In situ monitoring of nitrate content in leafy vegetables using mid-infrared attenuated total reflectance spectroscopy coupled with intelligent algorithm

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**ABSTRACT:**

**Background:** Vegetables are one of the most important nitrate sources of human diet. Establishing fast and accurate *in situ* nitrate monitoring approaches that could be used in the plant growth process and vegetable markets is essential.

**Results:** Incorporating the unique feature of N–O asymmetric stretch absorption in the mid-infrared region (1500-1200 cm⁻¹), portable Fourier-transform infrared attenuated total reflectance (FTIR-ATR) spectroscopic instruments, along with the Euclidean distance-modified intelligent algorithm extreme learning machine (ED-ELM) model, were employed to evaluate the nitrate contents in leafy vegetables. A total of 1224 samples of four popular vegetables (Chinese cabbage, swamp cabbage, celery, and lettuce) were analyzed. The results indicated that the nitrate contents (mean values: Chinese cabbage: 7550 mg/kg; swamp cabbage: 4219 mg/kg; celery: 4164 mg/kg; lettuce: 4322 mg/kg) highly exceeded the World Health Organization (WHO)-specified maximum tolerance limits. The ED-ELM model showed a better performance with the root-mean-square-error of 799.7 mg/kg, the determination coefficients of 0.93, the ratio of performance to deviation of 2.22, the optimized calibration dataset number of 100, and the number of hidden neurons of 30.

**Conclusion:** The results confirmed that FTIR-ATR, along with the suitable model algorithms, could be used as a potential rapid and accurate method to monitor the nitrate contents in the fields of agriculture and food safety.

**Keywords:** Leafy vegetables; nitrate, mid-infrared attenuated total reflectance; intelligent algorithm; extreme learning machine
Background

Nitrate is the most important form of nitrogen in the environment and human diet. Vegetables are one of the most important sources of vitamins, minerals, and biologically active compounds with regard to human nutrition [1]. Vegetables, in daily human diets, are also the primary sources of ingested nitrates, and about 80-85% of the daily nitrate intake comes from vegetables [2, 3], along with fruits, water, and additives in meat [4, 5].

Generally, nitrate is not treated as a directly toxic ion, but when it is mixed with food, it can be converted into nitrite by commensal bacteria in the mouth and gastrointestinal tract. Nitrites are further converted into nitrosamines, which are carcinogenic and teratogenic N-nitroso compounds [1, 6]. Previous studies have indicated that nitrate may be beneficial for human health [7, 8], and it was assumed that dietary nitrate could generate nitric oxide, which has antimicrobial effects on gut pathogens, thus providing gastric protection against microbial infections [8, 9]. Whether nitrate can be referred to as an essential nutrient or a food contaminant with potential adverse effects depends on its concentration [10]. Thus, the content of ingested nitrate is critical, and it is necessary to set a maximum limit value for nitrate intake [1].

On one hand, nitrate accumulation in plants, especially in most leafy vegetables, is a major concern [11]. Nitrates in the soil are the primary nutrients required for plant growth [12]. Moreover, nitrate fertilizers that have been used in agriculture result in the accumulation of high levels of nitrate in a variety of vegetables [3]. When nitrate uptake exceeds nitrate assimilation by the plant, accumulation of nitrate in plant tissues occurs. Other factors in the plant growth process that may influence nitrate concentration include the plant species, environmental conditions (e.g. light intensity,
temperature, and humidity), harvest time, and storage time [13-15]. For instance, a significant decrease in nitrate level is observed at ambient temperatures, but nitrate level remains constant over time during storage under refrigerated conditions [1, 16].

On the other hand, risk assessment of the safety of dietary nitrate intake and exposure from vegetables has been a major health concern in many countries in recent decades. According to the International Agency for Research on Cancer [17], vegetables can be divided into three levels based on their nitrate concentrations: low nitrate (< 100 mg/kg), medium nitrate (100–1000 mg/kg), and high nitrate (> 1000 mg/kg) [17]. The European Commission established the maximum level of nitrates in two leaf vegetables: 2000-3500 mg/kg in spinach and 2000-4500 mg/kg in lettuce [18].

Therefore, establishing fast and accurate nitrate monitoring approaches, especially nondestructive in situ methods that could be used in the plant growth process and vegetable markets, is essential. Over the years, spectrophotometry, photometry, potentiometry, spectrofluorimetry, ion chromatography, gas chromatography and high-performance liquid chromatography have been used widely to monitor the contents of various compounds in vegetables and other foods [14, 19-23]. However, these approaches require sample pre-preparation and need to be performed in the laboratory. Portable instruments could be used effectively for in situ nitrate monitoring. The use of Fourier-transform infrared (FTIR) spectroscopy techniques has been spreading widely owing to the advantages of this nondestructive, fast, and reliable approach for quality assessment in agro-food industries. Nitrogen distribution in the leaves of Chinese cabbage and nitrogen status in rice were measured based on FTIR photoacoustic spectroscopy [24, 25]. The nutritional and
functional components of leafy vegetables, including Chinese cabbage, beans, and pea seeds, can be successfully determined by the near-infrared and mid-infrared total and diffuse reflectance spectroscopy methods using the absorption feature at various wavelength/wavenumber [26-29]. Nitrate in the soil was reported to be detected by attenuated total reflectance (ATR) spectroscopy [30]. Thus, the use of portable FTIR instruments has extended the application of spectroscopy in the field of agriculture in recent years, making it possible for in-situ and real-time measurements of nitrate contents in vegetables in vegetable markets and the plant growth process.

Based on the analysis of the collected FTIR spectra data, a rapid, robust, computationally efficient artificial intelligence-based model framework was developed for the in situ and real-time monitoring of the nitrate content in leafy vegetables. To address complex data sets with many predictor variables, machine learning techniques are extensively used among researchers [31]. Here, estimating vegetable nitrate content using a proper machine-learning model was the basis for in situ measurements. Chinese cabbage (Brassica rapa subsp. chinensis), swamp cabbage (Ipomoea aquatica Forssk), celery (Apium graveolens L.), and lettuce (Lactuca sativa L.), the main vegetable species grown and consumed in southeast China, were chosen for this study. Fresh samples of the above-mentioned vegetables were purchased in local markets during harvest time (vegetative stage). The objectives of this research were to: i) investigate the nitrate contents in local leafy vegetables; ii) collect the spectral data of these vegetables using a portable FTIR-ATR device and analyze the features of nitrate absorption in vegetables; and iii) build an accurate in situ intelligent algorithm method for nitrate content measurements. The results provided a potential method on the development of a fast and feasible approach to monitor nitrate contents even in markets for the healthy management and
consumption of foodstuffs, making this research a good strategy for food safety assessment.

Methods

Collection of leafy vegetable samples

Four species of leafy vegetables, Chinese cabbage, swamp cabbage, celery, and lettuce, were purchased from four large local supermarkets and traditional wet markets in Nanjing, China, from August 2019 to September 2020. A total of 408 samples of the four species of vegetables were purchased (102 samples of each species). Thirty-two samples of each of the four species of vegetables were purchased from four markets on a single day, and they were analyzed by both spectral and laboratory methods on the same day to ensure that the vegetables were fresh and nitrate contents were relatively stable.

Chemicals

Hydrochloric acid ($\rho = 1.19$ g/mL, analytical reagent grade, AR), potassium nitrate (AR), ammonia (25% wt.), zinc sulfate heptahydrate (AR), potassium ferrocyanide (AR), octanol (AR), and activated carbon powder were purchased from the China National Pharmaceutical Group Corporation (Beijing, China). Deionized water ($15\Omega \text{ cm}^{-1}$) was prepared using a laboratory water system.

Nitrate content analysis

Nitrate content was analyzed following the national standard method (GB/T 5009.33–2016, China: Rapid determination of nitrate in vegetables), established by the Chinese National Institute of Metrology. The vegetable samples were mashed
using a planetary ball mill for 10 min. Then, vegetable homogenate (5 g), deionized water (10 g), ammonia buffer (5 mL) (pH = 9.6–9.7), and activated carbon powder were added to a conical flask, and the mixture was stirred (200 r/min) at 25°C for 30 min. The mixture was then transferred to a volumetric flask (250 mL) and mixed with 150 g/L potassium ferrocyanide solution (2 mL) and 300 g/L zinc sulfate solution (2 mL); deionized water was added to bring the volume of the resulting solution to 250 mL. This mixture was kept standing for 5 min and then filtered. Then, the filtered solution (2-3 mL) and deionized water were placed in a volumetric flask (25 mL), and the absorbance of the samples was measured at 219 nm by ultraviolet spectrophotometry. Meanwhile, nitrate standard solutions of different concentrations were measured to plot the standard curve. Finally, the nitrate content was calculated using the following formula:

\[
C_{\text{nitrate}} = \frac{\rho \times V_e \times V_u}{m \times V_a}
\]  

(1)

where \( C \) is the nitrate content in mg/kg; \( \rho \) is the value of nitrate obtained from the ultraviolet spectrophotometry standard curve; \( V_e \) is the constant volume in the volumetric flask used in the extraction process (250 mL); \( V_u \) is the constant volume for ultraviolet spectrophotometry measurement (25 mL); \( m \) is the mass of the vegetable homogenate (accurate to 0.01 g); and \( V_a \) is the volume of the filtered solution.

**FTIR-ATR spectroscopy**

Each vegetable was ground, and the vegetable juice obtained was scanned on a hand-held TruDefender FTIR spectrometer with an ATR spectra accessory (Thermo Fisher Scientific, USA). Spectra of the samples were recorded over the original range
from 4000 to 400 cm$^{-1}$, with a spectral resolution of 4 cm$^{-1}$. Atmospheric and instrumental noise was corrected by subtracting the background noise from each scan. Each ground sample was placed on a diamond reflection probe for three measurements, and a blank reference was scanned before the spectra for each sample were recorded. Spectra from the 1224 samples were collected. The spectra of a serial nitrate standard solution (with concentrations ranging from 0 to 15000 mg/kg) were recorded before each of the four samples was scanned.

**Pre-processing of the spectra**

The FTIR-ATR spectra were pre-processed with a Savitzky–Golay smoothing filter to eliminate baseline float and noise and improve the signal-to-noise ratio [24, 25]. Savitzky–Golay smoothing seemed to be superior to adjacent averaging because it reserved spectral features, such as peak height and width. The spectral range from 1500 to 1200 cm$^{-1}$ was selected based on the absorption characteristics of nitrate. Moreover, the second derivative spectra of nitrate in the range from 1500 to 1200 cm$^{-1}$ were obtained. Principal component analysis (PCA) was also performed. The MATLAB R2013a software (MathWorks, Natick, MA, USA) and related scripts were used to perform other statistical analyses.

**Theoretical overview**

The pre-processed spectra were divided into calibration and validation datasets. Then, the extreme learning machine (ELM) model, an intelligent algorithm, was employed to predict the nitrate contents. To improve the prediction accuracy, the calibration dataset was modified before being calibrated by the Euclidean distance (ED) method. Meanwhile, the partial least squares (PLS) model was used for comparison.
Subsequently, the performance of the models and prediction results were evaluated.

Calibration and validation datasets

The 1224 vegetable samples were randomly divided into a calibration dataset (training dataset) containing 74% (900 samples) of the spectra, and a validation dataset, containing the remaining 26% (324 samples) of the spectra.

Modified Euclidean distance

Each FTIR-ATR spectral curve represented the spectral features and concentration level of nitrate, and the linear matching of the nitrate concentration and spectra curves showed good correlation coefficients \([32]\). The spectral curves within a small intensity range were similar, and the nitrate concentrations were closer. Based on this observation, the ED method was employed to recognize similar spectral curves in the calibration data set for further modeling. The ED method was selected in this study based on our previous result, which reported that it was suitable for spectral identification \([33]\). ED between the calibration and target samples was computed using pairs of curves and their derivatives as a measure of similarity for clustering.

\[
ED_{ik} = \sqrt{\sum_{j=1}^{p} (x_{ij} - x_{kj})^2} \tag{2}
\]

where \(ED_{ik}\) is the Euclidean distance between the \(i\)th target sample \(x_i\) and each \(k\)th calibration sample \(x_k\), \(k \neq i\); and \(j\) is the variable index, \(j = 1, 2, \ldots, p\). The calibration data set sequence was re-ordered in an ascending manner based on the ED results, which meant that similar spectra were near-neighbors.

Extreme learning machine model
ELM belonged to a single hidden-based layer forward network [34]. For a sample set 
\((x_i, t_i)\), where \(x_i = (x_{i1}, x_{i2}, \ldots, x_{in})^T \in \mathbb{R}^n\) and \(t_i = (t_{i1}, t_{i2}, \ldots, t_{in})^T \in \mathbb{R}^k\), the standard single hidden-based layer forward network with \(L\) hidden nodes and activation function \(h(x)\) was mathematically modeled as [35] :

\[
\sum_{i=1}^{L} \beta_i h_i(x_j) = \sum_{i=1}^{L} \beta_i (w_i \times x_j + b_i) = o_j
\]  

(3)

where \(j = 1, 2, \ldots, n\); \(w_i = (w_{i1}, w_{i2}, \ldots, w_{in})^T\) is the weight vector connecting the \(i\)th hidden node to the input nodes, \(\beta_i = (\beta_{i1}, \beta_{i2}, \ldots, \beta_{ik})^T\) is the weight vector connecting the \(i\)th hidden node to the output nodes; and \(b_i\) is the threshold of the \(i\)th hidden node. Then,

\[
H \beta = T
\]  

(4)

\[
H = \begin{bmatrix}
h_i(w_{i1}x_1 + b_i) & \cdots & h_i(w_{iL}x_1 + b_i) \\
\vdots & \ddots & \vdots \\
h_i(w_{i1}x_n + b_i) & \cdots & h_i(w_{iL}x_n + b_i)
\end{bmatrix}_{N \times L}
\]

\[
\beta = \begin{bmatrix}
\beta_1^T \\
\vdots \\
\beta_L^T
\end{bmatrix}_{L \times K} \quad \text{and} \quad T = \begin{bmatrix}
t_1^T \\
\vdots \\
t_L^T
\end{bmatrix}_{N \times K}
\]  

(5)

The difference between conventional gradient-based solution methods and the ELM method was that the ELM method determined the function by using the formula:

\[
\beta = H^+T
\]  

(6)

where \(H^+\) is the Moore-Penrose generalized inverse of matrix \(H\).

In addition, the ELM input contained the training dataset and number of hidden neurons \(L\). The output included pre-processing the training data set (normalization), partitioning the available data set into the training and validation data sets, and computing the hidden layer output values of the ELM model. For the validation dataset, each validation spectrum possessed its re-ordered calibration dataset, which
was obtained using the ED method. Thus, the aim of this step was to obtain the
237 correct number of calibration datasets and the ELM hidden layer.

239 Partial least squares model

240 This was a bilinear model where a matrix $X$, containing the variables (spectra
241 wavenumber), and matrix $Y$, a function of the variables of $X$ (nitrate contents), were
242 used for the prediction of the smallest number of latent variables. In this study, the
243 optimal number of latent variables for each database model was determined based on
244 the minimal root-mean-square-error ($RMSE$) of cross-validation by leave-one-out
245 cross-calibration [36].

247 Model performance evaluation

248 The evaluation indices of predictive capability for the ELM and PLS models were
249 coefficients of determination ($R^2$), the ratio of performance to deviation ($RPD$), $RMSE$,
250 Willmott’s index ($WI$), and the Legates and McCabe index ($E_{LM}$) [35, 37].

$$R^2 = \frac{\sum_{i=1}^{n} (y_i' - \bar{y})^2}{\sum_{i=1}^{n} (y_i - \bar{y})^2} \quad (7)$$

$$RMSE = \sqrt{\frac{1}{n} \sum_{i=1}^{n} (y_i' - y_i)^2} \quad (8)$$

$$RPD = \frac{SD}{RMSE} \quad (9)$$

$$WI = 1 - \left[ \frac{\sum_{i=1}^{n} (y_i' - y_i)^2}{\sum_{i=1}^{n} (|y_i' - \bar{y}| + |y_i - \bar{y}|)^2} \right], 0 \leq WI \leq 1 \quad (10)$$

$$E_{LM} = 1 - \left[ \frac{\sum_{i=1}^{n} |y_i - y_i'|}{\sum_{i=1}^{n} |y_i - \bar{y}|} \right], (\infty \leq E_{LM} \leq 1) \quad (11)$$
where $y'_i$ and $y_i$ are the predicted data and data measured by the chemical analysis method, respectively; $n$ is the number of data sets; and $SD$ is the standard deviation. $RMSE_c$ and $RMSE_p$ represented the root-mean-square-error in the calibration and validation dataset models, respectively. The $RPD$, which is used for normally distributed data, represented prediction accuracy, and should be higher than 1.8. An $RPD$ value between 2 and 2.5 indicated a good quantitative prediction model, while a value higher than 3 suggested excellent performance of the model. A good performance model should have a $WI$ value close to 1. Hence, insensitivity could be overcome because the ratio of model errors, rather than the square of the model error difference, could be analyzed [38]. $ELM$, which was a more robust parameter than $WI$, predicted relatively higher values by squaring the differences [35, 37].

In addition, the ratio of $RMSE_p$ to $RMSE_c$ was used to judge the robustness of the model. A ratio lower than 1.2 was usually considered as a measure of robust performance [39, 40].

**Results**

**Nitrate content analysis**

A total of 408 vegetable samples were analyzed; the nitrate contents of Chinese cabbage, swamp cabbage, celery, and lettuce are listed in Table 1. The nitrate contents of Chinese cabbage (4063–14104 mg/kg), with an $SD$ of 1664 mg/kg, was extremely high. The highest value was 14104 mg/kg, which was more than four-fold higher than the level indicating serious contamination. The nitrate content of swamp cabbage (2111-6607 mg/kg), with an $SD$ of 1029 mg/kg, showed the narrowest range among the four species, but the average (4219 mg/kg) and median (4196 mg/kg) values were beyond the levels indicating serious contamination. The nitrate content of celery was
805–8643 mg/kg, with an SD of 1214 mg/kg; the average (4164 mg/kg) and median (1214 mg/kg) values were slightly lower than those of the other species. As reported by Kalaycıoğlu and Erim (2019), the fiber compounds in the vegetables could reduce the possible harmful effects of the high nitrate contents [1]. The nitrate content in lettuce (10485 mg/kg), which was two to three fold higher than the maximum nitrate content specified by the European Commission standard, was the highest. Fig. 1 shows the different nitrate contents in the four leafy vegetable samples. The “+” symbol represents the outliers with extreme nitrate contents. The lower and upper lines of the boxplot represent the first and third quartiles (25th and 75th percentiles), respectively, and the median value (50th percentile) was marked by the central line. Two horizontal lines were drawn out from the first and third quartiles to the smallest and largest non-outliners, respectively [35]. Additionally, all the nitrate contents followed the Gaussian distribution.

Spectral characterization

The characteristic absorption of nitrate, in the range from 1500 to 1200 cm\(^{-1}\), in the Chinese cabbage, swamp cabbage, celery, and lettuce samples is shown in Fig. 2(a), and the main peaks of nitrate were found at around 1401 cm\(^{-1}\) and 1350 cm\(^{-1}\). The absorption bands at 1401 cm\(^{-1}\) and 1350 cm\(^{-1}\) were associated with the N–O asymmetric stretching mode (\(v_3\)). They were generated by the splitting of the \(v\) generation mode into two bands labeled \(v_{3, \text{high}}\) and \(v_{3, \text{low}}\). Hudson et al. [41] reported that the peak at around 1400 cm\(^{-1}\) was not observed in a pure nitrate environment, but the spectra were similar to those of aqueous Ca(NO\(_3\))\(_2\). The peak at 1245 cm\(^{-1}\) was associated with nitrite (NO\(_2^-\)) [42].

Moreover, the second-order derivative spectra were then calculated and plotted,
as shown in Fig. 2(b). The peak at approximately 1460 cm\(^{-1}\) was associated with N=O vibration, the peaks at 1375 cm\(^{-1}\) and 1363 cm\(^{-1}\) were attributed to N=O and N–O, respectively, and the peak at 1300 cm\(^{-1}\) was associated with N–O vibration [30].

The first three principal components (PCs), PC1 (83.58 %), PC2 (9.42 %), and PC3 (4.05 %), containing 97.05% of the total spectral information, were investigated among the four vegetables, as shown in Fig. 3. These PCs for the four species were clustered, indicating that the main information was similar due to the characteristic absorption peaks of nitrate in the range from 1500 to 1200 cm\(^{-1}\). On the other hand, the clustered PCs also indicated that the nitrate spectra could not be separated by the general discrimination method.

### Optimization of parameters for calibration

Several parameters, including the numbers of calibration data sets, hidden neurons in the ELM model, and latent variables, are vital indices for the process of building a good model, and they should be determined and optimized before modeling. Here, the parameters of ELM, ED-modified ELM (ED-ELM), and PLS methods were investigated separately. The predictive capability of the models was determined by a low RMSE, large \(R^2\) (close to 1), and a value of \(RPD\) that was more than 1.8. For the ELM and PLS methods, there were 900 calibration samples and 324 validation samples. Optimal numbers of hidden neurons and latent variables were optimized, and are listed in Table 2. For the ED-ELM method, each of the 324 validation dataset samples had 900 re-ranked calibration datasets, in which the similarity degree sequence was arranged in a descending order. According to Ma et al. [40], a good calibration dataset should contain large variances and less interference. Thus, the optimal numbers of calibration data sets and hidden neurons were determined based
on the model performance parameters. Table 2 summarizes the performance indices, including the number of calibration data sets (900 samples). The optimal number of hidden layers was forty for the ELM model, while the optimal number of latent variables was seven for the PLS model. The values of $\text{RMSE}_C$ were $\approx 1089.91$ mg/kg in the ELM model and $\approx 1087.68$ mg/kg in the PLS model, indicating that the performance of the two models was similar. For the ED-ELM model, the optimal number of calibration datasets was determined according to the model performance parameters $R^2$, $\text{RPD}$, and $\text{RMSEP}$, as shown in Fig. 4. It was clear that with the increase in the calibration dataset number from 100 to 900, in intervals of 100, the values of $R^2$ decreased from 0.89 to 0.71, and the values of $\text{RPD}$ decreased from 3.05 to 1.76. Meanwhile, $\text{RMSEP}$ increased from 636.45 mg/kg to 1035.25 mg/kg. Thus, the optimal number of calibration datasets was in intervals of 100 in the ED-ELM model. This was probably because too large a dataset sample might introduce interference, which would reduce the performance of the model [40].

In any model, the number of neurons in a hidden layer is important to determine the ideal network architecture [38]. A small architecture could short sufficient degrees of freedom to correctly learn the predictor data, while an elaborately large architecture may not converge in a reasonable model execution time, or it may over-fit rather than generalized the data [35]. Thus, the number of hidden neurons was optimized to the range from 10 to 90, and the ratio of $\text{RMSEP}$ to $\text{RMSE}_C$, $\text{RPD}$, $R^2$, $WI$, and $\text{ELM}$ were calculated as functions of the number of hidden neurons. It can be noted in Fig. 5(a) that the ratio of $\text{RMSEP}$ to $\text{RMSE}_C$ increased with an increase in the number of hidden neurons. In contrast, the values of $\text{RPD}$ and $R^2$ decreased. Considering that the ratio of $\text{RMSEP}$ to $\text{RMSE}_C$ should be lower than 1.2, $\text{RPD}$ should be higher than 1.8, and $R^2$ should be close to 1, the optimal number of hidden neurons was found to be 30.
\( \text{RMSE}_p / \text{RMSEC} \approx 1.13; \quad \text{RPD} \approx 2.2, \quad \text{and} \quad R^2 \approx 0.93 \). Meanwhile, the WI and ELM values in Fig. 5(b) are presented in a parabolic shape, with an increase in the number of hidden neurons, and the values of WI \( \approx 0.85 \) and \( E_{LM} \approx 0.64 \) were the highest when the number of hidden neurons was 30. Thus, using the ELM model, the number of calibration datasets was set as 100, and the optimal number of hidden neurons was determined to be 30.

**Nitrate prediction model**

Based on the optimization parameters, the validation data set was predicted using the ELM, PLS, and ED-ELM models and the scatter plots are displayed in Fig. 6. The performance indices were \( \text{RMSE}_p \approx 995.77 \text{ mg/kg}, \quad R^2 \approx 0.70, \quad \text{and} \quad \text{RPD} \approx 1.76 \) in the ELM model and \( \text{RMSE}_p \approx 1172.01 \text{ mg/kg}, \quad R^2 \approx 0.65, \quad \text{and} \quad \text{RPD} \approx 1.66 \) in the PLS model. The RPD values predicted by both the ELM and PLS models were lower than 1.8, indicating that the models were not reliable. For the ED-ELM model, the \( \text{RMSE}_p \) was \( \approx 799.67 \text{ mg/kg} \), the \( R^2 \) was \( \approx 0.93 \), and the \( \text{RPD} \) was \( \approx 2.22 \), indicating that this model showed the best performance.

The Taylor diagram graphically depicted how closely the position of a prediction pattern matched the measured values, and it was used for evaluating the performance of multiple models [43, 44]. The position between the predicted and measured patterns was quantified in terms of the correlation of the \( (R^2) \), centered \( \text{RMSE}_p \) with the SD to evaluate the model that yielded values that were closest to the predicted values. The closer the predicted pattern to the measured pattern, the better the performance of the model is. The Taylor diagrams of the PLS, ELM, and ED-ELM models are shown in Fig. 7, showing the models that yielded data closest to measured data in the validation dataset. The predicted patterns that agree well with the measured
values are placed at the nearest position marked ‘measured’. The organ color contours indicate the centered $\text{RMSE}_p$ values. The red square at the bottom of the diagram represents the position of the measured nitrate content. It represented that: $R^2 = 1$, $SD = 1$, and centered $\text{RMSE}_p = 0$, when compared to itself. The blue diamond represents the position of the values ($R^2 \approx 0.65$, $SD \approx 1.11$, and centered $\text{RMSE}_p \approx 0.67$) predicted by the PLS model; the yellow rounded shape shows the position of the values ($R^2 \approx 0.69$, $SD \approx 0.98$, and centered $\text{RMSE}_p \approx 0.57$) predicted by the ELM model; and the green triangle represented the position of the values ($R^2 \approx 0.93$, $SD \approx 0.93$, centered $\text{RMSE}_p \approx 0.47$) predicted by the ED-ELM model. Thus, the PLS model-derived values were the farthest from the measured values, while the ED-ELM model-derived values were the nearest to the measured values.

**Nitrate status monitoring**

According to the WHO-specified tolerance levels of nitrate content in fresh vegetables, four scales were used: the low level (nitrate content $\leq 432$ mg/kg), allowing consumption of raw vegetables; medium level ($432$ mg/kg $< \text{nitrate content} \leq 785$ mg/kg), indicating that consumption of raw vegetables was not allowed; high level ($785$ mg/kg $< \text{nitrate content} \leq 1234$ mg/kg), indicating that consumption of raw or salted vegetables was not allowed; and serious contamination level ($1234$ mg/kg $< \text{nitrate content} \leq 3100$ mg/kg), indicating that consumption of the vegetable in any form was not allowed [45]. Considering the WHO-specified tolerance level of nitrate, nitrate contents in all four species of vegetables exceeded the maximum limit of consumption. The FTIR-ATR spectra could be used for detecting nitrate in various vegetables.
Discussion

To some extents, the prediction accuracy depends on the proper modelling algorithms and when optimal parameters are determined. Though the FTIR-ATR displayed that the nitrate spectral intensity positively coincided with the nitrate content, the PLS model results were not satisfied, which meant the FTIR-ATR spectra and the nitrate contents was non-linear in the range from 1500 to 1200 cm$^{-1}$. It was probably because other compounds in vegetable included in the range were seemed as interference; another reason might be the nitrate absorption peak split to two peaks at 1401 cm$^{-1}$ and 1350 cm$^{-1}$ at with high concentration were not following linear relationship with nitrate contents [41]. To deal with non-linear modeling, intelligent algorithms have been developed. ELM is a developed algorithm for both classification and regression [41]. The advantages of ELM are extremely fast speed, less human intervenes and great computational scalability [46-47]. To improve the prediction accuracy, the calibration dataset was modified before being calibrated by the Euclidean distance method based on the spectra feature of target sample, which belongs to self-adaptive models, to obtain a similar-sample dataset [33, 40]. Therefore, FTIR-ATR spectra couple with extreme learning machine model with proper parameters showed rapid, accuracy and in-situ measurement results, and it could be potentially used in plant growth management and food safety testing.

Conclusions

In this study, the nitrate content levels in four leafy vegetables, Chinese cabbage, swamp cabbage, celery, and lettuce, were investigated. Portable FTIR-ATR spectroscopic instruments, along with the ED-ELM model, were used to predict the nitrate contents. The unique feature of nitrate was its absorption in the mid-infrared region at 1500-1200 cm$^{-1}$, and the absorption bands at 1401 cm$^{-1}$ and 1350 cm$^{-1}$ were
associated with the $\nu_3$, N–O asymmetric stretch, which was generated by the splitting of the $\nu$ generate mode into two bands labeled $\nu_3$, high and $\nu_3$, low. The peak at 1245 cm$^{-1}$ was associated with nitrite (NO$_2^-$). The results indicated that the nitrate contents in the vegetables exceeded the corresponding WHO-specified maximum tolerance limits (Chinese cabbage: 4063–14104 mg/kg; swamp cabbage: 2111–6607 mg/kg; celery: 805–8643 mg/kg; lettuce: 2567–10485 mg/kg), and these vegetables could only be consumed after being completely boiled. Moreover, the ED-ELM model (with performance indices of $RMSE_p \approx 799.67$ mg/kg, $R^2 \approx 0.93$, and $RPD \approx 2.22$) showed the best performance, compared to that of the ELM ($RMSE_p \approx 995.77$ mg/kg, $R^2 \approx 0.70$, $RPD \approx 1.76$) and PLS ($RMSE_p \approx 1172.01$ mg/kg, $R^2 \approx 0.65$, $RPD \approx 1.66$) models. The results indicated that FTIR-ATR, along with the ED-ELM model method, was a rapid and accurate in situ method to estimate nitrate contents.

**Abbreviations**

ATR, attenuated total reflectance spectroscopy; ED, Euclidean distance; ED-ELM, Euclidean distance-modified intelligent algorithm extreme learning machine model; ELM, extreme learning machine; $E_L M$, Legates and McCabe index; FTIR, Fourier-transform infrared spectroscopy; FTIR-ATR, Fourier-transform infrared attenuated total reflectance spectroscopy; PCA, principal component analysis; PC1, the first principal component; PC2, the second principal component; PC3, the third principal component; PLS, partial least squares; $R^2$, coefficients of determination; $RMSE$, root-mean-square-error; $RMSE_c$, the root-mean-square-error of the calibration dataset; $RMSE_p$, root-mean-square-error of the validation dataset; $RPD$, the ratio of performance to deviation; $WI$, Willmott's index;

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**Notes**

The authors declare no competing financial interest.

**Author’s contributions**

CWD designed and directed the experiment, and made revisions of the manuscript. FM conducted the experiment, processed the data analysis and composed the manuscript. SLZ collected the vegetables, prepared the samples and recorded the spectra data. YXD collected the vegetables, prepared the samples and measured nitrate contents. All authors read and approved the final manuscript.

**Acknowledgements**

Not applicable.

**Competing interests**

All the authors declare that they have no competing interests.

**Availability of data and materials**

The datasets supporting the conclusion of this article are included within the article (additional files).

**Consent for publication**

Not applicable.

**Ethics approval and consent to participate**

Not applicable.

**Funding**

This work was supported by the Strategic Priority Research Program of Chinese Academy of Sciences (XDA23030107) and the Innovational Project in Agriculture
from Jiangsu Province (CX(17)3010).

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Table 1. Nitrate contents in vegetables analyzed by the chemical method

| Sample         | Range  | Mean (mg/kg) | Median (mg/kg) | Standard Deviation |
|----------------|--------|--------------|----------------|--------------------|
| Chinese cabbage | 4063–14104 | 7550         | 7543            | 1664               |
| Swamp cabbage  | 2111–6607  | 4219         | 4196            | 1029               |
| Celery         | 805–8643  | 4164         | 3991            | 1214               |
| Lettuce        | 2567–10485 | 4322         | 4163            | 1035               |
Table 2. The performance parameters of the calibration model

| Models   | Calibration data sets | Hidden layers | Latent variables | $RMSE_c$ (mg/kg) |
|----------|-----------------------|---------------|------------------|------------------|
| ELM      | 900                   | 40            | –                | 1089.91          |
| PLS      | 900                   | –             | 7                | 1087.68          |
| ED-ELM   | 100                   | 100           | –                | 322.52           |
Figure 1. Boxplot of distributions of the nitrate contents of Chinese cabbage, swamp cabbage, celery, and lettuce (306 samples each), following Gaussian distributions.
Figure 2. (a) Characteristic FTIR-ATR absorption spectra of nitrate, (b) the second derivative spectra of nitrate, in the range from 1500 to 1200 cm$^{-1}$, obtained from Chinese cabbage, swamp cabbage, celery, and lettuce.
Figure 3. Principal component (PC1, PC2, and PC3) score plots of the vegetable samples; the four types of vegetables were Chinese cabbage, swamp cabbage, celery, and lettuce.
Figure 4. Statistical distribution of the validation parameters $R^2$, RPD, and RMSEP with various numbers of calibration datasets in the ED-ELM model for the prediction of nitrate content.
**Figure 5.** The ED-ELM model performance indices of (a) the ratio of $RMSE_P$ to $RMSE_C$, RPD, and $R^2$, (b) $WI$ and $E_{LM}$ values for various numbers of hidden neurons from 10 to 90.
Figure 6. Scatterplots of measured nitrate values and prediction results obtained by (a) the ED-ELM model, (b) the ELM model and (c) the PLS model. The dotted line is the reference line (1:1), and the reference line corresponds to the exact prediction. The samples are distributed along the reference line.
Figure 7. Taylor diagram illustrating the parameters and positions for the measured and predicted obtained by the PLS, ELM, and ED-ELM models.
Leafy vegetables \( \rightarrow \) Nitrate molecule \( \rightarrow \) Attenuated total reflectance accessory

Adaptive extreme learning machine \( \rightarrow \) FTIR-ATR spectra of nitrate