INFLUENCE OF HONEY-ROASTING ON THE MAIN PHARMACOLOGICAL ACTIVITIES AND THE WATER-SOLUBLE ACTIVE GLYCOSIDES OF LICORICE

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Abstract

In traditional Chinese medicine (TCM), licorice is usually processed with honey and traditionally used in decoction form. However, the influence of honey-roasting on the main pharmacological activities and the water-soluble active constituents of licorice has not been reported. The aim of the present study is to determine whether honey-roasting can modify the main pharmacological activities and the active constituents of licorice. According to licorice clinical application and processing method, the mainly related pharmacological activities of crude licorice, processed licorice and refined honey, such as enhancing immune function, relieving cough, eliminating phlegm and detoxication, were compared. The results showed that honey-roasting obviously reinforced the licorice activity of enhancing Qi-deficiency mice’s immune function, and significantly weakened the licorice activity of relieving cough, removing phlegm and detoxication. However, honey didn’t show the significant activity of relieving cough, removing phlegm and detoxication. The influence of honey-roasting on the chemical compositions in licorice slice and licorice decoction was investigated by using HPLC. The results showed that the content and the decocting quantity of mainly 5 active glycosides in licorice, i.e. liquiritin apioside, liquiritin, licuraside, isoliquiritin and glycyrrhizin, obviously changed after processing; glycyrrhizin and liquiritin obviously decomposed during honey-roasting. In conclusion, honey-roasting obviously modified the main pharmacological activities and the water-soluble compositions of licorice. The modification was not caused by honey only. This finding may shed some light on understanding the differences in the therapeutic values of crude and processed licorice.

Key words: crude licorice, processed licorice, honey-roasting, pharmacological activity, active glycosides, modification.

Introduction

Licorice, a very famous herbal medicine, has been used for more than 2 millennia in China and referred as the “King of Herb” (Shang et al., 2010). Licorice is usually administered as decoction, and commonly classified into two decoction slices, i.e. processed licorice (roasting with refined honey), crude licorice (China Pharmacopoeia Committee, 2005). According to TCM theory, the efficacies of crude licorice and processed licorice are very different. Crude licorice is good at relieving cough, dissipating phlegm and detoxication, and usually used to curing bronchitis, sore throat and intoxication caused by drugs and food, while processed licorice is expert in invigorating vital energy, and usually used to treat immune deficit, dyspepsia and arrhythmia (Gong, 2007).

Modern research revealed that the roasted licorice (roasted without honey) had the stronger effect in antianaphylaxis (Majima et al., 2004), anti-inflammation (Kim et al., 2006; Kim et al., 2010), neuroprotection (Hwang et al., 2006) and anti-diabetic action (Ko et al., 2007) than crude licorice. In recently years, pharmacological research also showed that processed licorice had the better effect of anti-arrhythmia (Huang et al., 1984) and analgesia (Peng et al., 1989) than crude licorice. However, the research about the influence of roasting with honey on the main efficacy of licorice, such as strengthening immunity, relieving cough, dissipating phlegm and detoxification, and the processing influence on the water-soluble active constituents of licorice, has not been reported. In the present study, therefore, the mainly pharmaceutical evaluation and the chemical analysis of licorice before and after honey-roasting were carried out. The results showed that honey-roasting could obviously modify the main pharmacological activities and chemical compositions of licorice.

Materials and Methods

Animals

Kunming albino mice (clean grade), with an average body mass of 20.0 ± 2.0 g at the time of study initiation, provided by the Experiment Animal Center of Fudan University (Shanghai, Certificated number: SCXK (Hu) 2007-0026). Animals were kept under standard environmental conditions with free access to rodent diet and water. The animal experimental protocol was approved by the Institutional Animal Care and Use Committee of School of Pharmacy, Shanghai Jiao Tong University.

Materials and agents

Licorice was purchased from Inner Mongolia Yili Technology Company (Batch Number: GY 07020510) and identified as the root of Glycyrrhiza uralensis Fisch by one of authors-Prof. Xiaobo Li. The voucher specimen (SJTU
2007-09-18) was deposited in School of Pharmacy, Shanghai Jiao Tong University. Refined honey was provided by Shanghai Jing’an Drug Manufactory. Reference standards, liquiritin apioside, liquiritin, licuraside, isoliquiritin, glycyrrhizin, liquiritigenin, glycyrrhetic acid-3-O-β-D-glucuronide and glycyrrhetic acid were isolated from licorice by the authors (Figure 1). The relative purities were all more than 95% by HPLC analysis. Indian ink was provided by Shanghai Maisha Biotechnology Company and diluted to 5 times before use. Sodium bicarbonate, ammonia water, phenol red, chloral hydrate, methanol, methanoic acid and phosphoric acid were analytical grade from Shanghai National Pharmaceutical Co. Ltd. Acetonitrile was HPLC grade from Shanghai Xingke Biochemistry Co., Ltd (Shanghai, China). Water was purified by Milli-Q system (Massachusetts, USA).

\[ \text{glycyrrhetic acid} \quad R=H \]
\[ \text{glycyrrhetic acid-3-O-glucuronide} \quad R=\text{GluA} \]
\[ \text{glycyrrhizin} \quad R=\text{GluA (2-1)-GluA} \]
\[ \text{liquiritigenin} \quad R=H \]
\[ \text{liquiritin} \quad R=\text{Glu} \]
\[ \text{liquiritin apioside} \quad R=\text{Glu-(2-1)-Api} \]
\[ \text{isoliquiritin} \quad R=\text{Glu} \]
\[ \text{licuraside} \quad R=\text{Glu-(2-1)-Api} \]

**Figure 1:** Chemical structures of 8 compounds

**Instruments**

Agilent 1200 series HPLC system consisted of quaternary pump, on-line degasser, autosampler, thermostatic column compartment and DAD (Agilent, Germany), 754 UV-VIS spectrophotometer (Shanghai Precision Scientific Instrument Corporation, China), BS300S analytic balance (Sartorius, Germany), transferpetto (Eppendorf, Germany), 402 ultrasonic nebulizer (Shanghai Heli Medical Apparatus and Instruments Factory, China), DZF-6050 vacuum drying oven (Shanghai Yiheng Scientific Instrument Corporation, China).

**Sample preparation**

**Licorice processing:** Each 100 g of crude licorice slice was macerated with 25 g refined honey in 50 ml water for 8 h, then roasted at 180 °C in drying oven for 1 h (China Pharmacopoeia Committee, 2005).

**Sample for pharmacological test:** 500 g crude and processed licorice were decocted 2 times in 5000 ml of distilled water for 30 min, respectively. The decoctions were filtered and condensed to 1.0 g/ml. The honey was diluted with distilled water to 1.0 g/ml.

**Sample for chemical analysis of licorice decoction:** 1.0 ml of the decoction mentioned above was diluted to 25 ml with distilled water, filtrated through a 0.22 μm filteration and ready for the HPLC analysis (Liu et al., 2008).

**Sample for chemical analysis of active glycosides:** 1.0 mg of liquiritin and glycyrrhizin in powder form were mixed with 5.0 mg honey, and heated at 180 °C in drying oven for 1 h, respectively. The mixture was dissolved in 2 ml of 50% methanol solution and filtrated for the HPLC analysis.

**Assay for the influence on immune function of Pi-deficiency mice**

One hundred and ten mice, including 55 males and 55 females, were randomly divided into 11 groups. Mice in group 1 (normal group) were administrated intragastrically (i.g.) with distilled water. Mice in group 2-11 were administrated (i.g.) with rhubarb extract (10 g/kg body weight, twice a day) for 5 d to induce Pi-deficiency model (Zheng et al., 2006). Then, the mice in group 1 and group 2 (model group) received (i.g.) distilled water only. The remaining mice received (i.g.) crude licorice, processed licorice and honey once daily for 7 d, respectively. Twenty four hrs after the final administration, the mice were mainlined with the prepared India ink (0.01 ml/g body weight) in the cauda. Twenty microliter blood was taken from the eyebit vein plexus at 2 min and 20 min after the India ink injection, respectively. And then, the mice serum OD values at the wavelength of 680 nm were determined. The charcoal particle expurgation index (K) of macrophage count was done according to the following formula: \( K = (\lg \text{OD1} - \lg \text{OD2}) / (t_2 - t_1) \). Here \( t_1 \) and \( t_2 \) were the time points of taking blood from mouse eyes after injecting India ink (2 min and 20 min, respectively). Thirty minutes after the India ink injection, the thymus gland and spleen were taken out and weighed. The thymus gland index and spleen index was calculated according to the following formula: Organ Index (mg/g) = Organ Weight (mg)/bodyweight (g) (Chen, 1994).

**Antitussive test**

One hundred mice, including 50 males and 50 females, were divided into ten groups randomly. The mice in blank control group received (i.g.) distilled water, the remaining mice received (i.g.) crude licorice, processed licorice and honey once daily for 3 d, respectively. One hr after the final administration, the antitussive activity \textit{in vivo} was investigated on a

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classical mouse cough model induced by ammonia liquor. Briefly, mice were placed in a special glass chamber and exposed for 5 seconds to a 28% ammonia water aerosol which was produced through a nebulizer by compressed air at a pressure of about 400 mmHg. The cough incubation period and cough frequency produced during a 5 min exposure period were counted as reported by Xu et al. (2002).

Eliminating phlegm test

One hundred mice, including 50 males and 50 females, were divided into ten groups randomly. The mice in blank control group received (i.g.) distilled water, the remaining mice received (i.g.) crude licorice, processed licorice and honey once daily for 5 d, respectively. Half an hour after the final administration, the mice were administrated 0.5% red phenol (0.025 ml/g body weight) by intraperitoneal injection. Half an hour later, the trachea was separated and washed with 0.5 ml of 0.5% NaHCO₃ solution for two times. The rinsed solution was incorporated and centrifuged at 4000 × g for 30 min (4 ºC). The OD value of the supernatant was determined at the wavelength of 545 nm (Xu et al., 2002).

Detoxication test

Eighty mice, including 40 males and 40 females, were divided into 4 groups randomly. The blank control group of animals received (i.g.) distilled water, the remaining groups received (i.g.) crude licorice, processed licorice and honey, respectively. Five minutes later, they were administrated (i.g.) chloral hydrate (1200 mg/kg body weight). Mortality was recorded in 2 h (Sanders et al., 1982).

HPLC analysis

The chemical composition profiles of licorice slice, the contents of the 5 active compounds in licorice slice and the decocting quantity in water decoction were determined by HPLC analysis. The sample was separated on an Agilent Zorbax Elipse XDB-C₁₈ column (5 μm, 250 × 4.6 mm) connected with an Alltech Brava BDS-C₁₈ guard column (5 μm, 7.5 × 4.6 mm). Then, the contents of 5 active compounds in the licorice slice and their decocting quantity in water decoction were calculated using their standards.

Statistical analysis

The results were expressed as mean ± standard deviation (SD). Differences between groups were evaluated by one-way analysis of variance (ANOVA) followed by Dunnett’s test and χ² test with the aid of SPSS11.5 software package. A P value of less than 0.05 was considered statistically significant.

Results

Effect on immune function of Pi-deficiency mice

The results showed that crude and processed licorice obviously improved the immune function of Pi-deficiency mice, including elevating expurgation index, spleen index and thymus index. The crude licorice improved the expuration index was better than processed licorice, and the difference of expurgation index between crude and processed licorice group at the dosage of 5.0 g/kg was significant (P<0.05), indicating that honey-roasting could enhance the licorice immunomodulation action. Crude licorice and processed licorice both increased the spleen and thymus index of Pi-deficiency mice, but the significant difference between the same dosage was not observed (P>0.05). While in the honey-treated groups, only the high dosage group was effective in improving the expuration index of Pi-deficiency mice (Table 1).

Table 1: Effect of crude licorice, processed licorice and honey on the immune function of Pi-deficiency mice (mean ± SD, n=10)

<table>
<thead>
<tr>
<th>Group</th>
<th>Dose (g/kg)</th>
<th>Spleen index (mg/g)</th>
<th>Thymus index (mg/g)</th>
<th>Expurgation index</th>
</tr>
</thead>
<tbody>
<tr>
<td>model</td>
<td>/</td>
<td>3.24±0.54</td>
<td>2.29±0.32</td>
<td>5.31±2.28</td>
</tr>
<tr>
<td>normal</td>
<td>/</td>
<td>3.71±0.40*</td>
<td>2.47±0.54</td>
<td>7.59±2.40*</td>
</tr>
<tr>
<td>Crude licorice</td>
<td>1.25</td>
<td>3.01±0.23</td>
<td>2.13±0.49</td>
<td>6.04±1.75</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>3.84±0.67*</td>
<td>2.55±0.57</td>
<td>7.61±2.73</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>4.07±0.55**</td>
<td>2.68±0.37*</td>
<td>8.19±2.53*</td>
</tr>
<tr>
<td>Processed licorice</td>
<td>1.25</td>
<td>3.22±0.38</td>
<td>2.28±0.35</td>
<td>7.02±2.95</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>4.03±0.66**</td>
<td>2.48±0.46</td>
<td>8.73±3.02*</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>4.26±0.73**</td>
<td>2.77±0.41**</td>
<td>10.43±2.91**</td>
</tr>
<tr>
<td>Honey</td>
<td>1.25</td>
<td>3.42±0.44</td>
<td>2.30±0.32</td>
<td>5.25±1.62</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>3.37±0.39</td>
<td>2.44±0.46</td>
<td>6.64±1.95</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>3.54±0.45</td>
<td>2.39±0.21</td>
<td>7.56±1.20*</td>
</tr>
</tbody>
</table>

*P<0.05, ** P<0.01 vs model group; † P<0.05 vs crude licorice group of the same dosage.

Antitussive tests

The results in Table 2 showed that crude and processed licorice obviously prolonged the cough incubation period and inhibited the cough frequency caused by ammonia, indicating that they had the significant antitussive activity. The action

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of crude licorice was obviously stronger than that of the same dosage processed licorice (p<0.05), implying that honey-roasting obviously weakened the licorice antitussive activity. In the honey-treated groups, the significant activity of honey was not observed.

### Table 2: Effect of crude licorice, processed licorice and honey on mice cough induced by ammonia (mean ± SD, n=10)

<table>
<thead>
<tr>
<th>Group</th>
<th>Dose (g/kg)</th>
<th>Cough incubation period (s)</th>
<th>Cough frequency (time)</th>
</tr>
</thead>
<tbody>
<tr>
<td>water</td>
<td>/</td>
<td>35.9±11.8</td>
<td>99.4±12.7</td>
</tr>
<tr>
<td>Crude licorice</td>
<td>1.25</td>
<td>66.7±29.2 *</td>
<td>61.2±20.3 **</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>72.6±20.7 **</td>
<td>51.1±15.1 **</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>76.4±22.8 **</td>
<td>48.4±10.0 *</td>
</tr>
<tr>
<td>Processed licorice</td>
<td>1.25</td>
<td>42.7±8.6 *</td>
<td>79.7±28.5</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>47.2±11.7 *</td>
<td>68.3±20.4 **</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>55.8±21.3 *</td>
<td>59.7±16.7 **</td>
</tr>
<tr>
<td>Honey</td>
<td>1.25</td>
<td>36.9±15.4</td>
<td>94.3±19.5</td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td>43.0±7.7</td>
<td>83.5±21.6</td>
</tr>
<tr>
<td></td>
<td>5.00</td>
<td>40.9±13.8</td>
<td>85.4±13.9</td>
</tr>
</tbody>
</table>

*P<0.05, **P<0.01 vs vehicle group; *P<0.05, **P<0.01 vs crude licorice group of the same dosage.

### Table 3: Effect of crude licorice, processed licorice and honey on the death caused by chloral hydrate

<table>
<thead>
<tr>
<th>Group</th>
<th>Dose (g/kg)</th>
<th>Survival rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>water</td>
<td>/</td>
<td>4/20</td>
</tr>
<tr>
<td>Crude licorice</td>
<td>7.50</td>
<td>10/20*</td>
</tr>
<tr>
<td>Processed licorice</td>
<td>7.50</td>
<td>8/20</td>
</tr>
<tr>
<td>Honey</td>
<td>7.50</td>
<td>5/20</td>
</tr>
</tbody>
</table>

*P<0.05, **P<0.01 vs vehicle group.

**Eliminating phlegm**

The results showed that the crude and processed licorice significantly enhanced the red phenol secretion of mice trachea, indicating they had the obvious activity of eliminating phlegm. The action intensity of crude licorice was weakened after processing. Honey didn’t show obvious effect (Figure 2).

![Figure 2](http://dx.doi.org/10.4314/ajtcam.v9i2.2)
HPLC analysis

The HPLC profiles (showed in Figure 3) showed the chemical constituents between crude and processed licorice.

**Figure 3:** HPLC profiles of licorice (A. methanol extract of licorice; B. decoction of licorice; CL. crude licorice; PL. processed licorice; 1. liquiritin apioside; 2. liquiritin; 3. licuraside; 4. isoliquiritin; 5. glycyrrhizin). The mobile phase consisted of acetonitrile (a) and 0.1% phosphoric acid solution (v/v, b). The gradient and detection wavelength were as follows: 0-20 min, 13%-19% a, 276 nm; 20-37 min, 19%-35% a, 360 nm; 37-55 min, 35%-55% a, 254 nm; flow rate: 0.8 ml/min. The injection volume was 10 μL and the column was thermostated at 35°C.

**Figure 4:** Influence of processing on the content (A) and decocting quantity (B) of five active compounds in licorice (mean ±
were very different. The polar constituents of processed licorice and its decoction were more abundant than those of crude licorice. The ratio of peak areas, such as licurside (3)/isoliquiritin (4), obviously altered after processing. The result indicated that honey-roasting modified the chemical compositions in licorice. The contents of five compounds in licorice slice and the decocting quantity were obtained, according to the standard calibration curves established from six different concentrations of the standard solutions. The results showed that the contents and decocting quantity of liquiritin apioside, liquiritin and glycyrrhizin in processed licorice descended significantly, while the content and decoction of isoliquiritin obviously increased (Figure 4). In order to determine if honey-roasting had caused the decomposition of active glycosides, the standards were treated under the same processing condition. HPLC analysis results (Figure 5) showed that liquiritin and glycyrrhizin were obviously decomposed. Glycyrrhizin was decomposed to glycyrrhetic acid-3-O-β-D-glucuronide and glycyrrhetic acid. Liquiritin produced isoliquiritin and liquiritigenin. The result indicated that honey-roasting altered the chemical structure of active glycosides in licorice.

Discussion

Licorice is one of the most common medicines prescribed by TCM practitioners. TCM theory claims that its efficacy obviously changes after processing with honey. Since 200 BC, crude licorice and processed licorice have been used separately, and have been two of the most frequently employed botanicals in China (China Pharmacopoeia Committee, 2005). According to TCM theory, the crude licorice has the stronger potency of relieving cough, eliminating phlegm and detoxication. The complex prescriptions for cough and cold, such as Sangju drink, Yingqiao powder and Chaige Jieji decoction, often use crude licorice. Nowadays, the crude licorice has been used widely for bronchitis and asthma in Eastern and Western cultures. Our study showed that the honey-roasting weakened the crude licorice potency in relieving cough, eliminating phlegm and detoxication for the first time. The result was consisted with licorice clinical use.

It is reported that glycyrrhizin, liquiritin apioside and liquiritin have the significant activity of relieving cough and inhibiting allergic asthma (Kamei et al., 2005; Jayaprakasam et al., 2009; Xie et al., 2009; Shin et al., 2007). Their content decreases significantly after roasting with honey (Sung and Li, 2004; Wang et al., 1995; Wang et al., 2008). This study revealed that the activity of relieving cough and eliminating phlegm weakened obviously after honey processing; the contents and decocting quality of liquiritin apioside, liquiritin and glycyrrhizin descended obviously; honey did not show obvious activity. The activity decrease after honey processing was probably associated with the content decrease of flavonoid.
glycosides and glycyrrhizin.

In TCM theory, crude licorice is good at detoxication also. This study showed the crude product had the stronger activity of inhibiting death rate caused by chloral hydrate than that of processed product. It is reported that glycyrrhizin cloud relief acute intoxication caused by food and drug, and its content in crude licorice is higher than that in processed licorice (Li and Tong, 2007; He et al., 2009; Rahman and Sultana, 2007). Our study showed that the content and decocting quantity of glycyrrhizin in licorice descended after processing, and honey did not possess obvious intoxication action. So, the difference in detoxication activity between crude and processed licorice may be associated with the content of glycyrrhizin.

Processed licorice is believed to possess the better effect of invigorating vital energy than crude licorice, and often used to cure Pi-deficiency syndrome in TCM clinic. Some complex prescriptions for Pi-deficiency syndrome, such as Buzhong Yiqi decoction and Sijunzi decoction, all require using processed licorice rather than crude licorice. Although crude licorice and processed licorice had the obvious effect of enhancing Pi-deficiency mice’s immune function, the processed product showed the stronger activity than crude licorice in the present study. Glycyrrhizin and polysaccharides have been reported to possess significant activity of enhancing immune function (Yi et al., 1996; Cheng et al., 2008), but their content decrease after roasting with honey. It is to be noted that the content and decocting quantity of isoliquiritin increased obviously after processing in the present study. So, the immunomodulation activity of isoliquiritin is worth investigating.

The results of content analysis in the present study indicated that glycyrrhizin and flavone glycosides may be decomposed or isomerized in the processing condition. In order to test this hypothesis, the stability of glycyrrhizin and liquiritin, two major ingredients in licorice, was investigated under the same processing condition. The results indicated that the honey-roasting did cause the decomposition and isomerization of the active glycosides in licorice. The recently studies showed that roasting could caused glycyrrhizin thermal decomposition (Sung and Li, 2004; Hwang et al., 2006), consisted with the result of the present study. Previous research indicated that glycosidic flavonoid constituents were hydrolyzed stepwise during roasting (Kuwajima, 1999). Our study found that liquiritin isomerized to isoliquiritin and decomposed to liquiritigenin during honey-roasting for the first time. This conversion reaction was consisted with the content increase of isoliquiritin in processed licorice.

In conclusion, the honey-roasting obviously modified the main pharmacological activities and chemical compositions of licorice. This modification didn’t cause by honey only, and the roasting appeared to exert some effects. These findings provided a piece of scientific proof for the different therapeutic values of crude and processed licorice. In order to elucidate the mechanism of roasting with honey, the more researches, including the investigation of the processing influence on the lipid- soluble active ingredients, the absorption and metabolism of the active compounds in licorice, should be carried out.

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References


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