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Analyzing Cinnamon Spice Adulteration with Spectroscopy: The Influence of **Data Preprocessing on Multivariate Prediction Models**

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Abstract

Detecting fraud in the cinnamon supply chain is critical for ensuring consumer safety and maintaining product integrity. Recent advances in spectral data preprocessing techniques offer enhanced accuracy in identifying adulterants in spices like cinnamon. This study investigates the impact of different spectral preprocessing techniques on predicting adulterants—specifically soybean powder, hazelnut shell powder, and dry bread powder—mixed with cinnamon powder using spectroscopy combined with multivariate analysis. The transmittance spectra were collected across the mid-infrared range of 2-4000 cm⁻¹, and Partial Least Squares Regression (PLSR) was employed to model the adulteration levels based on these spectra. Various preprocessing methods were applied to optimize the spectral data. Among them, orthogonal signal correction (OSC) combined with detrending yielded the highest predictive accuracy, with a coefficient of prediction (R2p) ranging from 0.900 to 0.981. Conversely, Extended Multiplicative Scatter Correction (EMSC) and Savitzky-Golay second derivative (D²) were less effective, with R²p values between 0.115 and 0.931. Soybean powder was the easiest adulterant to detect, with a prediction error range of 5–10%. These findings underscore the importance of selecting appropriate preprocessing techniques to improve the accuracy of fraud detection in cinnamon powder using spectroscopic methods.

Keywords: Adulterants, Chemometrics, Data Preprocessing, Food Safety, Spectral Analysis.

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1. Introduction

Food fraud, particularly in the spice industry, has emerged as a significant global concern, jeopardizing consumer health and undermining market integrity [1,2]. Spices, including cinnamon, are highly valued not only for their flavor and aroma but also for their numerous health benefits [3,4]. The high market value of authentic spices makes them susceptible to adulteration, where cheaper substances are added to enhance weight or appearance [5,6]. This practice can mislead consumers, posing potential health risks due to the introduction of harmful additives or low-quality materials [7,8]. Furthermore, food fraud can severely impact the economy, as it erodes consumer trust and diminishes the reputation of legitimate producers. As the complexity and prevalence of food fraud continue to rise, effective detection methods become imperative to protect consumer interests and ensure food quality.

Fourier-transform infrared (FTIR) spectroscopy has gained prominence as a powerful tool for detecting food adulteration, thanks to its ability to provide detailed molecular information about food samples [9–11]. FTIR spectroscopy is based on the principle of measuring the absorption of infrared light by a sample, producing a unique spectral fingerprint that corresponds to its molecular composition [12,13]. This technique offers several advantages, including non-destructive analysis, rapid data acquisition, and minimal sample preparation requirements [14]. Additionally, FTIR can analyze solid, liquid, and gas samples across a broad range of wavelengths, making it highly versatile for various applications in food science [15–17]. Its ability to identify specific functional groups in chemical compounds enables researchers to differentiate between pure and adulterated products, making FTIR a valuable method for quality control in the food industry [18].

In recent years, the application of FTIR spectroscopy in food authentication has expanded significantly, particularly in the detection of adulterants in spices [19–21]. Numerous studies have demonstrated its effectiveness in identifying various adulterants in spice products, providing a rapid and reliable means of ensuring product integrity [22–24]. For instance, FTIR has been successfully employed to differentiate between pure spices and those mixed with cheaper fillers or artificial additives, thus safeguarding consumer interests. The integration of FTIR with advanced data analysis techniques, such as multivariate analysis, enhances its capability to classify samples based on spectral data [25–27]. By applying sophisticated algorithms to process the spectral information, researchers can improve the accuracy of fraud detection, enabling the identification of specific adulterants and their concentrations in spice products [28].

Cinnamon, a popular spice derived from the inner bark of Cinnamonum trees, is not only valued for its culinary uses but also for its health-promoting properties, including antioxidant and anti-inflammatory effects [29]. Unfortunately, the rising demand for cinnamon has led to increased incidents of adulteration, where lower-quality cinnamon varieties, such as Cassia, are often substituted for authentic Ceylon cinnamon [30]. This fraudulent practice can mislead consumers and diminish the health benefits associated with genuine cinnamon. The complexity of cinnamon's chemical composition and the presence of similar aromatic compounds in adulterants complicate detection efforts. Therefore, it is crucial to develop robust analytical methods that can effectively differentiate between pure and adulterated cinnamon, ensuring that consumers receive authentic products and promoting confidence in the spice market [31].

This study aims to investigate the influence of data preprocessing techniques on the predictive accuracy of multivariate models for detecting cinnamon adulteration using FTIR spectroscopy. By employing various preprocessing methods, the research seeks to enhance the reliability of

spectral data analysis, enabling more accurate identification of adulterants such as soybean powder, hazelnut shell powder, and dried bread powder in cinnamon products. This research presents a comprehensive evaluation of 14 data preparation methodologies employed in FTIR spectral analysis for the identification of adulteration in cinnamon powder. Furthermore, the inclusion of three adulterants (soybean powder, hazelnut shell powder, and dry bread powder) contributes to the broader generalizability of the results. based on authors' knowledge, these specific adulterants have very limitted previously investigated in the context of spice adulteration specially in cinnamon powder, making this research a novel contribution to the field. The findings of this study will contribute to improving food quality assurance practices, providing insights into the effectiveness of FTIR spectroscopy in combating food fraud, and ultimately supporting consumer health and safety.



Fig 1: Prepared samples, Ci (cinnamon powder), Ds (soybean powder), Hn (hazelnut shell powder), Db (dry bread powder).

2. Materials and methods

2.1. Collection and preparation of samples

In this study, various ingredients were acquired from local markets in Mashhad, Iran, specifically focusing on their authenticity and quality for food analysis. The primary ingredient used was Cinnamomum verum, commonly known as cinnamon, which was selected for its exceptional quality. The preparation involved a meticulous cleaning process to eliminate contaminants, followed by an air-drying phase that preserved its natural characteristics. After drying, the cinnamon was finely processed using a high-precision grinder to obtain a standard cinnamon powder, which served as the baseline for comparison in our experiments.

To simulate common adulteration practices, we also utilized several potential adulterants, including roasted soybean grains, cleaned and ground hazelnut shells, and powdered dried bread. Each adulterant was processed according to established methods documented in previous studies, including those by Hashemi-Nasab et al. (2023) and Khodabakhshian et al. (2021), with necessary adaptations made to suit the parameters of our investigation [32, 33]. The soybean was lightly toasted before grinding to reflect typical processing conditions, while the hazelnut shells were dried and pulverized into a consistent powder form. The dried bread was ground and sieved to achieve a uniform texture compatible with the other ingredients.

All the selected samples underwent a standard grinding procedure to ensure uniform particle size. Following this, they were passed through a mesh sieve to maintain consistent particle

distribution, which is critical for accurate analysis. To create mixed samples, the cinnamon powder was combined with varying proportions of the adulterants—5%, 10%, and 15% by weight. Figure 1 presents the various prepared mixtures. The composition of the sample groups, including pure and adulterated powders, along with the number of replicates, is summarized in Table 1. This blending was executed thoroughly to achieve a homogeneous mixture, which is representative of actual adulteration practices. Each adulterated mixture was prepared in triplicate, including controls of pure cinnamon powder. The final samples were securely packaged in airtight bags to shield them from external factors like moisture and light. They were stored in a controlled environment to preserve their integrity until the spectroscopic evaluations were conducted.

Table 1. Composition of sample groups and number of replicates.

Sample NO.	Description	Adulterant Level (% w/w)	No. of Unique Samples	Replicates per Sample	Weight of per sample	Total Measurements
1	Pure cinnamon powder (Ci)	0%	1	5	10g	5
2	Pure soybean powder (Ds)	100%	1	5	10g	5
3	Pure hazelnut shell powder (Hn)	100%	1	5	10g	5
4	Pure dried bread powder (Db)	100%	1	5	10g	5
5	Cinnamon + Soybean powder (CiDs)	5%, 10%, 15%	3	5	10g	15
6	Cinnamon + Hazelnut shell powder (CiHn)	5%, 10%, 15%	3	5	10g	15
8	Cinnamon + Dried bread powder (CiDb)	5%, 10%, 15%	3	5	10g	15
Total			13			65

2.2. Spectra data acquisition

In this study, an Avatar 370 FTIR spectrometer (Thermo Nicolet Corporation, Madison, WI, USA) equipped with a deuterated triglycine sulfate (DTGS) detector and a KBr beamsplitter was employed to detect adulteration in cinnamon powder. The spectrometer operates in the midinfrared range ($4000-400~\rm cm^{-1}$) with a spectral resolution of 4 cm⁻¹ and a wavenumber accuracy of $\pm 0.01~\rm cm^{-1}$. The system includes an attenuated total reflectance (ATR) accessory with a diamond crystal, allowing for minimal sample preparation and direct analysis of powdered samples.

Each spectrum was obtained by averaging 32 scans per sample to enhance the signal-to-noise ratio (SNR). The instrumental setup and spectral acquisition process are illustrated in Figure 2.

To ensure data reproducibility, the spectrometer was calibrated weekly using a polystyrene standard, and background scans were performed before each measurement session.

For data acquisition and initial preprocessing, OMNIC spectroscopy software (Thermo Nicolet Corporation, Madison, WI, USA) was used. This software supports real-time spectral correction, baseline adjustment, and noise reduction. Further advanced preprocessing was conducted using Unscrambler X V10.4 (Camo Analytics, Oslo, Norway), as detailed in the following sections.

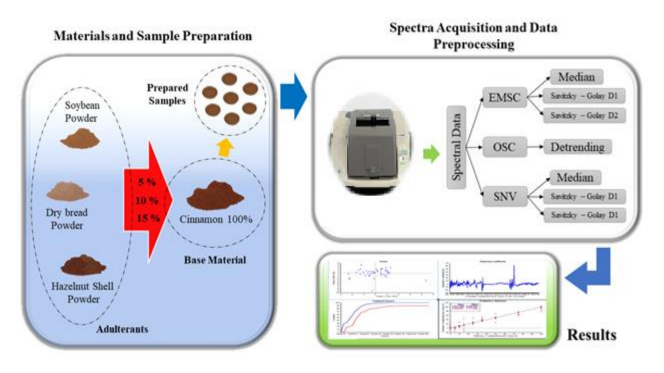
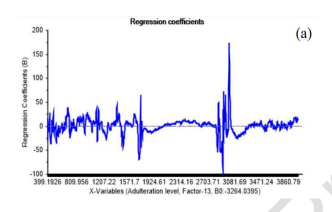


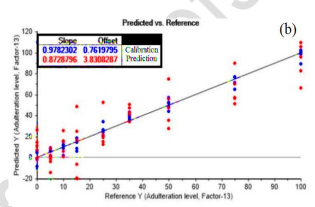
Fig 2: A schematic diagram of the proposed procedure.

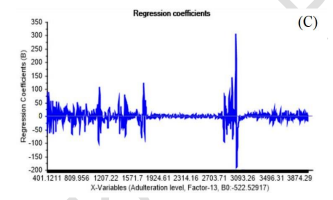
2.3. Spectral Preprocessing

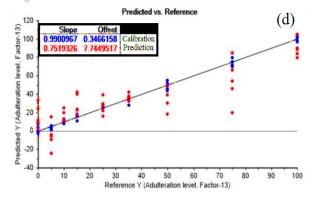
To ensure accurate multivariate modeling for detecting adulteration in cinnamon powder, several spectral preprocessing techniques were applied. These preprocessing steps were conducted using the Unscrambler X software (version 10.4, CAMO Software AS, Bedford, Massachusetts, United States). Initially, orthogonal signal correction (OSC) was used to remove variations unrelated to the cinnamon adulteration while preserving the primary information relevant to the spectral analysis. OSC is particularly effective in eliminating noise or irrelevant variability in spectral data, which could obscure the detection of adulterants such as dry bread powder. Following OSC, we implemented standard normal variate (SNV) transformation. SNV reduces scattering effects that may arise from inconsistencies in sample morphology or particle size, ensuring that the spectral data are more uniform and interpretable across different samples [37]. Additionally, extended multiplicative scatter correction (EMSC) was applied to further correct for scatter variations while incorporating adjustments for physical and chemical interactions affecting the absorbance spectra [38].

The preprocessing strategy also included smoothing techniques to reduce noise and enhance signal clarity. A median filter was first employed to remove outlier spikes, followed by Savitzky-Golay smoothing, which was applied in both the first (D¹) and second (D²) derivative forms to further improve peak resolution. The Savitzky-Golay algorithm, which fits successive polynomial functions to localized spectral segments, is particularly useful for reducing random noise while preserving important features such as peak intensity and width. By applying D¹ and D² derivatives, baseline shifts were effectively corrected, and overlapping spectral peaks were resolved, making it easier to distinguish between pure and adulterated cinnamon samples [34]. The combination of these preprocessing methods enabled the generation of a clean and reliable dataset, suitable for subsequent multivariate analysis, particularly in the context of the Partial Least Squares Regression (PLSR) model [39].









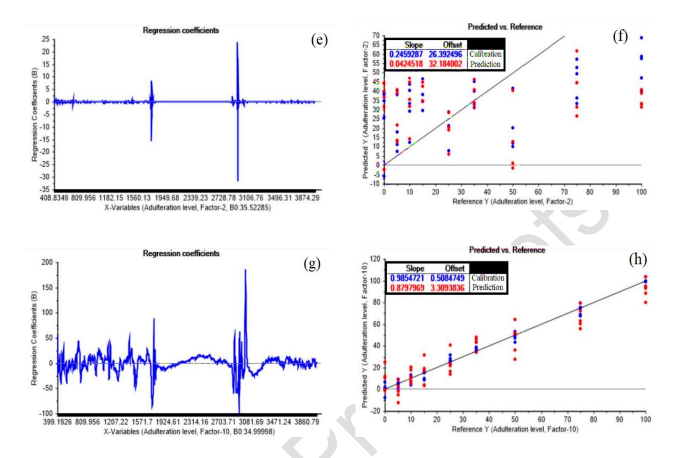


Fig 3: Spectra and quantification models (pure cinnamon and adulterated with dry bread powder) developed by PLSR, with (a, b) no preprocessing, (c, d) SNV with Savitzky-Golay D¹, (e, f) EMSC with Savitzky-Golay D², (g, h) OSC with Detrending.

In addition to scatter correction and smoothing, the preprocessing approach involved detrending to address any remaining baseline drifts. Detrending removes systematic baseline variations that may occur over the course of spectral data collection, particularly when dealing with complex matrices like powdered food products. This step was essential for ensuring that the PLSR model could accurately predict adulteration levels without interference from baseline noise. The final preprocessing pipeline consisted of a combination of OSC, SNV, EMSC, and derivative transformations, followed by mean-centering of the spectral data. These preprocessing techniques were chosen based on their proven effectiveness in previous studies and were fine-tuned to achieve optimal performance for this specific dataset [40–42].

Figure 3 illustrates the effect of preprocessing methods on pure cinnamon and cinnamon adulterated with dry bread powder. The dataset was divided into 70% calibration (n=84 samples) and 30% validation (n=36 samples) sets to ensure robust model training and testing. The grouping of data points in the scatter plot reflects the distinct adulteration levels (0–30% w/w), with each cluster representing a specific concentration. The PLSR model utilized 10 latent variables, selected based on minimizing the root mean square error (RMSE) of cross-validation. The processed data were validated using k-fold cross-validation (k=5) to assess the predictive capability of the PLSR model, confirming the preprocessing approach's impact on enhancing

model accuracy and robustness. The regression coefficients (slope = 0.98, offset = 0.51) indicate strong agreement between predicted and reference values ($R^2 > 0.90$).

2.4. Partial least squares regression (PLSR)

In this study, Partial Least Squares Regression (PLSR) was employed to establish a relationship between the spectral data and the level of adulteration in cinnamon powder samples. PLSR is widely used in chemometrics due to its ability to handle collinear and high-dimensional spectral data while predicting the concentration of target compounds. By projecting both the spectral matrix (X) and the adulteration levels (Y) into a new latent space, PLSR simplifies the complex relationship between the two variables, allowing for robust calibration models. This approach was chosen because of its effectiveness in handling multivariate data with overlapping spectral regions, as well as its ability to deal with noise and variability in spectral measurements [43]. One key factor in improving the accuracy of PLSR models is variable selection, which focuses on identifying the most relevant spectral regions. By excluding non-informative variables, the model is refined, improving both its predictive power and stability [44].

The performance of the developed PLSR models was evaluated using several statistical metrics. The calibration model's accuracy was assessed through the correlation coefficient (R²), which measures the goodness of fit between the predicted and actual adulteration levels. Additionally, the root mean square error (RMSE) was calculated for both the calibration (RMSEC) and prediction (RMSEP) datasets. Lower RMSE values indicate a more accurate and reliable model, while higher R² values suggest better predictive capability. Cross-validation techniques were also applied to minimize overfitting, ensuring that the PLSR models would generalize well to unseen data. This combination of metrics provided a comprehensive evaluation of model performance, confirming the effectiveness of the PLSR approach in predicting adulteration levels based on spectral information [45].

3. Results and Discussion

3.1. Spectral features of the samples

In Figure 4, raw spectra of samples including pure cinnamon powder, soybean powder, hazelnut shell powder, and dry bread powder in the range of 4000 – 400 cm-1 are shown. The differences between the spectra are attributed to the transmittance associated with various functional groups, which can increase or decrease depending on the sample composition. The absorption bands of water, carbohydrates, proteins, and lipids can explain this spectral treatment [46]. The known troughs in cinnamon spectra, corresponding to the stretching vibrations of C=O, C=C, and OH in aromatic compounds are approximately located at 1678 cm-1, 1626 cm-1, 1450 cm-1, and 1289 cm-1 [47]. Two prominent transmittance troughs in the nearby region of 2800 cm-1 in dry soybean suggest the symmetric stretching of fatty acid CH2 groups [48]. For hazelnut shell powder, valleys at ~2924 cm-1, ~2845 cm-1, ~1709 cm-1, and ~1613 cm-1 correspond to the vibrations of the alpha-hydroxy group (Alpha-CH), aromatic carbonyl/carboxyl C=O stretching, and aromatic C=C ring stretching, respectively [49]. The valley near 1373 cm-1 in dry bread powder is attributed to sulfone, a derivative of wheat flour [50]. As shown in Figure 4, the chemical composition of the investigated samples makes it difficult to distinguish between pure cinnamon powder and adulterated samples with the naked eye. This is because the spectra of all the samples appear very similar, with nearly parallel peaks.

To build upon this analysis, it's important to compare these findings with similar studies on food adulteration and spectral analysis. A similar study where they observed distinct spectral features

in various food powders, demonstrating that FTIR (Fourier-transform infrared) spectroscopy can effectively identify adulterants in food products [46]. In the current study, while the transmittance spectra for adulterated and pure cinnamon powder appear nearly identical, small differences in peak intensities and locations indicate the potential for advanced multivariate analysis techniques, such as PCA (Principal Component Analysis) or PLS-DA (Partial Least Squares Discriminant Analysis), to distinguish these samples more effectively.

Also another study confirmed that the presence of C=O and OH stretching vibrations in cinnamon is a reliable marker for the identification of pure samples [47]. However, the overlapping peaks with adulterated samples, as seen in this study, indicate that visual differentiation is insufficient. Similarly a research observed that hazelnut shell powder contains strong aromatic C=C and carbonyl groups, which can interfere with identifying certain adulterants [51].

In conclusion, while the spectral analysis shows similarities across samples, more refined data analysis methods can provide higher accuracy in distinguishing pure and adulterated samples. This comparison also highlights the need for complementary techniques like NIR (near-infrared) spectroscopy, which could further improve adulteration detection.

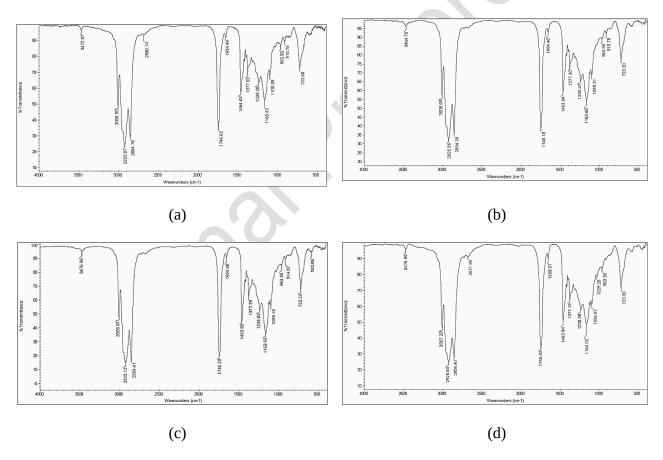


Fig 4: Transmittance spectrum of (a) pure cinnamon powder, (b) soybean powder, (c) hazelnut shell powder, and (d) dry bread powder samples.

3.2. Evaluation of PLSR model performance

Based on a literature review, we know that a PLSR model is considered suitable when it has a high coefficient of determination, low RMSE (<10%), and an R² value greater than 0.9 [52,53]. Based on the results (shown in Tables 2-4), the PLSR model demonstrated a high value of correlation coefficient of calibration (R2c) and a low root mean squared error of calibration (RMSEC) for all adulterations (R2c > 0.97 and RMSEC< 5%). However, it performed poorly in predicting adulterations, except for soybean powder (with an R²p value of 0.938 and an error of approximately 9%). For other adulterants, the R2p value was below 0.900 and the RMSEP was above 14%. These results highlight the importance of using preprocessing methods to enhance the accuracy of fraud prediction. In the following section, the effect of preprocessing methods on performance of the PLSR model for predicting the level of frauds is investigated.

These findings are consistent with other studies that have highlighted the challenge of achieving robust prediction for multiple adulterants without preprocessing techniques. For instance, Tan et al. (2024) reported similar difficulties in predicting adulteration in cumin powder without applying spectral preprocessing methods [54]. Studies like the one conducted by Khodabakhshian et al. (2023) also emphasize that while PLSR models can yield high calibration accuracy, the prediction accuracy often depends heavily on the use of preprocessing techniques to reduce noise and enhance the quality of the spectral data [34]. Thus, as seen in the present study, the relatively low predictive performance for most adulterants supports the argument that preprocessing methods are crucial for improving fraud detection accuracy in spectroscopy-based models. This will be further explored in the following section.

3.3. Effect of preprocessing methods on PLSR model

In this section, various preprocessing techniques – including 14 studied methods – and their effects on the PLSR model were investigated. The evaluation results of the PLSR model and the PLSR model combined with preprocessing methods for adulterated samples are presented in Tables 2-4.

Table 2. The results of PLSR for different preprocessing techniques (cinnamon powder adulterated with
hazelnut shell powder)

NO.	Preprocessing	Factor	R^2_{c}	RMSEC (%)	R^2_{p}	RMSEP (%)
1	None	15	0.993	2.823	0.867	14.800
2	OSC	5	0.995	2.355	0.971	7.571
3	SNV	18	0.996	1.562	0.841	15.827
4	EMSC	15	0.995	2.042	0.873	14.026
5	Detrending	15	0.995	2.212	0.881	14.063
6	Median filter	17	0.997	2.062	0.832	16.146
7	Savitzky-Golay D ¹	16	0.999	0.945	0.782	19.642
8	Savitzky-Golay D ²	15	0.999	1.258	0.632	25.531
9	OSC, Detrending	6	0.995	2.756	0.978	7.683
10	SNV, Median filter	16	0.998	1.563	0.846	15.892
11	SNV, Savitzky-Golay D ¹	15	0.999	1.405	0.768	18.376

12	SNV, Savitzky-Golay D ²	15	0.997	1.782	0.551	25.340
13	EMSC, Median filter	16	0.997	1.335	0.893	13.006
14	EMSC, Savitzky-Golay D ¹	16	0.998	1.582	0.773	18.621
15	EMSC, Savitzky-Golay D ²	17	0.998	1.593	0.571	25.367

With this vision, in cinnamon powder adulterated with hazelnut shell powder, the orthogonal signal correction (OSC) method with detrending (Figure 3g, h) yielded the most accurate prediction results, with an R_p^2 value of 0.978. It was able to estimate the level of hazelnut shell powder in cinnamon powder with an error of approximately 7%. Following OSC with detrending, OSC alone produced the second-best result with, an R_p^2 value of 0.971. The other results can be seen in Table 2.

It is worth noting that although models with a higher number of latent factors often show better fitting performance (higher R^2c and lower RMSEC), they are more prone to overfitting and multicollinearity, especially when dealing with high-dimensional spectral data. In contrast, reducing the number of factors generally leads to higher prediction errors (e.g., increased RMSEP), yet it also simplifies the model and reduces the dimensionality of the data matrix, which is desirable for robust model interpretation and generalizability. Therefore, a trade-off must be made between model complexity and predictive performance. For instance, in Table 2, while the raw data (no preprocessing) model used 15 factors and yielded a relatively high R^2c (0.993), its prediction performance ($R^2p = 0.867$) was lower than that of OSC with 5 factors ($R^2p = 0.971$). This indicates that using fewer factors with effective preprocessing (like OSC or OSC with detrending) can lead to more accurate and stable predictions despite lower apparent model fit during calibration.

Consequently, researchers must strike a balance between the number of latent variables and prediction accuracy. A model using 5–6 factors with lower RMSEP and higher R²p (such as OSC and OSC+Detrending) is often preferable over a highly fitted model using 15–17 factors with worse prediction errors, due to the risk of overfitting and reduced generalizability.

Table 3. The results of PLSR for different preprocessing techniques (cinnamon powder adulterated with soybean powder)

NO.	Preprocessing	Factor	R_{c}^{2}	RMSEC (%)	R^2_{p}	RMSEP (%)
1	None	13	0.991	3.095	0.938	8.537
2	OSC	5	0.985	3.978	0.971	6.169
3	SNV	11	0.993	2.917	0.963	7.352
4	EMSC	8	0.989	3.411	0.951	7.638
5	Detrending	10	0.988	3.587	0.932	8.922
6	Median filter	10	0.992	3.064	0.951	7.992
7	Savitzky-Golay D ¹	12	0.995	2.440	0.967	7.265
8	Savitzky-Golay D ²	12	0.991	3.101	0.942	8.230
9	OSC, Detrending	5	0.987	3.714	0.981	5.773
10	SNV, Median filter	9	0.993	3.023	0.951	7.393
11	SNV, Savitzky-Golay D ¹	11	0.995	2.698	0.963	6.696

12	SNV, Savitzky-Golay D ²	15	0.999	0.853	0.941	8.680
13	EMSC, Median filter	10	0.993	2.858	0.969	6.597
14	EMSC, Savitzky-Golay D ¹	11	0.994	2.672	0.964	6.782
15	EMSC, Savitzky-Golay D ²	16	0.999	1.015	0.931	9.279

Table 4. The results of PLSR for different preprocessing techniques (cinnamon powder adulterated with dry bread powder)

NO.	Preprocessing	Factor	R_{c}^{2}	RMSEC (%)	R^2_{p}	RMSEP (%)
1	None	12	0.972	4.488	0.811	14.159
2	OSC	10	0.971	4.518	0.882	11.287
3	SNV	15	0.994	1.767	0.878	11.500
4	EMSC	12	0.990	2.971	0.886	11.069
5	Detrending	11	0.981	4.010	0.864	12.109
6	Median filter	13	0.984	3.653	0.858	12.360
7	Savitzky-Golay D ¹	15	0.997	1.650	0.552	22.897
8	Savitzky-Golay D ²	3	0.321	25.880	0.212	30.996
9	OSC, Detrending	11	0.982	3.708	0.900	9.972
10	SNV, Median filter	12	0.988	3.098	0.874	11.626
11	SNV, Savitzky-Golay D ¹	12	0.987	3.100	0.771	15.716
12	SNV, Savitzky-Golay D ²	2	0.332	25.678	0.195	31.107
13	EMSC, Median filter	12	0.989	3.169	0.889	10.826
14	EMSC, Savitzky-Golay D ¹	15	0.989	1.122	0.793	14.796
15	EMSC, Savitzky-Golay D ²	3	0.351	25.316	0.115	31.266

As emphasized in section 2.3, we employed preprocessing manners to increase the accuracy of the prediction models. After examining the results shown in Table 2, we discovered that certain preprocessing methods, such as SNV, median filter, and SNV with median filter, underperformed when compared to the un-preprocessed data of adulterated cinnamon powder and hazelnut shell powder. Furthermore, some preprocessing methods, such as Savitzky-Golay D^2 , EMSC with Savitzky-Golay D^2 , and SNV with Savitzky-Golay D^2 , had a significantly negative impact on prediction accuracy (Table 2). The poorest result was obtained using SNV Savitzky-Golay D^2 which yielded an R^2_p value of 0.551.

Regarding cinnamon powder adulterated with soybean powder, almost all of preprocessing methods demonstrated suitable performance and improved the R^2_p value of the PLSR models. In some cases, such as detrending and EMSC with Savitzky-Golay D^2 , the accuracy was lower compared to the un-preprocessed data (Table 3). However, in this case, OSC with detrending showed the highest accuracy in prediction, with an R^2_p value of 0.981. It was able to estimate the level of soybean powder in cinnamon powder with an error of approximately 6%. Additionally, OSC produced the second-best results with an R^2_p value of 0.971. Furthermore, several methods including EMSC, median filter, and SNV with median filter illustrated similar results ($R^2_p = 0.951$) (Table 3).

For cinnamon powder adulterated with dry bread powder (Table 4), approximately half of the preprocessing methods had a negative effect, with some cases showing a significant negative impact (such as Savitzky-Golay D^2 , SNV with Savitzky-Golay D^2 , and EMSC with Savitzky-Golay D^2 , resulting in an $R^2_p < 0.22$). The poorest prediction accuracy was obtained using EMSC with Savitzky-Golay D^2 with an R^2_p value of 0.115 and an RMSEP of 31.266% (Figure 3e, f). However, in other cases, the preprocessing methods improved prediction accuracy compared to the un-preprocessed spectra of adulterated samples. The OSC with detrending method yielded an acceptable with a R^2_p value of 0.900 and an RMSEP=9.972%).

3.4. Comparison of preprocessing methods

It is important to keep in mind that the ultimate goal of a chemometrician is to develop a reliable model that can accurately predict results. As observed earlier, most preprocessing methods had a positive effect on prediction results and improved prediction accuracy. However, in some cases, this effect was negligible, and in some cases, it was significant. Some methods showed worse results than un-preprocessing data. Our results indicated that preprocessing techniques play a crucial role in enhancing the predictive performance of the model, but the effectiveness of each method varied. The OSC (Orthogonal Signal Correction) combined with detrending proved to be the most effective technique, reducing the prediction error to a range of 5-10%. This high accuracy suggests that OSC can effectively remove uninformative variation from the spectral data, making it highly suitable for this specific application. On the contrary, methods such as EMSC (Extended Multiplicative Signal Correction) with Savitzky-Golay D² did not perform well across the board. This result aligns with other studies that indicate the potential limitations of certain preprocessing techniques in handling complex datasets. For instance, a study by Liu et al. (2018) also reported poor results using EMSC in the context of powdered food adulteration, attributing the failure to the method's inability to adequately account for baseline shifts and noise [55].

When comparing our findings with other research in the field, such as the work by Wu et al. (2023), we observed that preprocessing methods like MSC (Multiplicative Scatter Correction) and SNV (Standard Normal Variate) often improve predictive accuracy, especially in cases of adulteration with lower levels of contaminants [56]. However, in our study, the application of MSC was not as effective as OSC, which might be due to the specific characteristics of cinnamon powder and the adulterants used (e.g., soybean powder, hazelnut shell powder). Similar conclusions were drawn by Khodabakhshian et al. (2022), who emphasized that the performance of preprocessing methods depends on the type of adulterant and the specific spectral properties of the sample [43].

This variability in effectiveness highlights the importance of selecting the appropriate preprocessing technique based on the characteristics of both the sample and the adulterant [57]. While prior studies offer valuable insights, relying solely on their results can lead to suboptimal outcomes. Our study confirms that preprocessing techniques must be evaluated and tailored to the specific adulteration scenario at hand, as supported by findings in related research. In summary, although there is a general consensus on the positive impact of preprocessing on prediction accuracy, our study demonstrates that not all methods are equally effective. Therefore, it is critical for researchers to carefully assess and select the most suitable preprocessing techniques for their specific case studies.

4. Conclusion

In this study, we investigated the use of a FTIR spectroscopy system and multivariate analysis to predict the level of adulteration in cinnamon powder and we examined the effects of several data preprocessing methods on the accuracy of this prediction. We evaluated the quantitative performance of 14 preprocessing techniques and assessed their efficacies on the prediction accuracy of the PLSR model. According to the results, OSC with detrending demonstrated the best results for all of the adulterants (soybean powder, hazelnut shell powder, and dry bread powder), with an R²_p value of greater than 0.90 and an RMSEP value of under 10% for predicting cinnamon powder adulteration. On the other hand, EMSC with Savitzky-Golay D² showed the lowest accuracy in predicting adulterants with an RMSEP range of 9-32%. However, in comparison with the other adulterants, the accuracy of soybean powder prediction was slightly superior. In summary, the PLSR model based on OSC with detrending could predict cinnamon powder adulteration with high accuracy and this combination performed best in all adulterants, while other integrations reduce the model's performance compared to using un-preprocessed data. Furthermore, given the variation in the effectiveness of preprocessing approaches across different adulterants, selecting an appropriate preprocessing technique is crucial, as otherwise, it may reduce the accuracy of fraud pediction. Given the nature of soft computing techniques, it is recommended that future research investigate the use of artificial neural networks and other machine learning methods on predictive accuracy.

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Data availability: The data that support the findings of this study are available from the corresponding author (Dr. Rasool Khodabakhshian) upon reasonable request.

Declarations

Ethical approval: No ethical clearance is required.

Human and animal rights: We declare that there are no animal studies or human participant involvement in the study.

Conflict of interest: The authors declare no competing interests.

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مقاله پژوهشی ارزیابی تقلب در ادویه دارچین بوسیله طیفسنجی: تأثیر پیش پردازش دادهها بر مدلهای پیش بینی چند متغیره

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تشخیص تقلب در زنجیره تأمین دارچین برای تضمین ایمنی مصرف کننده و حفظ یکپارچگی محصول بسیار مهم است. پیشرفتهای اخیر در تکنیکهای پیشپردازش دادههای طیفی، دقت بیشتری را در شناسایی تقلبات در ادویههایی مانند دارچین ارائه میدهد. این مطالعه به بررسی تأثیر تکنیکهای مختلف پیشپردازش طیفی بر پیشبینی تقلبات – بهویژه پودر سویا، پودر پوسته فندق و پودر نان خشک – مخلوط با پودر دارچین با استفاده از طیفسنجی همراه با تجزیه و تحلیل چند متغیره میپردازد. طیفهای عبوری در محدوده مادون قرمز میانی 1 ۴۰۰ cm دارچین با استفاده از رگرسیون حداقل مربعات جزئی (PLSR) برای مدلسازی سطوح تقلب بر اساس این طیفها استفاده شد. روشهای پیشپردازش مختلفی برای بهینهسازی دادههای طیفی اعمال شد .در میان آن ها، OSC همراه با Detrennding بالاترین دقت پیشبینی را با ضریب پیشبینی (R²p) از ۹۹۰۰ تا ۹۸۰۱ به دست آورد. در مقابل، Savitzky-Golay D² و Savitzky-Golay D² با مقادیر پیشبینی را با ضریب پیشبینی (۱۹۹۳) و ۱۹۹۲) با محدوده خطای پیشبینی ۵ تا ۱۰ درصد، آسان ترین تقلب برای تشخیص بود. این یافتهها بر اهمیت انتخاب تکنیکهای پیشپردازش مناسب برای بهبود دقت تشخیص تقلب در پودر دارچین با استفاده از روشهای طیفسنجی یافتهها بر اهمیت انتخاب تکنیکهای پیشپردازش مناسب برای بهبود دقت تشخیص تقلب در پودر دارچین با استفاده از روشهای طیفسنجی تأکید می کنند.

كليد واژه ها: تقلبات، شيمي سنجي، پيش يردازش دادهها، ايمني مواد غذايي، أناليز طيفي.

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