘spectrum-effect’ (19-29). Pearson’s correlation coefficient was used in the current study to calculate the relative peak areas, which corresponded to the antipyretic effect and anti-endotoxin activity of the samples. Based on these results, different species of LJT, LFH and LMH exhibited antipyretic effects and anti-endotoxin activities. Correlation coefficients >0.3 or <0.3 were considered significant (P<0.05), whereas correlation coefficients >0.5 or <0.5 were considered highly significant (P<0.01). Based on the comprehensive results of PCA and BA, these eight constituents (NCA, CA, CCA, SLA, SL, ICAB, ICAA and ICAC) were demonstrated to exert anti-endotoxin activity.

LC technology was used for the qualitative and quantitative analysis of eight main chemical constituents in different species of LJT, LFH and LMH. The present study illustrated how six constituents, namely NCA, CA, CCA, ICAB, ICAA and ICAC, were selected as representative variables for the antipyretic effect and anti-endotoxin activity. The constituents that significantly correlated with the concentration of cAMP and AVP were NCA, CA, CCA, SL, ICAB, ICAA and ICAC. The constituents that significantly correlated with ET levels were NCA, CA, CCA, SLA, SL, ICAB, ICAA and ICAC. Therefore, NCA, CA, CCA, ICAB, ICAA and ICAC were demonstrated to be the key substances that mediate the

Table II. cAMP, AVP and ET content in rats from different experimental groups (n=8 rats/group).

| No | Group            | cAMP content (nmol/ml) | AVP content (pg/ml) | ET content (EU/ml) |
|----|------------------|------------------------|---------------------|-------------------|
| 1  | LJT (6.75 g/kg)  | 24.40±1.598<sup>a,d</sup> | 23.28±2.12<sup>b</sup> | 164.13±12.98<sup>b,c</sup> |
| 2  | LJT (13.5 g/kg)  | 21.85±0.669<sup>b</sup>  | 24.96±0.91<sup>b,c</sup> | 146.62±10.03<sup>b,c</sup> |
| 3  | LJT (27 g/kg)    | 19.73±1.084<sup>b,c</sup> | 27.78±1.58<sup>b</sup>  | 131.15±10.51<sup>b,c</sup> |
| 4  | LFT (6.75 g/kg)  | 24.85±1.058<sup>a</sup>  | 24.22±1.87<sup>b</sup>  | 178.34±13.83<sup>b,c</sup> |
| 5  | LFT (13.5 g/kg)  | 22.16±0.934<sup>b</sup>  | 25.88±2.59<sup>b</sup>  | 159.20±14.28<sup>b</sup>  |
| 6  | LFT (27 g/kg)    | 20.66±0.762<sup>b</sup>  | 27.20±3.08<sup>b</sup>  | 146.60±5.29<sup>b</sup>   |
| 7  | LMT (6.75 g/kg)  | 22.45±1.719<sup>b</sup>  | 23.69±2.27<sup>b</sup>  | 153.23±12.93<sup>b</sup>  |
| 8  | LMT (13.5 g/kg)  | 20.63±1.068<sup>b</sup>  | 25.62±2.07<sup>b</sup>  | 138.38±6.33<sup>b</sup>   |
| 9  | LMT (27 g/kg)    | 18.71±1.553<sup>b</sup>  | 27.33±1.73<sup>b</sup>  | 122.82±10.44<sup>b</sup>  |
| 10 | Model (6.75 g/kg)| 25.95±1.175<sup>a</sup>  | 17.88±3.26<sup>b</sup>  | 208.68±12.87<sup>c</sup>  |
| 11 | Positive (6.75 g/kg) | 20.64±1.421<sup>b</sup>  | 28.59±1.74<sup>b</sup>  | 143.78±9.94<sup>b</sup>   |
| 12 | Blank (6.75 g/kg)| 22.84±0.677<sup>b</sup>  | 22.92±1.83<sup>b</sup>  | 102.80±8.23<sup>b</sup>   |

<sup>a</sup>P<0.05 and <sup>b</sup>P<0.01 vs. model control group; <sup>c</sup>P<0.01 and <sup>d</sup>P<0.05 vs. blank control group. LJT, Lonicera japonica Thunb.; LFH, Lonicera fulvotomentosa Hsu et S. C. Cheng; LMH, Lonicera macranthoides Hand.-Mazz.; AVP, arginine vasopressin; ET, endotoxin.
Table III. Eigen values and corresponding percentages of the variables of antipyretic effect (top two principal components).

| Component | Total | Variance (%) | Cumulative (%) | Total | Variance (%) | Cumulative (%) |
|-----------|-------|--------------|----------------|-------|--------------|----------------|
| 1         | 9.443 | 67.447       | 67.447         | 9.443 | 67.447       | 67.447         |
| 2         | 4.557 | 32.553       | 100.000        | 4.557 | 32.553       | 100.000        |

Figure 3. Bivariate analysis of eight main chemical constituents identified in different species of *Lonicera japonica* Thunb., *Lonicera fulvotomentosa* Hsu et S. C. Cheng and *Lonicera macranthoides* Hand-Mazz. with regards to the antipyretic effect. Correlation coefficients >0.3 or < -0.3 were considered significant (P<0.05), whereas correlation coefficients >0.5 or < -0.5 were considered highly significant (P<0.01). NCA, neochlorogenic acid; CA, chlorogenic acid; CCA, cryptochlorogenic acid; SLA, secologanic acid; SL, secoxyloganin; ICAB, isochlorogenic acid B; ICAA, isochlorogenic acid A; ICAC, isochlorogenic acid C; AVP, arginine vasopressin.

Table IV. Eigen values and corresponding percentages of the variables of anti-endotoxin activity (top two principal components).

| Component | Total | Variance (%) | Cumulative (%) | Total | Variance (%) | Cumulative (%) |
|-----------|-------|--------------|----------------|-------|--------------|----------------|
| 1         | 9.047 | 82.246       | 82.246         | 9.047 | 82.246       | 82.246         |
| 2         | 1.953 | 17.754       | 100.000        | 1.953 | 17.754       | 100.000        |

Figure 4. Bivariate analysis of the eight main chemical constituents identified in different species of *Lonicera japonica* Thunb., *Lonicera fulvotomentosa* Hsu et S. C. Cheng and *Lonicera macranthoides* Hand-Mazz. with regards to anti-endotoxin activity. Correlation coefficients >0.3 or < -0.3 were considered significant (P<0.05), whereas correlation coefficients >0.5 or < -0.5 were considered highly significant (P<0.01). NCA, neochlorogenic acid; CA, chlorogenic acid; CCA, cryptochlorogenic acid; SLA, secologanic acid; SL, secoxyloganin; ICAB, isochlorogenic acid B; ICAA, isochlorogenic acid A; ICAC, isochlorogenic acid C; ET, endotoxin.
antipyretic effect and anti-endotoxin activity of different species of *Lonicera japonica*.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

JXD wrote the manuscript and analyzed data. YZ designed the experiments. WNY established and evaluated the finger print. WPL carried out the ELISA experiments. HS and JXL performed the animal experiments. CLT recorded and analyzed the experimental data. CL and XWY analyzed the data. All authors read and approved the final manuscript.

Ethics approval and consent to participate

All experimental procedures were reviewed and approved by the Animal Ethics Committee of Guizhou University (approval no. GZU-2019-11).

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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