Accepted Manuscript

Title: Tissue-smashing based ultra-rapid extraction of chemical constituents in herbal medicines

Author: Yong Fan Chen-Pu Yan Cheng Chen Kwok-Fai So

Ping Li Lian-Wen Qi

PII: S0731-7085(14)00123-X

DOI: http://dx.doi.org/doi:10.1016/j.jpba.2014.03.004

Reference: PBA 9483

To appear in: Journal of Pharmaceutical and Biomedical Analysis

Received date: 2-1-2014 Revised date: 28-2-2014 Accepted date: 3-3-2014

Please cite this article as: Yong FanChen-Pu YanCheng ChenKwok-Fai SoPing LiLian-Wen Qi Tissue-smashing based ultra-rapid extraction of chemical constituents in herbal medicines (2014), http://dx.doi.org/10.1016/j.jpba.2014.03.004

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.



1

For Journal of Pharmaceutical and Biomedical Analysis

2	Tissue-smashing based ultra-rapid extraction of chemical constituents in herbal
3	medicines
4	
5	Yong Fan ^{a,1} , Chen-Pu Yan ^{a,1} , Cheng Chen ^a , Kwok-Fai So ^b , Ping Li ^{a,**} and Lian-Wen Qi ^{a,*}
6	
7	^a State Key Laboratory of Natural Medicines and Department of Pharmacognosy, China
8	Pharmaceutical University, No. 24 Tongjia Lane, Nanjing 210009, China.
9	^b Department of Anatomy, The University of Hong Kong, Hong Kong Special Administrative
10	Region.
11	
12	
13	
14	
15	* Corresponding author. Tel: +86 25 86185135, Fax: +86 25 83271379.
16	E-mail address: Qilw@cpu.edu.cn (L. W. Qi) *
17	¹ The first two authors (Yong Fan and Chen-Pu Yan) contributed equally to this work.

Abstract

18

19	Sample extraction is the first challenge in analysis of herbal medicines (HMs). Numerous methods
20	have been developed to improve extraction efficiency, use less solvent and short time. In this work,
21	a tissue-smashing based ultra-rapid extraction (TSURE) method has been proposed through the
22	designed particle crushing, drastic stir, and dynamic molecular permeation at normal temperature.
23	Factors in TSURE like extraction time, volts, and solvents were optimized for extraction efficiency
24	of salvianolic acid B, cryptotanshinone, and tanshinone IIA from Salvia miltiorrhiza. The TSURE
25	method was validated in terms of repeatability (RSD < 2.2%) and extraction recoveries (93-106%)
26	with RSD $< 5.0\%$). TSURE showed a comparable extraction efficiency to conventional heat reflux
27	extraction (HRE) and better than ultrasonic assisted extraction (UAE). The extraction time was
28	about 2.0-3.0 min for TSURE, 60 times faster than the performance of HRE and 20 times faster
29	than UAE. Microscopic analysis showed that the Krummbein diameter of plant particles after
30	extraction were about 600-1200 μm for HRE and UAE, and decreased to 50-80 μm for TSURE.
31	Subsequently, the developed TSURE was applied to high-throughput extraction of 19 S.
32	miltiorrhiza samples collected in different regions of China. Besides, application of TSURE to
33	other herbal medicines was also investigated, including Panax quinquefolius and Lonicera japonica
34	TSURE method provided an ultra-rapid and promising alternation for extraction of ingredients in
35	herbal medicines, and can be extended to pharmaceutics, foods and cosmetics.

38

36

37

medicines; quality control; ultra-rapid.

Keywords: Tissue-smashing based ultra-rapid extraction; High-throughput extraction; herbal

1. Introduction

39	Sample preparation is the first challenging step in analysis and quality control of botanicals and
10	herbal medicines (HMs) [1]. Efficient sample extraction strategy can improve extraction efficiency
4 1	and enrich the target analytes [2-3]. As stated in the previous studies, some conventional and
12	simple methods, such as ultrasonic assisted extraction (UAE), heating under reflux extraction
13	(HRE), are commonly used [4-5]. Methanol and ethanol are most widely referred as the solvents
14	[6, 7]. These methods, however, are usually time-consuming, solvent-consuming, and may have
15	low extraction efficiencies [8, 9].
16	In recent years, many ultra-pressure or ultra-temperature extraction methods have been

- introduced for extraction of analytes of interest present in plant materials, such as pressurized liquid extraction (PLE), microwave-assisted extraction (MAE), and supercritical fluid extraction (SFE) [10-12]. The newer methods use relatively less solvent, take shorter time and are more efficient [5]. Various sample preparation techniques are summarized and compared for the extraction of plant materials [13]. Each has advantages and limitations depending on the projected use of results and the properties of analytes [14]. Undoubtedly, methods that are simple, rapid and environmentally friendly will be preferred [15].
- Tissue-smashing based ultra-rapid extraction (TSURE) was first introduced as a new extraction technique in 1993 [16]. The operating process of TSURE method is similar to juice squeezing [17].

 The TSURE enables ultra-rapid extraction of target ingredients at normal temperature through the designed particle crushing, drastic stir, and dynamic molecular permeation [18]. An ultra-rapid extraction process provided by TSURE is meaningful for sample analysis, such as the qualification

and quantification of ingredients from herbal medicines. In TSURE process, plant particles were crushed into smaller ones with the help of high-speed shear force and mixing power. In addition, under the partial negative pressure permeation, soluble balance between solid materials and solvents can be rapidly achieved [19]. The main advantages of TSURE are its versatility, ultra high speed, flexibility and low cost [20]. Summary diagram of the TSURE method was shown in Figure 1.

In this work, a TSURE method was developed and its potential in rapid extraction of constituents in HMs was systematically investigated. Factors in TSURE like extraction time, volts, and solvents were optimized for extraction efficiency of salvianolic acid B, cryptotanshinone, and tanshinone IIA from *Salvia miltiorrhiza*, one of the best-selling and most studied natural products [21]. The TSURE method was validated in terms of repeatability and extraction recoveries. TSURE was compared with two conventional methods HRE and UAE in extraction efficiencies and extraction time. Microscopic analysis was performed to test the plant particle sizes after extraction. Subsequently, the developed TSURE was applied to high-throughput extraction of 19 *S. miltiorrhiza* samples collected in different regions of China. Besides, the TSURE was also applied to the other two botanical materials, extraction of ginsenosides Rb1, Rc, Rg1 and Re from *Panax quinquefolius*, and extraction of chlorogenic acid, 3, 5-dicaffeoylquinic acid and 4, 5-dicaffeoylquinic acid from *Lonicera japonica*. This work demonstrates the potential of TSURE method for extraction of compounds of interest in herbal medicines and opens perspectives for similar studies on pharmaceutical, cosmetic and food industries.

2. Experimental

2.1. Plant materials

- S. miltiorrhiza samples were collected from 19 different regions of China. Radix samples of
 American ginseng (P. quinquefolius) was purchased from Roland Ginseng, LLC (Wausau, WI,
 USA), and Flos samples of L. Japonica was obtained from Shandong Province, China. The
 botanical origins of the materials were identified by the authors. The sample specimens were
 deposited at room temperature in the stationary storage center with accession numbers named
 2013S-1 to 2013S-19, 2013P-1 and 2013L-1 in State Key Laboratory of Nature Medicines, China
 Pharmaceutical University.
- 88 2.2. Chemicals and reagents
- Acetonitrile was of HPLC grade from Merck (Darmstadt, Germany). Deionized water was further purified by a Milli-Q system (Millipore, Milford, MA, USA). Other chemicals were of analytical grade. All solvents and samples were filtered through 0.22 µm membranes before injecting into HPLC.
- Reference compounds, including salvianolic acid B, cryptotanshinone, tanshinone IIA, chlorogenic acid, 3, 5-dicaffeoylquinic acid and 4, 5-dicaffeoylquinic acid were bought from Must Bio-Tech Co. Ltd. (Chengdu, China). The reference ginsenosides Rg1, Re, Rb1 and Rc were purchased from Jilin University (Changchun, China). Their structures shown in Supplementary Figure S1 were further elucidated in the authors' laboratory by ¹³C NMR and MS data. The purity of each reference compound was determined to be higher than 95% by normalization of the peak areas detected by HPLC-UV.

100 *2.3. Apparatus*

- TSURE experiments were performed on a JHBE-50S Herbal Blitzkrieg Extractor (Henan Jinnai Sci-Tech Development Ltd.). The extractor contains five major parts, including integrated volt controller, lifting controller, high speed motor, tissue crushing head and extraction bottle.
- Chromatographic analyses were carried out on a Shimadzu HPLC system consisting of a pump (LC-20AB), an auto-sampler (SIL-20A), UV/VIS detector (SPD-20A) and automatic column temperature control oven (CTO-20AC). Separation was performed on an Amethyst C18-P column (5 μm, 4.6×250 mm). Shimadzu Labsolutions software were used for the chromatographic analysis.
- Microscopic test was performed by a Nikon Eclipse 50i microscope system and analyzed by
 NIS-elements F 3.0 version software.

111 *2.4. Analysis*

For HPLC analysis of S. miltiorrhiza sample, the mobile phase consisted of 0.1% formic acid 112 water (A) and acetonitrile (B) using a gradient elution of 27-30% B at 0-8 min, 30-70% B at 8-15 113 114 min, 70-85% B at 15-30 min and 85-100% B at 30-40 min. The detection wavelength was set at 115 286 nm for salvianolic acid B and 270 nm for cryptotanshinone and tanshinone IIA. The 116 chromatographic conditions for American ginseng were using 0.025% phosphoric acid water (A) and acetonitrile (B) with a gradient elution of 19-20% B at 0-25 min, 20-40% B at 25-60 min and 117 40-100% B at 60-70 min. The wavelength was set at 203 nm for ginsenosides analysis. The 118 chromatographic conditions for L. Japonica were using 0.1% formic acid water (A) and acetonitrile 119

- (B) as the mobile phase in a linear gradient program of 10-20% B at 0-15 min, then 20% B isocratic elution for 15 min, 20-30% B at 30-40 min and 30-100% B at 40-50 min. The detector wavelength was set at 350 nm. All the sample volume injected was 10 μ l and the flow rate was 1 ml/min with column temperature at 35 °C.
 - 2.5. Tissue-smashing based extraction

- All the dried samples of *S. miltiorrhiza*, *P. quinquefolius* and *L. japonica* were pulverized into powder through a 40 mesh sieve. Sample powder and solvent were mixed in an extraction bottle.

 The tissue crushing head was lifted under solvent surface. The designed extraction volt can be adjusted by twisting the integrated volt controller. Accurately weighed 0.5 g powder was extracted by TSURE method using solvents at different ratios to form a homogeneous solution. The loss of the solvent was supplemented. The sample solutions were then centrifuged at 13,000 rpm for 10 min and then filtrated through 0.22 μm filters before injecting into HPLC analysis.
- 132 2.6. Reference extraction procedures
- 133 *UAE:* Accurately weighed 0.5 g powder of *S. miltiorrhiza*, *P. quinquefolius* and *L. japonica*.

 134 The powders were then mixed with 40 ml of methanol and placed into a 150 ml conical flask.

 135 UAE was conducted on a KH-300DB digital ultrasonic cleaner (Kunshan Ultrasonic Instrument

 136 Co., Ltd. Kunshan, Jiangsu, China) with frequency set at 100 Hz for 40 min.
- 137 HRE: 0.5 g herbal powder of S. miltiorrhiza, P. quinquefolius and L. japonica were weighed
 138 accurately to a 100 ml round-bottom flask with 40 ml of methanol, and then the system was
 139 extracted for 120 min at 75 °C.

140	All of the obtained extracts were cooled to room temperature and the loss of the solvent was
141	replenished with methanol. After centrifugation at 13,000 rpm for 10 min, the supernatant was
142	filtered through 0.22 µm filters and then injected in to HPLC system for analysis.

3. Results and discussion

3.1. Optimization of the TSURE process

Sequential investigations of a number of main variables potentially affecting the TSURE procedure were conducted to obtain an efficient extraction. The univariate method was used to optimize the five major parameters including time, extraction volt, solid-liquid ratio, extraction solvent and number of extraction cycles. The *S. miltiorrhiza* sample from Shandong-4 was used to test extraction efficiencies. The results were graphically summarized in Figure 2.

3.1.1. Optimization of the extraction solvents

The extraction solvents in this study tested were methanol and ethanol. As shown in Figure 2, methanol offered better extraction efficiencies than ethanol, especially for water-soluble salvianolic acid B, probably because of the stronger cell wall breaking effect of methanol [22]. Addition of water as the solvents causes possible blistering under intensive stir. Thus water was not considered in this work.

3.1.2. Optimization of the extraction time

Extraction time plays a vital role in the TSURE method. To identify the optimal time, we evaluated TSURE method in a range from 0.5 to 5 min at normal temperature. Results indicated

- that extraction time more than 2 min did not show a significant increase in the extraction efficiency for salvianolic acid B, cryptotanshinone, and tanshinone IIA from *S. miltiorrhiza*. When the extraction time increased to 5 min, the extraction efficiency of salvianolic acid B decreased probably because of thermal degradation of the compound at an extraction temperature higher than 85 °C [23]. Therefore, the optimal extraction time in this study was chosen to be 2 min.
- 164 3.1.3. Optimization of the extraction volt

165

166

167

168

169

170

- TSURE method uses a high speed motor to deliver energy to the tissue crushing head in contact with the samples and solvents. Larger the extraction volt is, higher speed the head achieves. Extraction volt was evaluated in a range from 90-110 V with 2-min extraction time. The extraction efficiencies at 100 V were better than those at 90 V and 110 V for all the three target compounds. Similarly, the extraction volt higher than 100 V resulted in an obvious decrease of the yield of salvianolic acid B because of its thermal instability in the case of increased temperature to 90 °C induced by high-speed stir, indicating the difficulty to control temperature at high volt.
- 3.1.4. Optimization of the solid-liquid ratio
- Another major step in the development of the TSURE method was to select a suitable solid-liquid ratio. In this study, ratios varying from 1:40 to 1:120 were investigated. As shown in Figure 2, a solid-liquid ratio at 1:80 was observed to achieve acceptable efficiency for the TSURE procedure and no significant improvement was indicated with a higher ratio at 1:100 and 1:120.
- 3.1.5. Optimization of the extraction cycles

Selection of a suitable extraction cycle is critical in this study. Data are shown in Figure 2. Extraction cycle of the TSURE method is usually conducted once. With a purpose to increase the extraction yield of the three analytes, the TSURE method was performed for a total of 1, 2 or 3 cycles. As seen in Figure 2, no significant difference was observed with multiple extraction cycles than once. To save the time and solvents, the optimal extraction cycle was once.

On the basis of these results, the optimized conditions for TSURE method were established using methanol as the solvent, a solid/liquid ratio at 1:80, an extraction volt at 100 V, and a single extraction for 2 min.

3.2. Comparison of the TSURE approach with the reference extraction procedure

The performance of TSURE was compared with HRE and UAE in terms of extraction efficiency and extraction time. They were conducted at the optimal conditions according to the experimental section from the 2010th Chinese Pharmacopoeia [24]. The concentrations of the three target compounds extracted by the three methods were compared in Table 1. As shown in Table 1, in reference to the students' t test, the total yield obtained between the TSURE and HRE showed no significant difference with a p-value 0.12. The efficiency of the UAE was significantly lower than that of TSURE method with a p-value < 0.05. Similar conclusion was also observed from the results obtained in other two herbal materials, P. quinquefolius, and L. japonica. This is probably because the TSURE method combines the process of smashing, vibration and stir together to extract the bioactive ingredients in herbal materials. However, in UAE procedure, the smashing and stirring process were not contained. At the late stage of UAE, there was equilibrium of extraction for the solvent that was unchanged. The UAE is thus usually considered as a moderate extraction

technique.

The advantages of TSURE method over HRE are consumption of less time, low energy required and normal temperature operation. First, the extraction time was shortened from 120 min with HRE to 2 min with TSURE method, providing an ultra-rapid alternation for qualification and quantification of *S. miltiorrhiza*. Second, TSURE was operated at normal temperature. Though stir process will generate some heat, the temperature remains under 50 °C. For compounds like volatile oil or semivolatile organics, TSURE method may be a good choice. We demonstrate here that the extraction of salvianolic acid B, cryptotanshinone and tanshinone IIA in *S. miltiorrhiza* using TSURE occupied higher or equal extraction efficiency as compared to the conventional methods with a high-speed extraction process. Supplementary Figure S2 showed no significant difference of the chemical components in *S. miltiorrhiza* after TSURE, UAE and HRE, indicating the stability of ingredients during TSURE process.

3.3. Method validation

All compounds determined by HPLC were identified by comparison of retention times with those of the reference compounds. In addition, spiking the samples with the standard compounds further confirmed the identities of these peaks. Typical chromatograms obtained from *S. miltiorrhiza*, *P. quinquefolius*, and *L. japonica* extracted by TSURE were shown in Figure 3. To validate the novel TSURE-HPLC method, the linearity, repeatability, recovery and detection limits were investigated under the optimal condition.

Calibration curves for salvianolic acid B, cryptotanshinone and tanshinone IIA were

constructed using the areas of the chromatographic peaks determined at six increasing concentrations. As shown in Table 2, a good linearity was observed with $r^2 > 0.999$ within test ranges for the three analytes. Limits of detection and quantification (LODs and LOQs) were calculated at 6 times based on the signal-to-noise ratio of 3 and 10. The LODs and LOQs were 0.32 ng and 1.60 ng for salvianolic acid B, 0.03 ng and 0.06 ng for cryptotanshinone, 0.02 ng and 0.10 ng for tanshinone IIA. As consistent with publications, tanshinones are better than salvianolic acids in the sensitivity detected by UV [25]. The repeatability was estimated by five repetitive samples obtained by TSURE. As Table 3 shows, the RSDs of the analytes were in the range of 1.6-2.2%, indicating that the repeatability of the method is acceptable.

Recovery test of the optimized TSURE procedure was examined by analyzing spiked *S. miltiorrhiza* samples with certificated low, medium and high concentrations for salvianolic acid B, cryptotanshinone and tanshinone IIA. As presented in Table 3, excellent recoveries in a range of 93-106% were observed with a relative standard deviation (RSD) of 0.30-4.81% for the Shandong-4 samples, suggesting that the proposed TSURE method is reliable.

233 3.4. Application of the TSURE method to 19 S. miltiorrhiza samples and other herbs

Figure 4 summarized the results of extraction yields of salvianolic acid B, cryptotanshinone and tanshinone IIA using TSURE in 19 *S. miltiorrhiza* samples cultivated in different regions of China. To test the universality of TSURE, the present method was applied to extraction of two different botanical materials, roots of *P. quinquefolius* and flowers of *L. japonica*. As the data shows, the concentrations of the constituents varied remarkably among the 19 *Salvia* samples. The concentration ranged from 6.1 mg/g to 47.1 mg/g for salvianolic acid B, 0.1 mg/g to 2.0 mg/g for

- cryptotanshinone, and 0.7 mg/g to 2.6 mg/g for tanshinone IIA. Quality control is of great importance since the chemical content of herbal materials differs, depending on the geographic information of the locations and even the season of its collection, and the subsequent processing.

 As shown in Figure 4, 4 *Salvia* samples collected from Gansu Province possess the lowest yield of target compounds comparing with other regions. This may because the dry climate in Gansu is unfavorable for the compounds enrichment in *S. miltiorrhiza*.
 - For *P. quinquefolius*, we focused on four major bioactive ginsenosides, Rg1, Re, Rb1 and Rc. As shown in Supplementary Figure S 3(A), TSURE method showed no significant differences in extraction efficiency compared with UAE (p= 0.78), but a little lower than HRE in the extraction efficiency of ginsenoside Rb1 (p= 0.002). The possible reason is that in HRE, the use of heat may cause an increase in the solubility of materials and rate of mass transfer. Moreover, ginsenosides are relatively not easily dissolved in organic solvents such as methanol. As shown in Supplementary Figure S 3(B), for *L. japonica*, no significant differences were observed in extraction efficiency for chlorogenic acid, 3,5- dicaffeoylquinic acid, and 4,5-dicaffeoylquinic acid between TSURE and UAE (p= 0.46). HRE showed higher extraction efficiency than TSURE for chlorogenic acid (p= 0.0003).
- 256 3.5. Mechanism of TSURE

- 257 3.5.1. Tissue-smashing effect during TSURE
- To further investigate the mechanism of TSURE, microstructure analysis was used. Figure 5 indicated that the effects of different extraction methods on herbal particle sizes of *S. miltiorrhiza*.

Figure 5(A), 5(B) and 5(C) showed the microscopic graphs of samples after HRE, UAE, and TESRE respectively. In order to get a microscopic image without bubbles, the particles were soaked in cold water overnight. The Krummbein diameter of particles photographed in wet status after HRE, UAE and TSURE were offered as 600-1200 μm for HRE and UAE and 50-80 μm for TSURE, respectively. In this method, the rotating speed of tissue crushing head can achieve 15000-30000 r/min. With the ultra-high speed, sample tissues are crushed into smaller size in a moment.

3.5.2. Drastic stir

During the dissection between high-speed inner and stable outer edges, a strong vortex in the center of inner edge was developed to drive the stir of crushed samples, inducing a rapid concentration change to the whole system. With smaller particles, rapid exposure and transfer of extracted sample molecules to solvent environment will occur. In TSURE, an alternation of balance and unbalance between solvent and material particles was proceeded to achieve the final thorough smashing and fully balanced extraction.

3.5.3. Ultra-rapid dynamic molecular permeation

In working status, the whole instrument was an ultra-dynamic system. Due to the high-speed rolling between the inner and stable outer edges, a vortex negative pressure was developed and the dissection of samples occurred. Under this negative pressure, molecular permeation was observed between the inner and outer side of the outer edge, meaning that the smashed or extracted molecules were surrounded, dissociated and replaced by solvents. Finally, the molecules would be

separated from herbal materials to finish the extraction process.

3.5.4. Strong vibration effect

It is believed that ultrasonic wave can accelerate maceration process [26]. The destructive effect of vibration on the plant tissue and dispersion system had been well studied [27]. This instrument can produce a vibration equivalent to 1/60 of the ultrasonic wave with a high speed rotation [16]. Undoubtedly, the solubility equilibrium between the inner and outer smashed sample particles can be strongly accelerated with the vibration effect.

4. Conclusions

In this work, a simple tissue-smashing based ultra-rapid extraction method proved to be efficient and validated to extract chemical constituents from herbal materials. Comparing with the conventional methods, TSURE provides excellent acceleration of the extraction process and higher or equal extraction yields with the optimized conditions. The big noise generated by the instrument is a major drawback that should be overcome in the future research. In addition, this promising extraction method offers an alternative reference for the application of scale-up production in pharmaceutical, food and cosmetic industries.

Acknowledgements

Special thanks go to Dr Alolga Raphael N. for his correction of the grammar errors in this manuscript. This work was supported in part by the National Natural Science Foundation of China (Nos. 81173497, 81222052, and 81130068), and Program for New Century Excellent Talents in

University (NCET-11-0737).



300 References

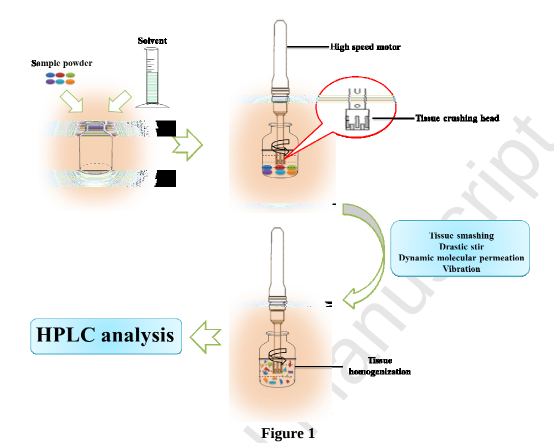
- 301 [1] L.-W. Qi, C.-Z. Wang, C.-S. Yuan, Isolation and analysis of ginseng: advances and challenges,
- 302 Nat. Prod. Rep. 28 (2011) 467-495.
- [2] R.-P.-S. Dassanayake, H. Wei, R.-C. Chen, A. Li, Optimization of the matrix solid phase
- dispersion extraction procedure for the analysis of polybrominated diphenyl ethers in human
- 305 placenta, Anal. Chem. 81 (2009) 5816-5822.
- 306 [3] X.-L. Cheng, J.-Y. Wan, P. Li, L.-W. Qi, Ultrasonic/microwave assisted extraction and
- diagnostic ion filtering strategy by liquid chromatography-quadrupole time-of-flight mass
- spectrometry for rapid characterization of flavonoids in *Spatholobus suberectus*, J. Chromatogr.
- 309 A 1218 (2011) 5774-5786.
- 310 [4] J.-L. Zhang, M. Cui, Y. He, H.-L. Yu, D.-A. Guo, Chemical fingerprint and metabolic
- fingerprint analysis of Danshen injection by HPLC-UV and HPLC-MS methods , J.
- 312 Pharmaceut. Biomed. Anal. 36 (2005) 1029.
- [5] Y.-J. Wei, L.-W. Qi, P. Li, H.-W. Luo, L. Yi, L.-H. Sheng, Improved quality control method
- for Fufang Danshen preparations through simultaneous determination of phenolic acids,
- saponins and diterpenoid quinones by HPLC coupled with diode array and evaporative light
- scattering detectors, J. Pharmaceut. Biomed. Anal. 45 (2007) 775.
- [6] S. Babić, M. Petrović, M. Kaštelan-Macan, Ultrasonic solvent extraction of pesticides from soil,
- J. Chromatogr. A 823 (1998) 3-9.
- [7] R. Metivier, F. Francis, F. Clydesdale, Solvent extraction of anthocyanins from wine pomace, J.
- Food Sci. 45 (1980) 1099-1100.
- [8] W. Huang, A. Xue, H. Niu, Z. Jia, J. Wang, Optimised ultrasonic-assisted extraction of

- flavonoids from *Folium eucommiae* and evaluation of antioxidant activity in multi-test systems *in vitro*, Food Chem. 114 (2009) 1147-1154.
- [9] X. Pan, G. Niu, H. Liu, Comparison of microwave-assisted extraction and conventional
- extraction techniques for the extraction of tanshinones from Salvia miltiorrhiza bunge,
- 326 Biochem. Eng. J. 12 (2002) 71-77.
- [10] S.-B. Hawthorne, C.-B. Grabanski, E. Martin, D.-J. Miller, Comparisons of Soxhlet extraction,
- pressurized liquid extraction, supercritical fluid extraction and subcritical water extraction for
- environmental solids: recovery, selectivity and effects on sample, J. Chromatogr. A 892 (2000)
- 330 421-433.
- [11] V. Lopez-Avila, R. Young, W.-F. Beckert, Microwave-assisted extraction of organic
- compounds from standard reference soils and sediments, Anal. Chem. 66 (1994) 1097-1106.
- [12] J.-J. Langenfeld, S.-B. Hawthorne, D.-J. Miller, J. Pawliszyn, Effects of temperature and
- pressure on supercritical fluid extraction efficiencies of polycyclic aromatic hydrocarbons and
- polychlorinated biphenyls, Anal. Chem. 65 (1993) 338-344.
- [13] C.-W. Huie, A review of modern sample-preparation techniques for the extraction and
- analysis of medicinal plants, Anal. Bioanal. Chem. 373 (2002) 23-30.
- 338 [14] Y. Dai, G.-J. Witkamp, R. Verpoorte, Y.H. Choi, Natural Deep Eutectic Solvents as a New
- Extraction Media for Phenolic Metabolites in Carthamus tinctorius L, Anal. Chem. 85(2013)
- 340 6272-6278.
- [15] F. Chemat, M. Lucchesi, J. Smadja, L. Favretto, G. Colnaghi, F. Visinoni, Microwave
- accelerated steam distillation of essential oil from lavender: A rapid, clean and
- environmentally friendly approach, Anal. Chim. Acta 555 (2006) 157-160.

- 344 [16] Y.-Z. Liu, K. Yuan, C.-R. Ji, New method of extraction on the chemical components of
- Chinese medicinal plants-extracting method by smashing of plant tissue, Henan Science 11
- 346 (1993) 265-268.
- [17] W. Tang, Y. Sun, Y. Liu, Y. Wei, Y. Zhao, Fast extraction of bioactive fatty acids from the
- perilla seeds by smash tissue extraction, Pakis. J. Bot. 45 (2013) 513-520.
- 349 [18] L.-F. Zhang, Z.-L. Liu, Optimization and comparison of ultrasound/microwave assisted
- extraction (UMAE) and ultrasonic assisted extraction (UAE) of lycopene from tomatoes,
- 351 Ultrason. Sonochem. 15 (2008) 731-737.
- [19] Y. Chen, X. Gu, S.-Q. Huang, J. Li, X. Wang, J. Tang, Optimization of ultrasonic/microwave
- assisted extraction (UMAE) of polysaccharides from *Inonotus obliquus* and evaluation of its
- anti-tumor activities, Int. J. Biol. Macromol. 46 (2010) 429-435.
- [20] S.-M. Shi, Y.-Z. Liu, W. Tai, C.-Q. Chen, Y.-Q. Zhao, Smashing Tissue Extraction and HPLC
- Determination of Active Saponins from Different Parts of *Panax notoginseng*, Chinese Herbal
- 357 Med. 4 (2012) 340-344.
- 358 [21] P. Tian, Convergence: Where west meets east, Nature 480 (2011) S84-S86.
- 359 [22] X. Ren, T.-E. Springer, T.-A. Zawodzinski, S. Gottesfeld, Methanol transport through nation
- membranes. Electro-osmotic drag effects on potential step measurements, Electrochem. Soc.
- 361 147 (2000) 466-474.
- 362 [23] X.-T. Zheng, H.-B. Qu, Characterisation of the Degradation of Salvianolic Acid B Using an
- On line Spectroscopic Analysis System and Multivariate Curve Resolution, Phytochemical
- Analysis 23 (2012) 103-109. [24] Chemical Industry Press, Beijing, The State Pharmacopoeia
- Commission of PR China, in: Pharmacopoeia of the People's Republic of China, vol. I, 2010,

366	122 and 205.
367	[25] J. Cao, YJ. Wei, LW. Qi, P. Li, ZM. Qian, HW. Luo, J. Chen, J. Zhao, Determination of
368	fifteen bioactive components in Radix et Rhizoma Salviae Miltiorrhizae by high-performance
369	liquid chromatography with ultraviolet and mass spectrometric detection, Biomed.
370	Chromatogr. 22 (2008) 164-172.
371	[26] W. Huang, A. Xue, H. Niu, Z. Jia, J. Wang, Optimised ultrasonic-assisted extraction of
372	flavonoids from Folium eucommiae and evaluation of antioxidant activity in multi-test
373	systems in vitro, Food Chem. 114 (2009) 1147-1154.
374	[27] M. Toma, M. Vinatoru, L. Paniwnyk, T. Mason, Investigation of the effects of ultrasound on
375	vegetal tissues during solvent extraction, Ultrason. Sonochem. 8 (2001) 137-142.
376	

376	Figure Legends
377	Figure 1. Summary diagram of the TSURE method. Accurately weighted samples were added into
378	an extraction bottle and mixed with solvent. Then the tissue crushing head was lifted under solven
379	surface. After several minutes, a homogeneous mixture contains analytes of interest was obtained.
380	Figure 2. Extraction efficiencies of three target analytes in S. miltiorrhiza under differen
381	conditions. The condition with an asterisk was chosen for the optimized TSURE procedure. The
382	error bars represent standard deviation of the triple analyses.
383	Figure 3. Representative HPLC chromatogram of <i>S. miltiorrhiza</i> (A), <i>P. quinquefolius</i> (B), and <i>L</i>
384	japonica (C) extracted using TSURE (peak 1, salvianolic acid B; peak 2, cryptotanshinone; peak
385	3, tanshinone IIA; peak 4, ginsenoside Rg1; peak 5, ginsenoside Re; peak 6, ginsenoside Rb1
386	peak 7, ginsenoside Rc; peak 8, chlorogenic acid; peak 9, 3,5-dicaffeoylquinic acid; peak 10
387	4,5-dicaffeoylquinic acid).
388	Figure 4. Extraction efficiencies of three target analytes in 19 individual S. miltiorrhiza samples
389	collected from different regions of China by TSURE method. The error bars represent standard
390	deviation of triple analyses.
391	Figure 5. Micrographs of S. miltiorrhiza particles after different extraction methods: (A) HRE, (B)
392	UAE, (C) TSURE. The images were obtained using a Nikon Eclipse 50i microscope system.



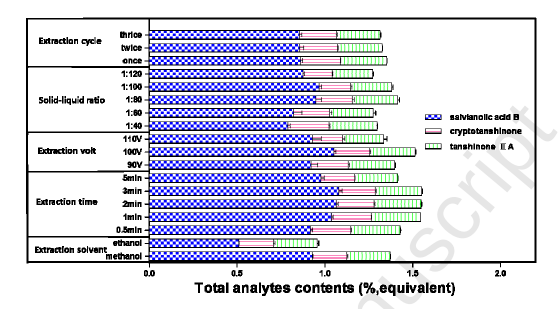
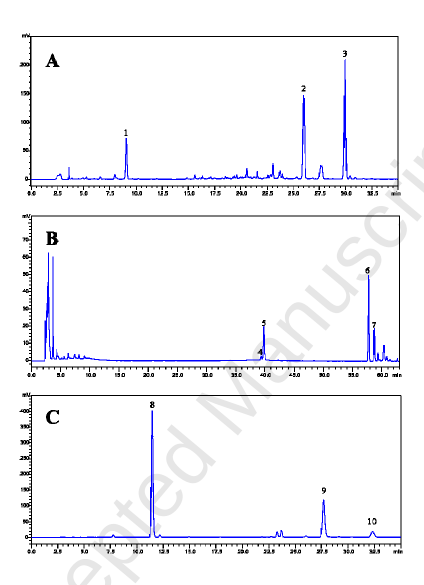


Figure 2

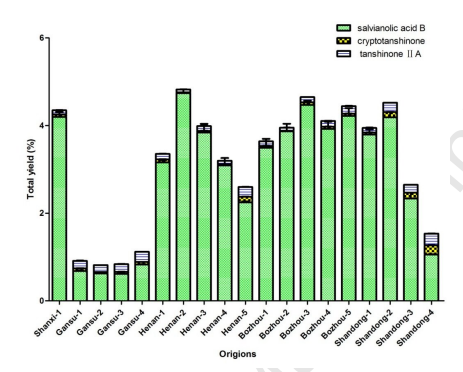
395

396



397

398 399 Figure 3



399400

401

Figure 4

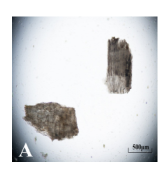
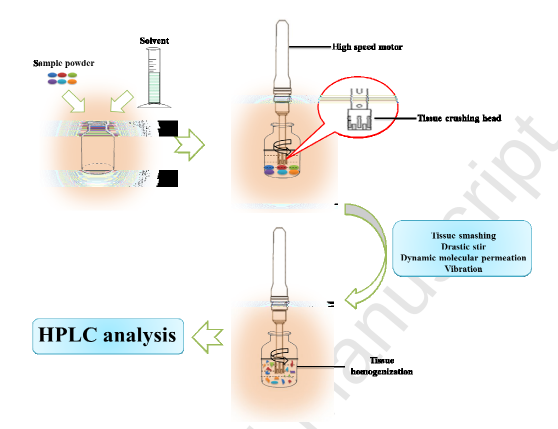




Figure 5





405	Highlights
406	
407	■ Three chemical constituents from Salvia Miltiorrhiza were extracted by new designed
408	TSURE
409	 Optimized TSURE offered good extraction efficiency that was comparable or even better
410	than conventional methods
411	 TSURE method was extended to the quality control of Salvia Miltiorrhiza
412	 Application of TSURE to other herbal medicines was also explored and similar results were
413	discovered
414	 Mechanisms of TSURE were discussed in detail for the first time
415	

415 **Table 1**

Comparison of extraction efficiencies of target compounds in S. miltiorrhiza by TSURE and

conventional extraction methods under the optimal conditions (mean \pm S.D., n = 6).

Method	Extraction	Extraction	Salvianolic	Cryptotan-	Tanshinone	Total	RSD
	time(min)	volume	acid B	-shinone	IIA	yield	(%)
		(ml/g)	(mg/g)	(mg/g)	(mg/g)	(mg/g)	
TSURE	2	80	1.158±0.031	0.208±0.056	0.291±0.106	1.657±0.077	4.673
UAE	40	80	0.950±0.016	0.208±0.007	0.286±0.005	1.444±0.042	2.877
HRE	120	80	1.083±0.018	0.298±0.003	0.270±0.004	1.589±0.039	2.426

TSURE: tissue-smashing based ultra-rapid extraction;

420 UAE: ultrasonic assisted extraction;

421 HRE: heat reflux extraction

422

422 **Table 2**

Calibration curves, test range, and LODs for 3 analytes in *S. miltiorrhiza* by LC-UV.

No.	Analytes	Calibration curve	r^2	Lincor rongo (ug)	LOQs	LODs
				Linear range (μg)	(ng)	(ng)
1	salvianolic acid B	y = 11962532x - 496388	0.9991	0.32-10.10	1.60	0.32
2	cryptotanshinone	y = 51551852x + 208610	0.9996	0.13-4.30	0.06	0.03
3	tanshinone IIA	y = 59718035x + 102353	0.9996	0.07-2.10	0.10	0.02

LOD: The limitation of detection (S/N > 3); LOQ: The limitation of quantification (S/N > 10)

Table 3
 The precisions and recoveries of 3 target components in *S. miltiorrhiza* obtained by TSURE method.

Analytes	Repeatability	Recovery (n=3)					
	(n=5, %)	O (mg)	S (mg)	F (mg)	Recovery (%)	RSD (%)	
Salvianolic acid B	1.65	2.16	1.60	3.71	97.07	2.60	
		2.16	2.15	4.36	102.43	0.30	
		2.15	3.20	5.31	98.53	1.86	
cryptotanshinone	1.90	0.50	0.25	0.74	93.47	1.99	
		0.50	0.49	0.99	100.53	4.81	
		0.57	0.73	1.29	99.33	2.13	
tanshinone IIA	2.18	0.52	0.38	0.92	105.20	2.63	
		0.65	0.64	1.27	96.00	3.71	
		0.66	0.89	1.56	101.60	1.53	

O: original contained; S: spiked; F: found

430