A study on the identification of habitats and determination of sulfur dioxide residue of Radix Astragali by UV-vis-SWNIR diffuse reflectance spectroscopy

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The feasibility of identifying habitat of sulfur fumigated Radix Astragali and of determining its sulfur dioxide residue using UV-vis and short wave near-infrared diffuse reflectance spectra coupled with chemometrics was first investigated. The results show that by using UV-vis-SWNIR DRS combined with PCA, the habit of Radix Astragali is identified definitely, whether it is sulfur-fumigated, or not sulfur-fumigated; after waveband selection, SG smoothing and excluding outliers, PLS modeling results were improved with a \( R^2 \) of 0.9238, 0.9104 and 0.9322 for calibration set, cross validation set and prediction set, respectively. A convenient and rapid method for identification of the habitat of sulfur fumigated Chinese herbal medicines and determination of sulfur dioxide residue by UV-vis-SWIR DRS was proposed.

1 Introduction

Identification of herbal medicines is an important part of their quality control. Traditional identification of herbal medicines is based on the difference in source, character identification, physiochemical identification and microscopic identification.  But these methods depend on experience to some extent and may not fully reflect the medicinal quality.  Afterwards, thin-layer chromatography, 2–5 HPLC, 6,7 and fingerprint technology, 8–11 based on the extraction of related indexes, were developed and widely used for the identification and quality evaluation of herbal medicines. However, they are complicated, laborious, time-consuming and solvent-consuming due to requiring chemical sample-pretreatments such as extraction, separation, and concentration. Therefore, in recent years, rapid and nondestructive methods without chemical pretreatment have gained fast development and wide acceptance. Infrared spectroscopy, 12–14 Fourier transform infrared (FTIR) spectra, 15 near infrared reflectance spectroscopy (NIR), 16,17,18 FT-Raman spectroscopy 19 and THz-TDS 20,21 methods have been reported for this objective. However, they have some disadvantages such as using sophisticated and expensive instruments. Additionally, they do not include UV-vis spectral region which contains a wealth of information regarding electron-energy level transition of molecules. THz-TDS also has other shortcomings: a lot of Chinese medicines have no obvious characteristic absorption peak in the THz frequency region, 22,23 the strong interference from water absorption makes it need to set the instrumental system in an airtight enclosure with nitrogen gas infusion to keep the relative humidity as less than 8% even 1%, to overcome the effects of water absorption. 2,20,23

On the other hand, sulfur fumigation was a traditional processing way used widely for some kinds of Chinese medicinal materials (CMM), which can make CMM mothproof and mildewproof, have beautiful appearance and bright color, 24,25 and is helpful for storage. However, sulfur fumigation may have a great impact on the chemical compositions and pharmacological effects of CMM, 24,25 make more residue of SO2 26,27 and more residues of toxic and harmful substances such as lead, arsenic, mercury and chromium. 28,29,30 Therefore, control of SO2 residue in CMM is necessary and it has been becoming more and more strictly. China has stipulated that residue amount of sulfur dioxide in herbal medicines should not exceed 150 mg kg \(^{-1}\), sulfur dioxide residue amount in 10 kinds of herbs and slices such as Rhizoma Dioscoreae, Radix Achyranthis Bidentatae, Kudzu root, Radix Asparagi, Rhizoma Gastrodiae, Radix Trichosanthis, Rhizoma Bletillae, Radix Paeoniae Alba, Rhizoma Atractylodis Macrocephalae and Radix Codonopsis should not exceed 400 mg kg \(^{-1}\). 24

Chinese official determination methods of SO2 residue in herbal medicines include acid-base titration, gas chromatography and ion chromatography. The first and third need a pre-treatment of steam distillation, they are complicated in operations and time-consuming. Therefore, effective and rapid determination of sulfur dioxide residue in herbal medicines becomes a key of quality supervision. It is expected to establish fast and efficient methods. FTIR spectroscopy has been
reported to be used to identify sun-dried and sulfur-fumigated *Radix Paeoniae Alba,*23 *Radix Angelicae Sinensis,*26 but they needed extraction and didn’t determine SO2 residue.

UV-vis-shortwave near infrared diffuse reflectance spectroscopy (UV-vis-SWNIR DRS) has advantages containing the information regarding electron-energy level transition as well as vibrational motion of molecules, being simple, fast and non-destructive. We have reported its applications on qualitative and simultaneous quantitative analysis of cimetidine polymorphs,27 qualitative analysis of chiral alanine,28 simultaneous determination of amiloride and hydrochlorothiazide in a compound tablet,29 qualitative and quantitative analysis of chiral alanine,28 rapid and simultaneous determination of soil properties,30 in which the importance of UV-vis spectral region for analysis has proven. But its use for the identification of habitats of herbal medicines and determination of SO2 residue in herbal medicines has not been reported.

*Radix Astragali,* a kind of Chinese traditional medicine mainly distributed in North China, has following clinical efficacies: tonifying qi and lifting yang, consolidating exterior for arresting sweating, inducing diuresis for removing edema, creating body fluid and nourishing blood, dissipating stagnation and promoting arthromyodynia, expelling toxin and eliminating pus, restraining ulceration and growing skin.31 It can enhance the immune function of patients, viral interferon induction ability, hypoxia tolerance of the patients’ body, promote metabolism and the antihypertensive and liver-protecting function. It is widely used in clinical treatment of many diseases. *Radix Astragalus* is prone to pests during storage.

Therefore, the objective of this study is to establish a simple, fast and non-destructive method to identify habitats of herbal medicines and to determine their SO2 residues, by using UV-vis-SWNIR DRS combined with chemometrics, and *Radix Astragali* was used as a research object.

## 2 Experimental

### 2.1 Sample and sulfur fumigation

*Radix Astragali* from three provinces Sichuan, Shanxi and Gansu were purchased from Wu Kuaishi market of Chinese herbal medicines in Chengdu, Sichuan province. In appearance, *Radix Astragali* from Sichuan is partial white, *Radix Astragali* from Gansu and Shanxi are yellowish, as shown in Fig. 1.

About 150 g of *Radix Astragali* of each habitat was weighed, wetted by spraying 40 mL of water and kept in a plastic bag for 12 h, then fumigated for 2 days by the smoke generated from the combustion of a certain amount of sulfur in a homemade device. After that the fumigated *Radix Astragali* was dried in an air dry oven of 30 °C for 2 days, milled with a household food disintegrator and sieved. The powder samples through 80-mesh sieve were collected and kept under seal.

During milling and sieving, it was found that more fibrous residue was not able to pass through 80-mesh sieve in *Radix Astragali* from Sichuan than from Gansu and Shanxi. The fibrous residue was cut with scissors and milled repeatedly until they can pass.

Five levels of sulfur fumigation were implemented for *Radix Astragali* of each habitat. The accurate sulfur dioxide residues in all the samples of *Radix Astragali* (three habitat, 5 levels of sulfur fumigation) were determined by acid–base titration method listed in Chinese Pharmacopoeia (2015)34 and the obtained values were treated as the reference values of sulfur dioxide residues.

The powder samples of certain appropriate levels of sulfur fumigation from a same habitat were weighed accurately, mixed according to a certain ratio to obtain 80 samples with different contents of SO2, so 240 spectral samples obtained for the three habitats.

### 2.2 Apparatus

S3000 fiber optic spectrometer (Race-Technology Co., Ltd, Hangzhou, China) equipped with a 3648-element linear silicon CCD array detector (Toshiba TCD 1305), a light source (Oceans Optics Inc., USA), a Y-type optical fiber probe and a home-made sample cell made from dark gray PVC was used to measure the UV-vis-SWNIR diffuse reflectance spectra.37

Sieves (Surwit Co., Ltd, Hangzhou, China) of 80, 100 and 120-mesh sieves (0.200, 0.150 and 0.125 mm nominal diameters) were used to screen the milled samples.

### 2.3 Procedures

#### 2.3.1 Spectral measurement

0.150 g of each powder sample of *Radix Astragali* of each habitat was weighed and filled in the home-made sample cell and flattened with a flat round bar. Then the optical probe was placed vertically on the upper surface of the powder sample to measure UV-vis-SWNIR diffuse reflectance spectra in the range of 200–1100 nm under the following measuring conditions: exposure time of 250 ms, integral sampling time of 1 ms, average number of 15 and smoothing number of 15, with a resolution of 0.33 nm and a spectralon as background reference. Each sample packed in the sample cell was measured twice before and after the sample was re-filled and the optical probe was reset, and their average was treated as a raw spectrum of the sample. In total, 240 spectra for the powder samples of *Radix Astragali* from three habitats were obtained.

#### 2.3.2 Identification of habitats

Identification of habitats was carried out by applying principle component analysis (PCA) on the UV-vis-SWNIR diffuse reflectance spectra, using Unscrambler ver 9.7.

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**Fig. 1** Appearance of *Radix Astragali.*
### 2.3.3 Determination of SO₂ residues.

Determination of SO₂ residues was implemented by applying partial least square regression (PLSR) modelling on the UV-vis-SWNIR diffuse reflectance spectra, using Unscrambler ver 9.7.

### 2.3.4 Evaluation of model performance.

Root mean square error of prediction (RMSE), determination coefficient of prediction ($R^2$) and relative predictive determinant of prediction (RPD) were used as model performance indicators. Their calculation can be found in the literature.³⁸

According to Saeyes et al.³⁴ and Vohland et al.,³⁹ $RPD > 3.0$ and $R^2 > 0.90$ are considered to be indicative of an excellent prediction; $2.5 < RPD < 3.0$ and $0.81 < R^2 < 0.90$ indicate good prediction; $2.0 < RPD < 2.5$ and $0.66 < R^2 < 0.81$ make approximate quantitative predictions possible; $1.5 \leq RPD < 2.0$ and $0.50 \leq R^2 < 0.66$ reveal a possibility to distinguish between high and low; $RPD < 1.5$ or $R^2 < 0.50$ indicate that prediction is bad.

### 3 Result and discussion

#### 3.1 Spectra

The original UV-vis SWNIR DRS showed that the absorbance in the range above 1090 nm and below 213 nm is negative. It might be due to instrument noise. Therefore, spectrum below 213 nm and above 1090 nm was cut off, leaving the spectral range of 213–1090 nm for analysis use, as shown in Fig. 2. It can be seen that the spectral waveforms were similar, but spectral intensity differs to some extent with habitats or SO₂ residue levels, especially as for the samples from Sichuan in the ranges of 240–450 nm and 1000–1100 nm.

#### 3.2 Identification of habitats

Identification of habitats was carried out on the base of wavelength selection. The aim of wavelength selection is to select the informative wavelengths which are relevant to the property of interest, in other words, to remove the uninformative and/or interfering variables to construct a reliable and interpretable calibration model with good prediction accuracy.

The forward evolving window correlation coefficient method was used to select characteristic wavelength range for PCA. The intra-class correlation coefficient of every kind of habitat and the average of three inter-class correlation coefficients between every two kinds of habitat samples were calculated, the results are shown in Fig. 3.

The larger the intra-class correlation coefficient and the smaller the inter-class correlation coefficient, the more conducive the realization of sample classification and discrimination. Fig. 3 shows that the correlation coefficients were all greater than 0.97 in the full spectral range, and in the range of 213–530 nm the average of inter-class correlation coefficients was smaller than the intra-class correlation coefficients. This provides the possibility to classify habitats. Therefore, besides the full spectral range of 213–1090 nm, various ranges of 213–1090 nm, 213–590 nm, 213–510 nm, 320–510 nm, 213–350 nm, 213–310 nm, 240–310 nm and 213–530 nm were tried as modelling wave bands in following PCA modelling.

In PCA modelling, the percentage of cumulative variance of the first N principle components vs. the total variance was calculated. Applying PCA in full spectral range of 213–1090 nm gave the first cumulative variance of 60%, the second cumulative variance of 86% and the third cumulative variance of 94.5%, while PCA in the range of 213–310 nm gave the best results, with the first cumulative variance of 72.5%, the second cumulative variance of 95% and the third cumulative variance of 99%. The three-dimensional projection using the range of 213–310 nm is shown in Fig. 4, which demonstrates the better effect of identification of habitats.

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Fig. 2 Diffuse reflectance spectra of powder samples.

Fig. 3 Forward evolving window correlation coefficient method (1) Sichuan; (2) Shanxi; (3) Gansu; (4) average of inter-class.

Fig. 4 Three-dimensional projection of the three habitats of *Radix Astragali* on the first three principle components of PCA in spectral range of 213–310 nm.
In Fig. 4, compared with *Radix Astragali* from Gansu and Shanxi, *Radix Astragali* from Sichuan dispersed in a larger scope, which may be due to the powder size of *Radix Astragali* from Sichuan. As has been mentioned above, more fibrous residue in Sichuan *Radix Astragali* powder was found than in Gansu *Radix Astragali*, Shanxi *Radix Astragali*. Therefore, it was considered that there were more fibrous or long-shaped particles in Sichuan *Radix Astragali* powder samples. This is disadvantageous to obtain a smooth sample surface when filled in the sample cell. Rough surface of a packed sample may cause larger fluctuations of the measured spectra. But in this case, this still permit a clear discriminant of the three habitats of *Radix Astragali*. This implies that the difference of natures or compositions between the three habitats of *Radix Astragali* is larger than the difference caused by smoothness of sample surface.

The results show that UV-vis spectrum plays an important role in identification of origin of *Radix Astragali*; by using UV-vis-SWIR DRS combined with PCA, habitat identification of *Radix Astragali* is feasible, whether it is sulfur-fumigated, or not sulfur-fumigated.

### 3.3 Determination of SO₂ residues

The total 240 spectra of *Radix Astragali* from 3 habits were divided into a calibration set of 180 samples and a test set of 60 samples by K/S algorithm.

**3.3.1 Wavelength selection.** Wavelength selection for PLSR model was carried out by correlation coefficient method and iPLS method, respectively.

Fig. 5 is the plot of the correlation coefficient between SO₂ residue and absorbance vs. wavelength. The bands with a $R^2$ greater than 0.2421, that is, the mean value of absolute values of all $R^2$ in the full spectrum, were used to build PLSR model for prediction of SO₂ residue. But the results were not satisfactory: the square correlation coefficient $R^2$ between predicted values of SO₂ residue and their reference values for the calibration set and prediction set was 0.7805 and 0.7406, respectively.

Then, wavelength selection for PLSR model was carried out by iPLS method. Fig. 6 and 7 is the plot of RMSECV vs. wavelength when the full wavelength range was divided into 2 intervals (2-iPLS) and 3 intervals (3-iPLS), respectively.

Comparing Fig. 6 and 7, it can be known that the RMSECV with the band (213–691 nm) selected by 2-iPLS and the RMSECV with the band (213–542 nm) by 3-band iPLS were both lower than that (horizontal line) with the whole band, but the former was lower. So the band of 213–691 nm selected by 2-iPLS method was used in further study, and the modelling effects were listed at the fourth line in Table 1, better than those using the wavelength selected by correlation coefficient method (Fig. 5).

**3.3.2 Spectral data pre-treatment.** Ultraviolet-visible spectra contain not only sample information but also some irrelevant information and noise, such as instrument noise, sample background and stray light. Therefore, it’s significant to select spectral pre-treatment methods to build robust quantitative model.

In this study, spectral data pre-treatment was performed by the 18 pre-treatment methods shown in Table 1, using the optimum modelling waveband (213–691 nm) selected in Section 3.3.1. It is found that modelling with SG smoothing pre-treatment has the best effect.

**3.3.3 Outliers exclusion.** Generally, as for quantitative analysis based on spectra, there may be some outliers consisting of spectral outliers and chemical value outliers. $F$-Test and PCA coupled with Mahalanobis Distance (MD) is applied on...
identifying chemical value outliers and spectral outliers, respectively. In the calculation process of PCA and MD, the number of PCs is selected according to the cumulative contribution of PCs set as 95%, and the threshold \( \Delta t \) of M Di s decided by setting the weight coefficient \( t \) as 2.

In this study, the outlier removal was carried out on the basis of SG smoothing pre-treatment having the best modelling effect. By computation, the value of \( \Delta t \) was 2.7789. The chemical value outliers were determined by the threshold value \( F_a = 4.0099 \) computed by \( F \) distribution with 1 degree of freedom in the numerator and 239 degrees of freedom in the denominator and 95% of confidence level. As a result, there were 21 outliers shown in Fig. 8. The results of modelling before and after outlier exclusion were showed in Table 1.

### Modelling effects
After waveband selection, SG smoothing and excluding outliers, modelling results were improved with a \( R^2 \) of 0.9238 for calibration set, a \( R^2 \) of 0.9104 for cross validation set and a \( R^2 \) of 0.9322 for validation set as shown in Table 1. Additionally, RPD by calculation was 3.8405. According to Saey et al. and Vohland et al., \( R^2 > 0.90 \) and RPD > 3.0 at same time are considered to be indicative of an excellent prediction. The results demonstrate that an excellent prediction has been achieved. The plots of predicted values of SO2 residue vs. their reference values for the calibration set and prediction set are gave in Fig. 9 and 10.

### Table 1 Modelling and verification of determination of residual SO2 in Radix Astragali by PLS

| Pretreatment          | N  | \( R^2 \) | RMSEC | RPD   | \( R^2 \) | RMSECV | RPD   | \( R^2 \) | RMSEP | RPD   |
|-----------------------|----|------------|-------|-------|------------|--------|-------|------------|-------|-------|
| Raw spectrum          | 8  | 0.8577     | 214.97|       | 0.8311     | 234.73 |       | 0.8526     | 213.05|       |
| 2-iPLS                | 9  | 0.8776     | 199.31| 2.858 | 0.8423     | 227.48 | 2.518 | 0.8711     | 201.56| 2.785 |
| SG                    | 9  | 0.877      | 199.81| 2.851 | 0.8461     | 224.66 | 2.549 | 0.8702     | 194.43| 2.776 |
| NOR                   | 8  | 0.863      | 210.90|       | 0.8374     | 231.30 |       | 0.8613     | 207.87|       |
| Spectroscopic         | 10 | 0.883      | 194.93|       | 0.8191     | 243.55 |       | 0.8336     | 1600.7|       |
| MSC                   | 6  | 0.8382     | 229.17|       | 0.8092     | 250.49 |       | 0.8431     | 217.54|       |
| Noise                 | 6  | 0.9645     | 107.30|       | 0.558      | 380.76 |       | 0.4631     | 393.93|       |
| 1st D                 | 6  | 0.9743     | 91.37 |       | 0.7332     | 296.41 |       | 0.0136     | 1402.7|       |
| 2nd D                 | 1  | 0.8272     | 236.84|       | -0.2557    | 641.72 |       | 0.0681     | 1055.2|       |
| Baseline              | 8  | 0.8617     | 211.92|       | 0.8311     | 235.67 |       | 0.8633     | 204.45|       |
| SNV                   | 6  | 0.8424     | 226.17|       | 0.8185     | 244.43 |       | 0.8464     | 215.54|       |
| Center and scale      | 9  | 0.8776     | 199.31|       | 0.8457     | 224.94 |       | 0.8711     | 201.56|       |
| Reduce                | 9  | 0.8771     | 199.76|       | 0.8486     | 222.87 |       | 0.8711     | 201.54|       |
| SNV + SG              | 7  | 0.8536     | 217.98|       | 0.8282     | 237.40 |       | 0.8642     | 299.46|       |
| SG + SNV              | 8  | 0.8743     | 201.98|       | 0.8426     | 227.26 |       | 0.6794     | 3951.1|       |
| SG + MSC              | 8  | 0.8692     | 206.08|       | 0.8416     | 229.06 |       | 0.7165     | 304.12|       |
| SG + reduce           | 9  | 0.877      | 199.84|       | 0.8479     | 223.08 |       | 0.871      | 201.62|       |
| SG + noise            | 6  | 0.9674     | 102.92|       | 0.5888     | 367.29 |       | 0.5935     | 346.05|       |
| SG + 1st D           | 8  | 0.9653     | 106.20|       | 0.801      | 256.00 |       | 0.0002     | 3800.9|       |
| MSC + SG              | 6  | 0.8504     | 220.38|       | 0.8286     | 237.90 |       | 0.8532     | 211.20|       |
| 2-iPLS + SG + O-E     | 9  | 0.9238     | 153.38| 3.623 | 0.9014     | 175.81 | 3.185 | 0.9322     | 142.55| 3.840 |

\( a \) N: number of latent variable for modelling, O-E: Outliers Exclusion.

Fig. 8 Outliers exclusion.

Fig. 9 Plot of predicted value vs. reference value of calibration set by PLS.
The limit of detection (LD) was calculated to estimate the lowest content of residual SO$_2$ that can be measured. The limit of detection was determined from multiple measurements ($n = 9$) of the spectral response of samples containing 60 mg residual SO$_2$ per kg of Radix Astragali from Gansu and Shanxi, 267 mg residual SO$_2$ per kg of Radix Astragali from Sichuan. The LDs were calculated by multiplying the standard deviation by three and then dividing by the slope of the calibration curve, as shown by eqn (1):

$$LD = 3 \times \text{S.D.}/\text{slope}$$  

The LD of residual SO$_2$ was found to be 360 mg kg$^{-1}$ for Radix Astragali from Gansu and Shanxi, 550 mg kg$^{-1}$ for Radix Astragali from Sichuan. Although they are not as low as 150 mg kg$^{-1}$, but less than or near 400 mg kg$^{-1}$, the limit of sulfur dioxide residues in ten traditional Chinese medicines allowed by China. Compared with the qualitative analysis of sulfur dioxide residues in the report which used FTIR, the sensitivity of our method is greatly improved. It is believed that in future with the improvement of optical fiber spectrometer hardware system, especially the improvement of the stabilities of light source and spectrometer, the precision of the proposed method will be improved, consequently, the detection limit will be reduced.

4 Conclusions

The application of UV-vis and short wave near-infrared diffuse reflectance spectrum coupled with chemometrics on the identification of habitat and determination of sulfur dioxide residue of sulfur fumigated Chinese herb medicine was investigated first. UV-vis spectrum plays an important role in both the identification of CHM and determination of SO$_2$ residue. By using UV-vis-SWNIR DRS combined with PCA, habitat of Radix Astragali is identified definitely, whether it is sulfur-fumigated, or not sulfur-fumigated; through wavelength band selection, SG smoothing and excluding outliers, modelling results were improved with a $R^2$ of 0.9238, 0.9104 and 0.9322 for calibration set, cross validation set and validation set, respectively.

The results imply that a convenient, rapid method for the identification of habitat of sulfur fumigated Chinese herbal medicines and determination of sulfur dioxide residue by UV-vis and short wave near-infrared diffuse reflectance spectrum is feasible.

Acknowledgements

This work was supported by the Science and technology special fund of Sichuan Provincial Administration of Traditional Chinese Medicine, China (No. 2016Q058) and the Scientific Research Foundation of the Education Department of Sichuan Province, China (No. 16ZB0122).

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