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ATR-FTIR combined with chemometrics to distinguish geographical indications from non-geographical indications *Gastrodia elata* Bl

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ABSTRACT

Gastrodia elata Bl (*G. elata*), a medicinal and edible homologous variety, has been artificially cultivated in different regions of China to meet the growing demands of human beings. In this study, attenuated total reflection/Fourier transform infrared spectroscopy (ATR/FTIR) combined with chemometrics [Principal component analysis (PCA), Partial least squares discrimination (PLS-DA), Support vector machines (SVM), and Data-driven soft independent modeling of class analogy (DD-SIMCA)] was used to differentiate between geographical indications and non-geographical indications of *G. elata*. PLS-DA, after the application of SNV+SD spectral preprocessing, achieved 100% accuracy on the training set and 88.89% on the test set, respectively. Under SG+SD conditions, SVM outperformed PLS-DA with 100% training set accuracy and 94.74% for the test set. A ResNet model that used synchronous 2DCOS data successfully distinguished *G. elata* from Yunnan and Guizhou, achieving 100% accuracy across training, test, and external validation sets. These findings support that ATR-FTIR and chemometrics can be utilized to effectively identify the geographical origin of *G. elata*, with potential applications for other medicinal and edible plants.

1. Introduction

The dried tuber of the Gastrodia elata Bl (G. elata) is a kind of perennial parasitic plant widely distributed in China, Bhutan, Nepal, India, Japan, Korea, Siberia, and other countries, which possesses both high medicinal and edible value [1]. In traditional Chinese medicine (TCM), G elata calms wind, stops spasms, soothes liver-yang, dispels wind, clears collaterals, etc., and can be leveraged for children's convulsions, epilepsy, tetanus, headache, dizziness, paralysis of hands and feet, numbness of limbs, rheumatism, and arthralgia [2]. It has been found that the chemical composition of G. elata encompasses phenols (e.g. Gastrodin, 4-hydroxybenzenol and balisin), polysaccharides, sterols (e.g. Beta-sitosterol), and organic acids (e.g. Citric acid and succinic acid) [3,4]. Also, modern pharmacological studies have proven that G. elata has anti-inflammatory, anti-epileptic, anti-stroke, antianxiety, and anti-depression pharmacological effects [5]. Beyond its role as one of the commonly used TCMs, G. elata is also a functional food. The Chinese government officially announced in 2019 that it can be applied as a "medicinal food homologous" plant [6]. In daily life, G. elata is employed to make various tonic diets, such as fish head soup, stewed chicken and ham, yam congee, etc., which indicates the valued and promoted trend in the field of TCM and food.

Geographical indication (GI) refers to products that grow in a certain geographical area, with a specific quality, reputation, or other characteristics, and are mainly determined by natural and human factors in the area. *G. elata* grows in Xiaocaoba Town, Yiliang County, Zhaotong

City, Yunnan Province, and Dafang County, Bijie City, Guizhou Province. Lv et al. [7] proposed that it is necessary to distinguish between GI and non-GI G. elata since GI products may attract consumers and influence their purchasing decisions. Experts can distinguish between GI and non-GI G. elata through visual perception and sensory evaluation, while this analysis is highly subjective and lacks reproducibility due to the variability of individuals and samples [8]. The traditional identification method is to collect G. elata samples from different origins and harness instruments like high-performance liquid chromatography (HPLC), mass spectrometry (MS), gas chromatography-mass spectrometry (GC-MS), etc., for analysis. After detecting the content of certain chemical components of G. elata, the equipment quantitatively uses it as a testing parameter to evaluate the quality level and category. Through using UPLC-ESI-QTOF-MS/MS and HPLC-UV, Su et al. [9] determined the content of 7 components of G. elata. Instruments including HPLC and GC-MS can provide highly accurate analytical results and then aid in identifying the differences between G. elata samples of different origins and assess their quality, whereas HPLC and GC-MS are cumbersome to operate and use chemicals harmful to the environment and human health. Consequently, non-destructive testing technology possesses an important application prospect in the analysis of G. elata and other Chinese medicinal materials.

Spectroscopy is the most widely applied method in geographical origin classification, and attenuated total reflectance-Fourier transform infrared spectroscopy (ATR-FTIR) analysis is characterized by the advantages of simple operation, fast analysis speed, no damage to

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samples, and requiring fewer samples and represents a pivotal technical approach widely used in various fields [10]. Spectroscopic techniques, especially ATR-FTIR, contribute much to the field of modern analytical chemistry and biomaterials. ATR-FTIR technology is used in basic chemical analysis and is also widely employed in exploring the homology of medicine and food and the quality identification of Chinese medicinal materials. ATR-FTIR has been utilized for the identification of *G. elata* origin [11,12], accurate identification of *G. elata* GIs [13], and the identification of Fritillaria species from various plant sources [14]. Being an efficient analytical method, ATR-FTIR can generate high-throughput, non-specific, and redundant data [15]. This feature can still bring some challenges in data analysis, even it is a significant advantage in data acquisition. With the development of chemometrics, an increasing number of efficient data processing methods have been proposed to tackle the complexity of infrared spectral data. Combined with Principal component analysis (PCA), Partial least squares discrimination (PLS-DA), Support vector machines (SVM), and other analysis methods, He et al. [16,17] used FT-NIR spectral data to identify the drying temperature of Amonum tsao-ko, and determined the quality grade of wine by ML model based on RF and K-Nearest Neighbors (KNN) algorithms. To address adulteration and abuse in the aboveground parts of herbal medicines, Song et al. constructed and evaluated ten different machine learning (ML) models based on ATR-FTIR spectroscopy [18]. Correspondingly, it is not difficult to find from the literature that the combination of ATR-FTIR and chemometrics gives full play to its advantages in identification.

This research aims to use ATR-FTIR technology, combined with chemometric methods, to quickly and accurately identify GI and non-GI G. elata. First of all, a total of 248 G. elata samples from Yunnan and Guizhou were dried under the same conditions. Secondly, the effectiveness of G. elata sample classification for GI and non-GI was evaluated separately through PCA. Thirdly, after the implementation of 10 spectral preprocessing methods, PLS-DA, SVM, and Data-driven soft independent modeling of class analogy (DD-SIMCA) models were constructed to compare the adaptability of ATR-FTIR technology integrated with chemometrics in distinguishing GI from non-GI G. elata. Finally, a Resnet model was established based on G. elata 2DCOS data to distinguish the G. elata production areas in Yunnan Province and Guizhou Province. In this study, a reliable G. elata identification model was set up for GI and non-GI, which provided a scientific basis for analyzing the geographical sources of G. elata, exhibiting important application prospects in the medicinal and edible industries, quality control of medicinal materials, etc.

2. Materials and Methods

In 2023, fresh cultivated *G. elata* samples were collected in Yunnan Province (Zhaotong Xiaocaoba and Not xiaocaoba (Yongshan County and Zhenxiong County) and Guizhou Province (Dafang County and Not dafang County (Hezhang County and Jinsha County) in China, totaling 248 samples (Table 1). They were identified by Professor Huang Hengyu of Yunnan University of Traditional Chinese Medicine. The relevant sample information has been shown in Figure 1. All tubers were washed, dried, crushed, and sieved through a 100-mesh sieve. The powder was stored in a plastic sealed bag at room temperature for further analysis.

Table 1.	Gastrodia	elata Bl.	Sample	information.
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Origin		Abbr.	Total	Train set	Test set
Zhaotong	Xiaocaoba town of yiliang county	XCB	59	41	18
city, Yunnan	Yongshan county	YS			
Province	Zhenxiong county	ZX	60	42	18
Bijie city,	Dafang county	DF	68	48	20
guizhou Province	Hezhang County	HZ			
	Jinsha County	JS	61	43	18

Note: The geographical indication production areas are XCB and DF, while the nongeographical indication production areas comprise YS, ZX, HZ, and JS, corresponding to NXCB (comprising YS and ZX) and NDF (comprising HZ and JS).

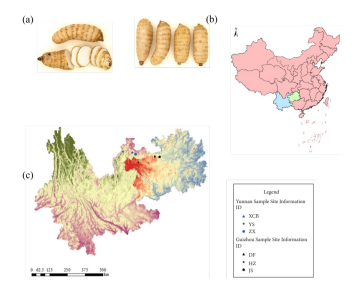


Figure 1. (a) Gastrodia elata Bl. Sample. (b) and (c) Sample site information.

2.1. Spectral collection

A Fourier transform infrared spectrometer (Perkin Elmer, USA) equipped with ZnSe attenuated total reflection attachment and OMNIC 9.77 (Thermo Fisher Scientific, USA) software was used for spectral acquisition and analysis. *G. elata* powder was placed on ZnSe crystal material for spectral scanning. ATR-FTIR spectral data within the wavenumber range of 4000-400 cm⁻¹ was recorded. Each sample was scanned 64 times with a resolution of 8 cm⁻¹ and repeated 3 times under the same conditions to verify accuracy and obtain the average spectrum for further analysis.

2.2. Spectral data preprocessing

The collected sample spectra were converted into a dataset using SIMCA-P+14.1 software (Umetrics, Umea, Sweden). The original ATR-FTIR spectrum was pre-processed to remove interference information, such as background, noise, and baseline drift. This study applied preprocessing methods, such as savitzky-golay smoothing (SG), second derivative (SD), multiple scattering correction (MSC), and standard normal variable (SNV) smoothing separately and in combination. SG smoothing is a commonly used spectral data smoothing method that smoothens curves by local polynomial fitting of spectral data, reduces the effect of noise, and maximizes the retention of the peak characteristics of the original spectral signal [19]. The SD can enhance the slope change of spectral data, extract fine features, and is often used for preprocessing to reduce random noise and improve signal frequency resolution [20]. The MSC corrects the scattering effect in the spectral data, making the spectral intensity more consistent between different samples and helping to improve the comparability of the data [21]. The SNV corrects the spectral baseline shift, eliminating differences due to equipment drift, sample placement, and particle size changes. It improves the ability of spectral comparison and understanding, highlights the main characteristics of the spectrum, and reduces the impact of scale differences between samples on the analysis results, helping to find small spectral changes [22]. SD with nine smoothing points is used to amplify FTIR and NIR spectral differences and eliminate baseline bias [23]. Chen et al. [12,24] performed SG smoothing pretreatment on the selected NIRS band, and applied SNV and MSC to NIRS and ATR-FTIR spectral pretreatment for honey adulteration.

2.3. Soft

ATR-FTIR transmittance was converted to absorbance using the OMNIC 9 software. Spectral data was preprocessed, and PLS-DA modeling was performed using SIMCA 14.1 software. Origin 9.1 software was used to draw. MATLAB 2023a software was used to establish SVM and DD-SIMCA models, as well as acquire of 2DCOS images. ResNet was built in the Spyder (Anaconda 3) software.

2.4. Model construction

Before data modeling, the sample was randomly divided into 70% training and 30% test sets by the Kennard-Stone algorithm, and stoichiometric analysis included multivariate statistical techniques, such as PCA, PLS-DA, SVM, and Random forests (RF). The performance of these classification algorithms on GI and non-GI spectral preprocessing data sets was compared.

Principal component analysis (PCA) is a kind of multivariate statistical method that transforms the collected data dimension and data of dimension reductions after further linear classification, a reduction in the number of features and complexity, while retaining most of the original data information [25]. Thus, the classification and differentiation of samples from different places can be realized, and the potential hidden characteristics and rules can be found.

PLS-DA is a method that combines Partial Least Squares Regression (PLS) and Discriminant Analysis (DA), commonly used for processing high-dimensional data with classification labels [26,27]. PLS-DA aims to find the component with the greatest covariance between the observed variable (X) and the response variable (Y) while achieving the best distinction between classes. PLS-DA is suitable for processing high-dimensional data with classification labels, helping to solve classification and prediction problems in data analysis while providing solutions for data reduction and feature selection.

Support vector machine (SVM) is a classical nonlinear supervised learning modeling method, which has outstanding advantages for the analysis and modeling of nonlinear relations of high-dimensional input variables. By finding an optimal hyperplane, the distance between sample points and this hyperplane is as far as possible [28]. The goal of SVM is to maximize the spacing to ensure better classification of different classes. This study employs the grid search algorithm (GS) to optimize the values of parameters c and g. To be specific, parameter c represents the degree of error acceptance of the model, in which too large or small c can bring poor generalization of the SVM model. Besides, the parameter g attached to the RBF function chosen as the kernel determines the distribution of the data set mapped to the high-latitude factor space.

DD-SIMCA is a multi-variable data analysis method based on the soft independent modeling of class analogy (SIMCA) algorithm. First, the scoring distance (SD) and orthogonal distance (OD) of each sample in the training set were calculated. Then, the acceptance region of the target class was established based on different methods. The purpose of the acceptance region is to determine whether the test set is considered to belong to the target class. Finally, the non-target class samples were substituted into the trained model for classification prediction [29,30].

A residual network (ResNet) is a deep convolutional neural network structure. Compared with ordinary neural networks, ResNet does not need to learn the entire input and output process but only the difference between output and input (residual) to minimize the training difficulty [31]. By introducing residual blocks, the ResNet enables the network to learn residuals directly rather than the difference between the original input and output. In this way, the problems of gradient disappearance and explosion can be effectively alleviated, and the training efficiency and generalization ability of the network can be improved [32].

2.5. The acquisition of 2DCOS image dataset

The generalized 2DCOS algorithm can obtain and analyze a series of dynamic spectra by applying external interference to the sample and finally carry out correlation analysis to obtain the 2DCOS image. The argument changes from continuous time to v. t is the external disturbance, and m is the spectrum measured by m steps at the same disturbance time interval t, as shown in Eq. (1).

$$\mathbf{y}(v) = \begin{vmatrix} \mathbf{y}(v, t_1) \\ \mathbf{y}(v, t_2) \\ \mathbf{y}(v, t_3) \\ \vdots \\ \vdots \\ \mathbf{y}(v, t_4) \end{vmatrix}$$
(1)

The intensity of synchronous two-dimensional correlation between v_1 and v_2 is expressed as formula (Eq. 2).

$$\phi(v_1, v_2) = 1/m - 1.\left(y(v_1)^T, y(v_2)\right)$$
(2)

The strength of the asynchronous two-dimensional correlation between v_1 and v_2 is expressed as formula (Eq. 3).

$$\psi(\boldsymbol{v}_1, \boldsymbol{v}_2) = 1 / m - 1.\left(\boldsymbol{y}(\boldsymbol{v}_1)^T \cdot \boldsymbol{N}_{jk} \cdot \boldsymbol{y} \begin{pmatrix} \boldsymbol{y}_2 \\ \boldsymbol{y}_2 \end{pmatrix} \right)$$
(3)

In formula (Eq. 4), where *N* is the Hilbert-Noda matrix.

$$N_{jk} = \begin{cases} 0, j = k\\ 1/m - 1, j \neq k \end{cases}$$

$$\tag{4}$$

The comprehensive two-dimensional correlation spectrum between ν_1 and ν_2 is defined as formula (Eq. 5).

$$I(v_1, v_2) = \phi(v_1, v_2) \cdot \psi(v_1, v_2)$$
(5)

2.6. Model performance evaluation parameters

17

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In the model performance evaluation, the evaluation indicators such as RMSEE, RMSECV, RMSEP and accuracy (Acc) are selected, and their respective calculation methods are as the following formula (Eqs. 6-9).

$$RMSEE = \sqrt{\frac{\sum_{i=1}^{m} (y_{i} \cdot \hat{y}_{i})^{2}}{m - N}}$$
(6)

$$RMSECV = \sqrt{\frac{1}{m-1} \sum_{i=1}^{m} (\hat{y}_{i} - y_{i})^{2}}$$
(7)

$$RMSEP = \sqrt{\frac{\sum_{i=1}^{m} (y_{i-} \hat{y}_{i})^{2}}{m}}$$
(8)

$$Acc = \frac{TP + TN}{TP + TN + FP + FN}$$
(9)

 \mathbf{y}_i is the measured value, $\hat{\mathbf{y}}_i$ is the predicted value, m is the sample size, and N is the number of variables considered in the model.

TP: Number of true positive TN: Number of true negatives

FP: Number of false positives

FN: Number of false negative

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3. Results and Discussion

3.1. ATR-FTIR analysis

The original and average mid-infrared spectra of GI and non-GI *G. elata* in Yunnan and Guizhou have been presented in Figure 2. Although artificially cultivated *G. elata* comes from different production areas in Yunnan and Guizhou, the spectra of different regions usually show similar trends. However, absorption intensity presents certain differences, which suggests that *G. elata* may differ slightly in the chemical composition of different regions.

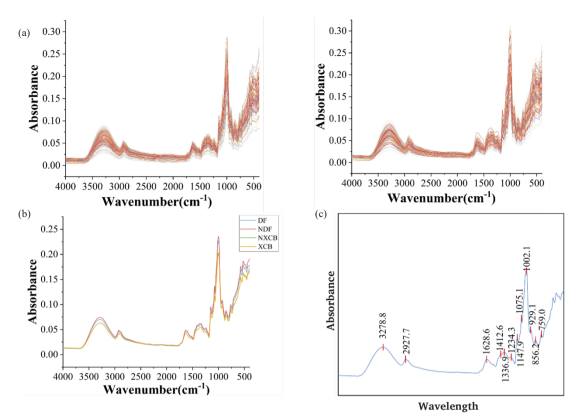


Figure 2. ATR-FTIR spectrum. (a) The original spectra (Left: Yunnan, Right: Guizhou). (b, c) Average spectrogram.

The strong broad band centered on 3278.8 cm⁻¹ may correspond to hydroxyl (-OH) stretching vibrations of phenolic compounds in G. elata, including hydroxyl absorption in phenolic compounds [33]. The presence of 2927.7 cm⁻¹ may be the symmetric and asymmetric stretching vibrations of C-H in aliphatic hydrocarbons in terpenoids [34]. Phenolic compounds in G. elata or other compounds containing carbonyl (C=O), including carbonyl absorption in phenolic compounds, are likely to be present at 1628.6 cm⁻¹, and 1412.6 cm⁻¹ usually corresponds to the methyl (CH₃) functional group, which indicates compounds containing methyl functional groups in G. elata. It is possible that C=O, thioether bond (S-S, S-C, and C-S) correspond to 1336.9, 1234.3, 1147.9, and 929.1 cm⁻¹, respectively [35,36]. What's more, 1075.1 cm⁻¹ may demonstrate compounds that contain hydroxyl functional groups in G. elata. The peaks corresponding to 856.2 cm⁻¹ and 759 cm⁻¹ are associated with C-H bending vibrations in aromatic compounds. Due to the very similar metabolic composition of G. elata, it is challenging to distinguish the original spectrum, and more sensitive machine-learning techniques must be employed to identify GI and non-GI G. elata.

3.2. Exploratory PCA analysis

In order to explore the similarities and differences between GIproducing areas and non-GI-producing areas, the principal component analysis was conducted to reduce the infrared spectral data of GI and non-GI G. elata. The results of PCA (Figure 3) showed the distribution of samples from Yunnan and Guizhou provinces in the principal component space, which facilitates the realization of GI and non-GI G. elata identification. PC1 and PC2, the first two principal components of the PCA score plot (Figure 3a), accounted for 88.4% and 9.0% of the total variance, respectively. Taken together, there is a high cumulative variance contribution of these two principal components, indicating that they contain most of the data variability. However, for the original ATR-FTIR of the samples, there was a classification overlap of the two types of samples in the score map, meaning that less difference existed in chemical composition between the GI and non-GI regions within the space constructed by the first two principal components. Figure 3(b) depicts the PCA results of different regions in Guizhou. The first two principal components account for over 90.0% of the variance explanation rate, with PC1 contributing 90.8% and PC2 contributing 6.7%, which implies that these two principal components not only more effectively explain the variance of the original data but also possess strong information preservation ability.

Although PCA can effectively explain the variance of the original data, there remains an overlap in the classification of GI and non-GI *G. elata* in Yunnan and Guizhou. Certain GI areas and non-GI areas probably exhibit similarities or differences. Due to the high similarity of data features and the uneven distribution of data, overlap of GIS and non-GI *G. elata* samples in principal component space in different provinces may be caused. It was identified that PCA does not well describe the differences between samples, which is consistent with geo-traceable studies of porcini, green tea, and canola oil [37-39]. Hence, the establishment of supervised machine learning models plays a necessary role in attaining more accurate traceability, including PLS-DA, SVM, and ResNet.

3.3. Chemometrics

3.3.1. PLS-DA results

Spectral preprocessing aims to improve data quality, highlight useful information, and eliminate interference factors, which can provide a reliable basis for subsequent data analysis and model. As can be seen from Table 2, the spectral data of G. elata in Yunnan Province obtained poor results without preprocessing, which still has room for improvement. After employing single or combined spectral preprocessing, improvement was witnessed in PLS-DA model R² and Q² compared with the original spectra, while a slight decrease was seen in the values of RMSEE, RMSECV, and RMSEP. SD is often used in spectral preprocessing to enhance peak value, improve signal-tonoise ratio, and correct baseline drift. The results of the preprocessing methods in Table 2 demonstrated that when harnessing SD alone, its R² and Q² values were the highest compared with the other four spectral preprocessing methods, implying that SD is a more suitable spectral preprocessing method for Yunnan G. elata in specific datasets and models. After SG smoothing pretreatment, the equivalent values of R², Q², RMSEE, RMSECV, and RMSEP were close to those of the

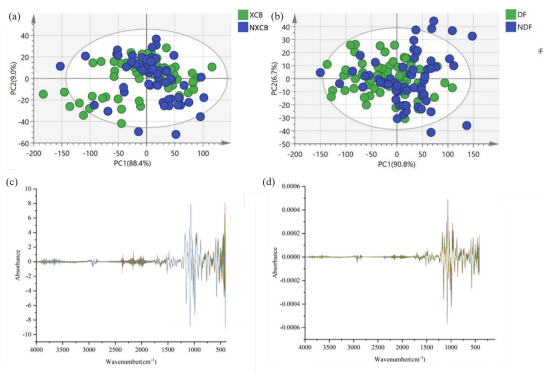


Figure 3. The establishment of PCA model to analyze the PCA results of geographical indication production areas and non-geographical indication production areas. (a) Yunnan (XCB, NXCB); (b) Guizhou (DF, NDF). Best Spectral Preprocessing Image. (c) Yunnan SNV + SD; (d) Guizhou SG + SD.

original spectrum. SG smoothing pretreatment lacks in improving the stability and reliability of modeling, though it aids in reducing noise and fluctuations in the spectrum. There is the largest R^2 (0.903) and Q^2 (0.653) in SNV combined with SD. SD can excel at the subtle features and changes in the spectral data, while SNV can eliminate the scale effect in the data, which enables the data to be more consistent with the normal distribution and further verifies that the combined spectral pretreatment can better enhance the model performance (Figure 3c). In summary, both single and combined spectral preprocessing exhibited improved PLS-DA prediction ability, albeit to different degrees of improvement.

Table 3 shows the spectral preprocessing results of GI and non-GI *G. elata* in Guizhou. The results obtained by a single method are consistent with those in Yunnan, in which excellent findings are presented by SD. The spectral pretreatment of MSC+SNV, MSC+SG, and SNV+SG indicates close equivalent values of R². Q², RMSEE, RMSECV, and RMSEP. The results imply that SG+SD serves as the best spectral preprocessing method to obtain the maximum accuracy of the training

Table 2. Parameters of PLS-DA model based on ATR-FTIR (Yunnan).

Pretreatment	R ²	Q ²	RMSEE	RMSECV	RMSEP	Train set Acc (%)	Test set Acc (%)
Raw	0.0473	0.0178	0.494	0.4955	0.5072	60.24	61.11
MSC	0.521	0.278	0.3568	0.445	0.487	87.95	58.33
SNV	0.522	0.277	0.3565	0.447	0.4876	89.16	55.56
SG	0.145	0.06	0.4940	0.4955	0.5072	60.24	61.11
SD	0.857	0.609	0.1941	0.3122	0.3291	94.96	83.33
MSC+SNV	0.522	0.278	0.3567	0.4453	0.4871	87.95	58.33
MSC+SG	0.589	0.252	0.3329	0.4514	0.5058	91.57	61.11
SNV+SG	0.521	0.277	0.3568	0.4462	0.4896	86.75	55.56
MSC+SD	0.806	0.571	0.2242	0.3316	0.3999	97.59	75.00
SNV+SD	0.903	0.653	0.1595	0.3058	0.3205	100.00	88.89
SG+SD	0.854	0.603	0.1957	0.3149	0.3278	100.00	83.33

Note: R²: Explains the ability of the model to fit the data; Q²: Indicates the prediction ability of the model to new data; RMSEE: Root mean square error of estimation; RMSECV: Root mean square error of cross-validation; RMSEP: Root mean square error of prediction; Acc: Accuracy. The bold text indicates the optimal preprocessing method.

set and the test set (Figure 3d). Consequently, suitable pretreatment methods should be selected according to their characteristics and data performance for different spectral data sets, which can therefore improve data quality, underline sample characteristics, and provide a reliable basis for subsequent analysis and model. Hence, it is essential to flexibly select the best pretreatment when processing spectral data. The results imply that the accuracy of both of them in the training set is 100% (Table 2, Table 3), whereas the accuracy of Yunnan in the test set is 88.89% (Table 2), which is lower than the 94.74% of the Guizhou test set (Table 3), and the confusion matrix is shown in Figure 4. Some differences are likely to be exhibited by the sample data of Yunnan Province and Guizhou Province, including the growth environment, climate, soil, and other factors, potentially resulting in some differences in the spectral data of G. elata in different regions, which enables the established model to perform differently on the test set in different regions.

On the basis of the best pretreatment spectral data, the PLS-DA model was established, and a good classification was demonstrated by G. elata in GI-producing areas and non-GI-producing areas in Yunnan Province (Figure 5a). The cumulative variance explanation rate of the first two principal components in PLS-DA score chart accounted for 17.5% of the total variance, of which PC1 explained 13.8% and PC2 explained 3.7%, respectively. The cumulative variance explanation rate is an important indicator to evaluate the degree of variation of the model's interpretation data, in which the higher value indicates the stronger interpretation ability of the model to the data. Under such circumstances, the explanatory power of PC1 is significantly higher than that of PC2 despite the relatively low cumulative variance interpretation rate, indicating that PC1 plays a more significant role in distinguishing samples. However, the PLS-DA scoring plot (Figure 5b) clearly reveals the distribution differences of G. elata samples in the principal component space of DF and non-GI regions. PC1 and PC2 explained 20.5% and 6.2%, respectively, of the variables in the spectral data, which may also mean that it is difficult to distinguish G. elata samples from GI and non-GI regions of these two principal components.

Regarding *G. elata* from GI and non-GI production areas in Yunnan Province and *G. elata* from Guizhou Province, the importance of substitution is leveraged to evaluate the fitting degree of the model. The phenomenon that the corresponding model training set has better

Pretreatment	R ²	Q ²	RMSEE	RMSECV	RMSEP	Train set Acc (%)	Test set Acc (%)
Raw	0.46	0.304	0.3775	0.45	0.331	81.32	86.84
SD	0.949	0.837	0.1165	0.2171	0.2442	100.00	94.74
MSC	0.444	0.32	0.3842	0.4125	0.3357	78.02	86.84
SNV	0.307	0.245	0.4203	0.4339	0.3981	79.12	84.21
SG	0.485	0.388	0.4751	0.4779	0.4655	67.03	57.89
MSC+SNV	0.427	0.32	0.3842	0.4125	0.3357	78.02	86.84
MSC+SD	0.888	0.694	0.1702	0.2779	0.2589	98.9	94.74
MSC+SG	0.428	0.321	0.3839	0.4121	0.3357	78.02	86.84
SNV+SD	0.96	0.736	0.1033	0.2661	0.2361	100.00	94.74
SNV+SG	0.428	0.321	0.3839	0.4122	0.3351	76.92	86.84
SG+SD	0.945	0.828	0.12	0.222	0.2469	100.00	94.74

The bold text indicates the optimal preprocessing method.

predictive performance and poorer validation for unknown external samples is explained. For the PLS-DA model, the Q^2 and R^2 values generated by each iteration are lower than the Q^2 and R^2 values in the upper right corner. All the above results demonstrate that the model is not overfitted for each category of discrimination (Figure 5c and 5d).

3.3.2. Analysis of DD-SIMCA results

To improve the prediction accuracy of the model, DD-SIMCA attempted to directly distinguish between geographical indications and non-geographical indications in Yunnan Province and Guizhou Province and investigate the impact of different spectral data processing techniques on the performance of the model. The DD-SIMCA model is a modeling method that can be used to analyse and identify anomalies, which primarily focuses on anomaly detection and classification in dynamic data processes. The acceptance area (the area within the green curve), outlier limitation (the red curve), rule samples (the green dots), extremum (the yellow square), and outliers (the red square) for the samples used in model construction have been provided in the acceptance graph of TrainSet. In Figure 6, sub-figures Figure 6(a) and 6(c), it can be seen that under the best preprocessing method,

the *G. elata* spectral data from Yunnan and Guizhou possess 1 and 2 samples located in the "extreme region" (yellow dots), respectively, and no samples located in the "abnormal region" (above the red threshold). When ATR-FIR is leveraged for data, all models cannot achieve 100% sensitivity regardless of preprocessing techniques, revealing that non-geographical indications *G. elata* in Yunnan and Guizhou cannot be correctly identified. However, the highest number of correctly classified non-GI *G. elata* using SD was 33 for *G. elata* from Yunnan, in which 27 out of 60 samples were projected to the geographical indication *G. elata* area. Consequently, the SD/D-SIMCA model obtained the best results, with a sensitivity of 96.67% and specificity of 55.00% for non-target classes (Figure 6b). By using the SG/D-SIMCA model, 52.46% of Guizhou's non-geographical indication *G. elata* and the sensitivity was 93.44% for non-target categories (Figure 6d).

In summary, the model cannot identify all non-geographical indication G. elata samples from Yunnan and Guizhou as non-target classes, which reveals that it cannot be harnessed to reject nongeographical indication samples. The possible reasons for this are as follows. First of all, the DD-SIMCA model relies on the characteristics of the input data. The model may not be able to effectively distinguish them if the geographical indication production areas and nongeographical indication production areas possess a high similarity or overlap in data features, including close distances between production areas. Secondly, the DD-SIMCA model requires a large amount of data for effective training and modeling. In case of insufficient sample data ON Yunnan and Guizhou, the model may not be able to capture the small differences between them. Being a linear model-based method that may have limitations in handling nonlinear features, DD-SIMCA can cause lower classification performance compared to PLS-DA and SVM models. SEN represents the model's ability to correctly classify target classes (regular samples), while SPE serves as the model's ability to correctly classify non-target classes (foreign samples).

3.3.3. SVM analysis

SVM refers to a classical model commonly utilized to solve nonlinear problems. As a two-classification model, its basic model is a linear classifier that establishes the maximum interval in the feature space to

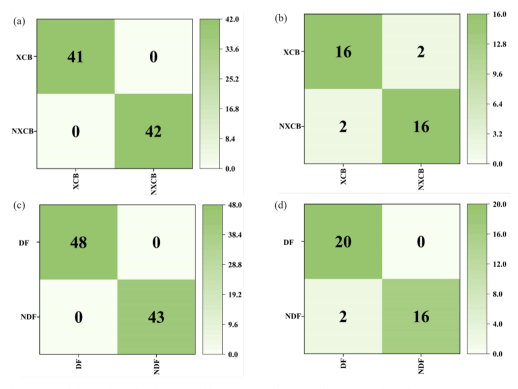


Figure 4. Confusion matrix results of PLS-DA model training set and test set. (a, b) Yunnan; (c, d) Guizhou.

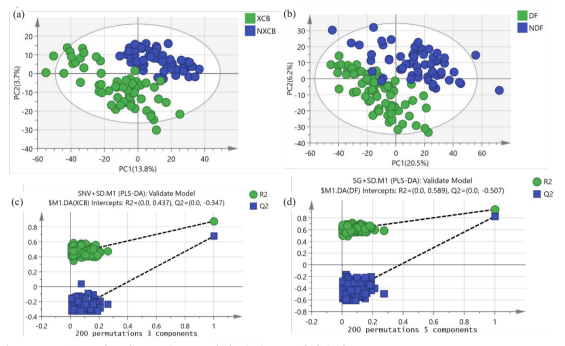


Figure 5. PLS-DA score plot and permutation test result plot. (a, c) Yunnan; (b, d) Guizhou.

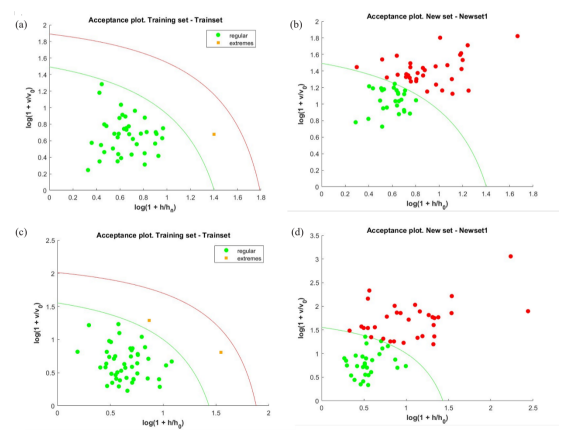


Figure 6. The DD-SIMCA results. Acceptance maps for training sets (a) Yunnan and (c) Guizhou; (b) Yunnan and (d) Guizhou are the predicted results of the test set.

reduce overfitting. The larger the *g* is, the higher the mapping dimension is, the better the training result is, whereas the more likely it is to cause overfit. In Table 4, the optimization results of the important parameters and the accuracy of the training and test sets have been shown. Prior to building the SVM model, the raw spectral data were preprocessed in 10 different ways. The optimal values of penalty factor *c* and kernel

parameter *g* were obtained by the grid search method, and the SVM model was also constructed. The model results have been shown in Table 4. However, it is essential to specify the reasonable range of the two parameters of SVM according to the situation since the places of *G. elata* exhibit some differences. After undergoing MSC+SD and SNV+SD spectral preprocessing, the original spectra of *G. elata* from Yunnan

Table 4. SVM modeling results based on different preprocessing methods (Yunnan).

Data set	Best c	Best g	Training set Acc (%)	Test set Acc (%)
Raw	524288	1.079×10^{-5}	91.57	69.44
MSC	32768	1.079×10 ⁻⁵	91.57	75.00
SG	262144	2.1579×10 ⁻⁵	91.57	69.44
SNV	23170.475	1.5259×10 ⁻⁵	91.57	75.00
MSC+SG	46340.95	7.6294×10-6	91.57	75.00
MSC+SNV	32768	1.079×10 ⁻⁵	91.57	75.00
SNV+SG	32768	1.079×10 ⁻⁵	91.57	75.00
SD	45.2548	0.00024414	92.77	88.89
MSC+SD	2	0.0039062	93.98	91.67
SG+SD	5.6569	0.0013811	90.36	88.89
SNV+SD	2	0.0039062	93.98	91.67

Best c

(Guizhou).

Raw	262144	2.1579×10^{-5}	93.41	94.74
SD	5.6569	0.0013811	97.80	94.74
MSC	8192	$4.3158 imes 10^{-5}$	94.51	92.11
SNV	16384	$2.1579 imes 10^{-5}$	94.51	92.11
SG	741455.2002	1.079×10^{-5}	92.31	94.74
MSC+SNV	5.6569	0.0013811	100.00	94.74
MSC+SG	23170.475	$1.5259 imes 10^{-5}$	94.51	89.47
MSC+SD	5.6569	0.0013811	100.00	94.74
SNV+SD	16	0.00097656	98.90	52.63
SNV+SG	32768	1.079×10^{-5}	94.51	89.47
SG+SD	5.6569	0.0013811	97.80	94.74

Table 5. SVM modeling results based on different preprocessing methods

showed excellent performance, yielding consistent c and g values of 2 and 0.0039062, respectively (Figure 7a), which indicates that the parameter adjustment effect of the model is good and can better fit the data. At the same time, the accuracy of the training set reached 93.98% (Figure 7b), and the results of the confusion matrix demonstrated classification errors (Figure 7c). Nevertheless, the accuracy of the test set reached 91.67%, which implies that the model not only possesses good generalization capability to a certain extent but also achieves a relatively considerable accuracy on the test data.

Being a supervised learning algorithm with classification and regression capabilities, SVM constructs hyperplanes in K-dimensional feature spaces to maximize the spacing between adjacent classes [10]. After the model parameters are adjusted, it can better align with the data characteristics, and under the best spectral preprocessing, the best c and g obtained are 5.6569 and 0.0013811, respectively (Figure 7d). In the ATR-FTIR model, the accuracy of the training set in the SVM model was as high as 100.00% (Table 5), and the accuracy of the test set was 94.74% (Figure 7e). The two type 2 samples in the test set were incorrectly set up (Figure 7f). Generally speaking, the SVM model both presents robust explanatory power and is capable of effectively distinguishing G. elata samples from GI-producing areas and non-GIproducing areas in Yunnan and Guizhou.

3.4. Deep learning

3.4.1. The 2DCOS spectra dataset

Two-dimensional correlation infrared (2D-IR) spectroscopy was first developed by Dr.IsaoNoda and subsequently extended to other analytical methods, and the concept of generalized 2DCOS was further proposed. To identify G. elata samples from different producing areas in Yunnan and Guizhou, this research compared three kinds of 2DCOS images, namely synchronous, asynchronous, and integrated 2DCOS (i2DCOS). By mapping one-dimensional (1D) information to two-dimensional space, 2DCOS improves spectral resolution and effectively solves the problem of overlapping absorption peaks of the same functional group in one-dimensional spectra. The mathematical methods were conducted to obtain the synchronous, asynchronous, and i2DCOS images (128×128 pixels) of G. elata in different production areas. Figure 8 shows the representative synchronous (Figure 8a(A1-A6)), asynchronous (Figure 8b(B1-B6)), and i2DCOS (Figure 8c(C1-C6)) of the six producing areas. In 2DCOS, the characteristic peaks are primarily divided into automatic peaks and cross peaks. Specifically, automatic peaks usually appear on the diagonal of the spectrum and, therefore, are also called diagonal peaks. In the area of the automatic peak in the spectrum, the degree of spectral intensity change under this wave number is reflected. The automatic peak is always positive, which means that the trend of increasing or decreasing spectral intensity is consistent at that wave number. There are usually cross peaks on both sides of the diagonal, and both positive and negative peaks appear simultaneously, indicating the type of autocorrelation (positive or negative correlation) of the spectral intensity change. Nevertheless, it may be subjective to identify G. elata from different origins in Yunnan and Guizhou provinces by two-dimensional spectrograms alone, as spectrograms can only provide

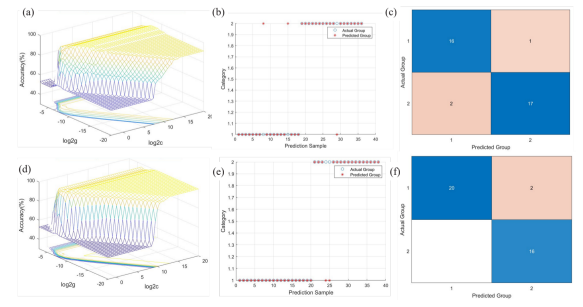


Figure 7. SVM results. SVM parameter selection results, test set accuracy, and confusion matrices. (a-c) Yunnan Province; (d-f) Guizhou Province.

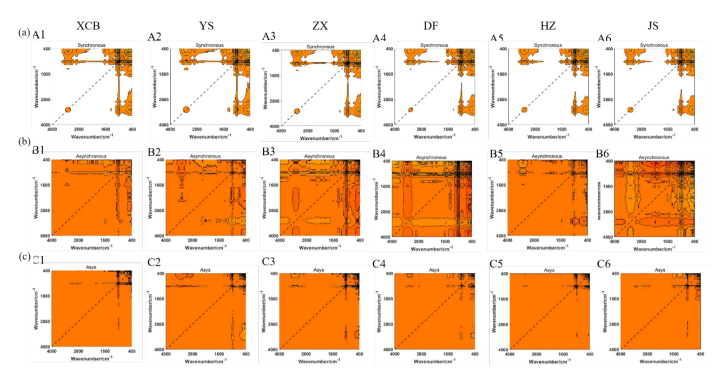


Figure 8. (a) Synchronous (A1-A6), (b) Asynchronous (B1-B6), and (c) Integrated (C1-C6) two-dimensional correlation spectra.

limited information, and deep learning techniques like ResNet can be used to more accurately identify the origin.

3.4.2. ResNet model analysis

The RseNet model, based on synchronous, asynchronous, and i2DCOS, was established since it is inaccurate to distinguish between GI production areas and non-GI production areas of G. elata solely based on the difference in 2DCOS recognized by the naked eye. Under the same number of iterations (200 epochs) and learning rate (0.01), Figure 9(a-c) reveals the accuracy curves of the training set and the test set of the three ResNet models with 2DCOS, with significant differences in the cross-entropy loss function curve and the confusion matrix of the external verification set. An accuracy curve is utilized to assess the model's recognition ability, and a cross-entropy loss function is employed to demonstrate the model's convergence effect. The closer the loss value is to 0, the closer the accuracy is to 100.00%, and the stronger the recognition ability of the model is. In Table 6, the detailed parameters of the ResNet model (number of iterations, loss value, accuracy of training set, test set and external validation set) were summarized. After 200 epochs, the accuracy of the ResNet model based on the 2DCOS spectral images of the training set and the test set was 100.00%, the loss value converged to 0.019, and the accuracy of the external validation set reached 100.00%. It is speculated that the ResNet model based on synchronous 2DCOS images not only extracted the most feature variables but also presented the strongest recognition ability, which holds the potential to quickly trace the geographical origin of G. elata. As the number of iterations increases, the accuracy of the training set of the ResNet model for synchronizing and i2DCOS images established under the same number of iterations can reach 100%, with different loss values and accuracy of the test set, which are 0.769, 25.00% and 0.272, 23.00%, respectively. In the results, there was the poorest external validation accuracy of ResNet based on i2DCOS, with DF G. elata misclassified to YS and JS G. elata not accurately identified. The model is unable to accurately distinguish G. elata from different places since the i2DCOS images may not fully extract the key differences in the appearance characteristics of DF, YS, and JS G. elata.

To sum up, the ResNet model training set of synchronous 2DCOS images attained 100% accuracy. It is hypothesized that asynchronous

Table 6. Parameters of ResNet model based on 2DCOS
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Data (2DCOS)	Epoch	Loss value	Train set Acc (%)	Test set Acc (%)	External validation Acc (%)
Synchronous	200	0.019	100.00	100.00	100.00
Asynchronous	200	0.769	98.00	25.00	100.00
Asys	200	0.272	100.00	23.00	87.50

and i2DCOS images may have more noise, changes, and complex features and, therefore, make the model more difficult to learn and recognize. In contrast, synchronous 2DCOS images are likely to have more regular and simple features, which are relatively clear and stable, and the model can more easily understand and capture key feature information, thus achieving better accuracy on the training set.

4. Conclusions

In this study, the application effect of several models in the identification of G. elata GIS and non-GIS in Yunnan and Guizhou was systematically evaluated. ATR-FTIR has the advantage of nondestructive testing to realize rapid sample analysis while requiring high stability of results. DD-SIMCA performs well in category recognition but is prone to recognition errors. In addition, PLS-DA shows strong capability in processing high-dimensional data, whereas overfitting should be prevented. Relatively speaking, SVM performs well in small samples and high-dimensional space and can effectively tackle sample imbalance and identify G. elata geographical indication and non-geographical indication production areas. Regarding the proposed methods, the synchronous 2DCOS spectral ResNet model exhibited obvious advantages in the identification of G. elata from 6 production areas in Yunnan and Guizhou, both outperforming other models and achieving 100% classification accuracy, which can effectively capture the small differences between samples and reduce the loss value and is suitable for application in a complex background. In this study, the combination of chemometrics and deep learning can aid in further improving the accuracy and stability of geographical indication identification, and the results can generally provide crucial guidance

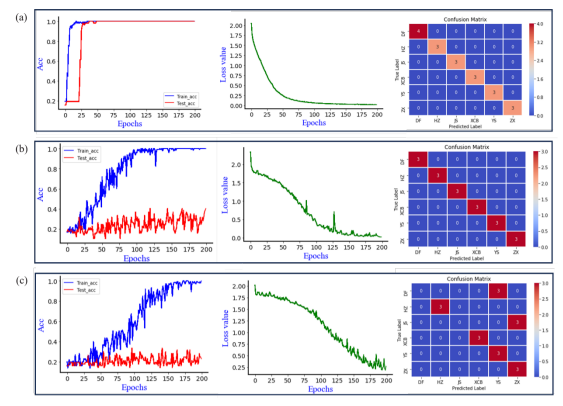


Figure 9. The accuracy curves of the training and testing sets of the ResNet model, as well as the cross entropyloss function curve and the confusion matrix of the external validation set. (a) Synchronize two-dimensional correlation spectra; (b) Asynchronous two-dimensional correlation spectroscopy; (c) Integrated two-dimensional correlation spectroscopy.

and reference for the future application of non-destructive testing of medicinal and edible homologous plants.

CRediT authorship contribution statement

Qiong He: Conceptualization, Formal analysis, Writing-original draft, Writing-review & editing. Hengyu Huang: Investigation, Resources. Yuanzhong Wang: Funding acquisition, Investigation, Resources, Supervision.

Declaration of competing interest

The authors declare that they have no known competing interest or personal relationship that could have appeared to influence the work reported in this paper.

Data availability

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

Declaration of Generative AI and AI-assisted technologies in the writing process

The authors confirm that there was no use of artificial intelligence (AI)-assisted technology for assisting in the writing or editing of the manuscript and no images were manipulated using AI.

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