Quality difference study of six varieties of *Ganoderma lucidum* with different origins

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INTRODUCTION

*Ganoderma lucidum* Karst a medicinal fungus, belonging to Basidiomycetes, Aphiyllophorales, Ganodermataceae, is widely used in Oriental medicine to maintain health. With both edible and medicinal value, it has more than 2000 years of history in China. And the annual output of *G. lucidum* is over 10000 tons. *G. lucidum* has the function of anti-aging, enhancing immunity, radioprotective, and liver detoxification as well as inhibiting malignant tumor growth (Zhao et al., 1999; Lin, 2007; Lü et al., 2011). The chemical composition of *G. lucidum* is complex, which contains 11 categories of active substances, such as polysaccharides, triterpenoids, fats and oils, organic germanium, inorganic ions, and sterols. These ingredients are closely related to their pharmacological activity (el-Mekkawy et al., 2007). Polysaccharides and triterpenoids are considered to be its main medicinal components (Wang and Sun, 1990; Zhao et al., 2002; Lin, 2007). The quality of *G. lucidum* is evaluated though the content of polysaccharide in “Chinese Pharmacopoeia,” but Ganoderic acid in Japan (Zhang and Yang, 2006). Ganoderic acid belongs to triterpenoids, which has a wide range of pharmacological active components. It has become a hot study subject in *G. lucidum* (Chen and Yu, 1990; Yang et al., 1995; Zhou et al., 2004). Ganoderic acid A and B content account for more than half of *G. lucidum* (Ding et al., 2009), so the determination of ganoderic acid A and B content can be used as the scientific basis for judging quality of *G. lucidum*.

Because wild fungus resources are limited and artificial cultivation of *G. lucidum* is affected by origin, cultivation, harvesting conditions, and so on. These factors lead to different quality productions of *G. lucidum*. We tested the *G. lucidum* samples from some main producing areas in Shandong Liaocheng, Jiangsu Nantong, Fujian Wuyi Mountain Zhejiang Longquan, Jilin Changbai Mountain, and Anhui Dabie Mountain. The test is focused on the contents of polysaccharide, triterpenoid, and ganoderic acid A and B. The result is to provide the basis of procurement for using *G. lucidum* as main raw materials.

MATERIALS AND METHODS

MATERIALS

*Ganoderma lucidum* karst is used for this experiment which is respectively from the Shandong Liaocheng, Jiangsu Nantong, Fujian Wuyi Mountain, Zhejiang Longquan, Jilin Changbai Mountain, and Anhui Dabie Mountain. In addition to *G. lucidum* from Shandong Liaocheng cultured on cotton seed, others are wood cultured.

INSTRUMENTS AND REAGENT

Waters Acquity ultra performance liquid chromatography (UPLC), QK-250E type ultrasonic cleaner (Kunshan Ultrasonic Instrument Co., Ltd.), AB265-S/100000 electronic balance (Mettler Toledo), TU-1900 type ultraviolet-visible spectrophotometer (Beijing Purkinje General Instrument Co., Ltd.). Standard ganoderic acid A, ganoderic acid B (purity > 98%) provided by Shanghai Tong Tian Biotechnology Co., Ltd. Urolic acid reference substance provided by the national institute for the control of...
pharmaceutical and biological products. Acetonitrile (chromato-
graphic grade), double distilled water, phosphoric acid, chloro-
form, petroleum ether, ethyl acetate, methanol, and other reagents
are all analytically pure.

METHODS
Ganoderic acid A and ganoderic acid B content determination
Chromatographic conditions. The chromatographic column was
Waters X-BridgeC18 (4.6 mm × 150 mm, 3.5 μm); the detection
wavelength was set at 252 nm; gradient elution, liquid phase gra-
dient ratio, and time relationships as shown in Table 1; the column
temperature was kept at 40˚C, and the flow rate was 0.4 ml/min;
the injection volume was 10 μl. In these chromatographic condi-
tions, ganoderic acid A and ganoderic acid B Mixed reference
substance, and G. lucidum extraction UPLC spectrum diagram as
shown in Figure 1.

PREPARATION OF STANDARD SOLUTION AND CALIBRATION CURVES
Precisely weighing amount of each reference substance and then
put it into a 10-ml volumetric flask respectively, adding methanol
to dissolve and to the constant volume, reaching concentrations
of 1.802 mg/ml of ganoderic acid A and 1.020 mg/ml of ganoderic
acid B. Precisely weigh the liquid reserves each 8.0 ml in 25 ml
volumetric flask, add methanol to scale, and shake to make mixed
standard stock solution of ganoderic acid A and ganoderic acid B
whose concentration are 0.577 and 0.326 mg/ml. Precisely weigh
standard stock solution 2.0–10 ml volumetric flask, dilute to the
mark with methanol, and then shake to get mixed standard solu-
tion of ganoderic acid A and B whose mass concentration are 0.100
and 0.065 mg/ml.

According to the method of Zhao et al. (2009), precisely mea-
sure mixed control solution 0.5, 1, 2, 4, 6, 8.0 to 10 ml volumetric
flask, dilute to the mark with methanol, and shake to get a series of
standard solution. Respectively take a 10-μl sample of the mixed
standard solution to analyze under the chromatographic condi-
tions. Draw a standard curve and make regression calculation
with mass concentration of the reference as abscissa, peak area
as the ordinate, the results show that, the regression equation of
ganoderic acid A is $Y = 1.9E + 07X-122000$ and ganoderic acid
B is $R^2 = 0.9996; Y = 2.0E + 07X + 139583, R^2 = 0.9991$. Gan-
oderic acid A and ganoderic acid B respectively in 28.85–400.8,
16.30–260.8 μg/ml are in good linear relation.

Preparation of sample solution
According to the method of Liu (2008), the accurately weighed
powder sample (250 mg) was extracted with 100 ml chloroform by
the heating reflux for 1 h. The extract was filtered with filter paper
which washed by methanol. After evaporating chloroform to dry-
ness by a rotary evaporator, residue was dissolved in methanol in
a 5-ml flask, and then filtered through a 0.45-μm membrane. Ten
microliters of sample solution were injected into the UPLC system
for analyzing.
**Precision test**
According to the chromatographic conditions, taking ganoderic acid A and ganoderic acid B mixed reference solution to successive injection six time, and recording the peak area. The RSD of peak area of ganoderic acid A and ganoderic acid B was 1.6 and 2.4% respectively.

**Repetitive test**
Accurately weighed G. lucidum samples of six from Dabie Mountain, according to methods of above, the RSD of contents of ganoderic acid A and ganoderic acid B is 3.1 and 1.8% respectively. The result shows that this method has a good repetitiveness.

**Stability experiment**
Take sample solution from Anhui Dabie Mountain for the test, which is in 0, 2, 4, 6, 8, 10, 12, 24 h at room temperature and record peak area. The results show that the sample solution has good stability in 24 h and the RSD of peak area of ganoderic acid A and ganoderic acid B were 0.9 and 1.4%.

**Recovery rate test**
Take nine portions of G. lucidum (0.25 g) from the Wuyishan which the content of ganoderic acid A and ganoderic acid B is known. Divide the portions into three groups and add the control solution of low, middle, high concentrations of ganoderic acid A and ganoderic acid B to each portion, then calculate the recovery rate follow the method above. The result is as shown in Table 2.

**Determination of polysaccharide**

**Sample preparation.** Accurately weighing 2.0 g of power sample, extracted by Soxhlet extractor with 90 ml water in the round-bottom flask, and heated under reflux for 6 h, then transfer the extract to a 100 ml flask, add water to the scale. Precisely measured 10 ml extract, added ethanol 150 ml, placed for 12 h at 4˚C. The extract separated by centrifugal precipitation, the precipitate is dissolve in water in a 50-ml flask as the sample solution.

**Preparation of standard curve.** D-Glucose anhydrous (25 mg) is accurately weighed and then dissolved in 25 ml of double distilled water, 1 ml solution is drawn to dilute 100 times with double distilled water to produce corresponding stock standard solution (0.01 mg/ml). Accurately draw glucose control solution 0.2, 0.4, 0.6, 0.8, 1, 1.2 ml to the 10 ml test tube, add water to the volume of 2.0 ml, precisely add anthrone–sulfuric acid [1.0 g of anthrone was dissolved in sulfuric acid (80%) in a 100 ml flask] 6 ml, heated for 15 min, then remove and put in ice-water to cool for 15 min, with the corresponding reagent as control. Determine the absorbance in the 625 nm wavelength and make it as the ordinate, concentration as abscissa to establish a standard curve.

Precisely measure the sample solution 2 ml, put it into 10 ml test tube, Follow the method of establishing the standard curve, as the "precisely add anthrone–sulfuric acid 6 ml" begin to determine absorbance. Then calculate the content of the polysaccharide according to the standard curve.

**Triterpenoid determination**

**Preparation of standard curve.** Accurately weigh 1.15 mg of the ursolic acid, dissolve in 10 ml ethyl acetate to produce corresponding stock standard solution. Take 0, 0.10, 0.20, 0.40, 0.60, 0.80, 1, and 1.20 ml control solution to dryness in a water bath at 100˚C. Then add 0.40 ml 5% vanillin–acetic acid solution and 1 ml perchloric acid, at 60˚C water bath heating for 15 min then move it into ice-water bath, add 5 ml acetic acid, place it at room temperature for 15 min. Determine its absorbance in the 548.1 nm.

Draw standard curve based on the determination result. Standard weight in 0–0.14 mg range showed a good linear relationship with the absorbance value, the linear regression equation was 

\[ Y = 0.2158X - 0.0018, \text{ correlation coefficient } r = 0.9991. \]

**Extraction of triterpenoids.** Triterpenoid extracts were prepared by 95% alcohol extraction as described before (Hou and Liu, 2010). Accurately weigh 200 g of dry G. lucidum powder for extraction. Take G. lucidum extracts about 10 mg to dissolve in 10 ml ethyl acetate, and determine its absorbance following the method above.

**RESULTS**

**CONTENT DETERMINATION OF DIFFERENT ORIGIN OF GANODERIC ACID A AND GANODERIC ACID B**
The content of ganoderic acid A and ganoderic acid B is as shown in Table 3. The content of ganoderic acid A of Dabie Mountain is the highest (7.254 mg/g); the followed behind is Longquan (6.658 mg/g), Shandong (1.959 mg/g). Ganoderic acid B content for Longquan (4.574 mg/g) is the highest.

**CONTENT DETERMINATION OF POLYSACCHARIDE**
The content of polysaccharides has significant differences. The highest content of G. lucidum polysaccharides is in Shandong, followed by Wuyi Mountain (7.38%); the lowest (1.85%) is in Dabie Mountain (see Table 4).
CONTENT DETERMINATION OF TRITERPENOID
The content of triterpenoid from different origins has been shown in Table 5. The highest content of triterpenoid of *G. lucidum* is cultivated in Dabie mountain (5.38%), the lowest is in Longquan (2.07%). The difference between them is significant.

DISCUSSION
Triterpenoid and polysaccharide are as the basis of quality of *Ganoderma* product. Triterpenoid has significant effect in immune regulation and antitumor (Morigiwa et al., 1986; Ceng and Bao, 2004; Huang and Xiao, 2008); its content decides the antitumor effect of *G. lucidum* products. Polysaccharide has the function of improving immunity, antitumor effects, removing free radical, hypoglycemic, lipid-lowering (Xu and Xu, 2003), and other functions. In recent years, the study has attracted many researchers (Lin et al., 2002; Cao and Lin, 2004). *Ganoderic* acid B and lucidenic acid A have the inhibitory activity against HIV-1 protease (Min et al., 1998). For the reason of above, we got the main medical ingredients of *G. lucidum* from six origins.

Through this experimental data, both the triterpenoid and ganoderic acid B are the highest in *G. lucidum* from Dabie mountain. But the highest polysaccharide of artificial Cultivation *G. lucidum* is from Liaocheng. There are several reasons impacting the accumulation of polysaccharides and triterpenoids. First, the same species of *G. lucidum* from different origins due to the culture medium, the growth environment, different stages of growth, and covered soil or not, will have different polysaccharide content (Li et al., 1997; Ding et al., 1999; Wei et al., 2006; Chen et al., 2009; Ye et al., 2010). Second, the content of polysaccharide and triterpenoid is different in varieties of *G. lucidum* (Liu et al., 1999; Xing and Jiang, 2001; Xing et al., 2004; Zheng et al., 2007). Third, some research suggests that the different drying methods have some effect in the content of polysaccharide of *G. lucidum*. The own drying for *G. lucidum* is significantly higher than the direct drying in polysaccharide content, which is considered to be hydrolysis, induced by hydrolytic enzymes (Xing and Jiang, 2001). The content of polysaccharide is higher in asporogenous *G. lucidum* than in sporiparous *G. lucidum*. The different parts of *G. lucidum* have different content of polysaccharide (Shi et al., 2010). Maybe due to the superior cultivation environment in Dabie Mountain, the triterpenoid content is highest in *G. lucidum*. There maybe some relations between the high polysaccharide content and the culture medium in Liaocheng where the only *G. lucidum* were cultured on cotton seed.

CONCLUSION
This study gives a comprehensive assessment of the *G. Lucidum* in terms of its efficacy and material, it provides shallow datum for the *G. Lucidum* quality from different areas. The result of the experiment indicated that there was no distinction correlation between polysaccharide and triterpenoid contents. Because of the difference in active ingredient from different origins, we can choose the *G. lucidum* according to our purposes.

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