A New Local Modelling Approach Based on Predicted Errors for Near-Infrared Spectral Analysis

Haitao Chang,1 Lianqing Zhu,2 Xiaoping Lou,2 Xiaochen Meng,2 Yangkuan Guo,2 and Zhongyu Wang1

1School of Instrumentation Science & Opto-Electronics Engineering, Beihang University, Beijing 100191, China
2Beijing Key Laboratory for Optoelectronic Measurement Technology, Beijing Information Science & Technology University, Beijing 100192, China

Correspondence should be addressed to Lianqing Zhu; zhulianqing@sina.com

Received 10 March 2016; Accepted 7 June 2016

Academic Editor: Karoly Heberger

Copyright © 2016 Haitao Chang et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Over the last decade, near-infrared spectroscopy, together with the use of chemometrics models, has been widely employed as an analytical tool in several industries. However, most chemical processes or analytes are multivariate and nonlinear in nature. To solve this problem, local errors regression method is presented in order to build an accurate calibration model in this paper, where a calibration subset is selected by a new similarity criterion which takes the full information of spectra, chemical property, and predicted errors. After the selection of calibration subset, the partial least squares regression is applied to build calibration model. The performance of the proposed method is demonstrated through a near-infrared spectroscopy dataset of pharmaceutical tablets. Compared with other local strategies with different similarity criterions, it has been shown that the proposed local errors regression can result in a significant improvement in terms of both prediction ability and calculation speed.

1. Introduction

Near-infrared (NIR) spectroscopy plays an important role in the analysis of complex samples or chemical process due to its simplicity, rapidity, and nondestructive measurements [1–4]. As a result, multivariate calibration methods that relate property \(Y\) and spectra \(X\) have been extensively used in the quantitative analysis of NIR spectroscopy. Many authors have stated that the choice of the appropriate calibration method is one of the key factors that influence the performance in prediction of the property \(Y\) of query samples [5].

The multivariate calibration methods, such as multiple linear regression (MLR), principal component regression (PCR) [6], and partial least squares (PLS) [7] regression, have been adopted for the first time [2, 8]. However, these methods are based on statistical linear models which are not always met in real-life situations and therefore are not able to efficiently model the nonlinear relationship between \(Y\) and \(X\). To solve this problem, different authors have demonstrated that nonlinear algorithms such as Artificial Neural Network (ANN) [9] and Least Squares Support Vector Machines (LS-SVM) [10] can produce better results than traditional linear methods especially used together with large NIR spectral libraries [11].

The approach presented in this paper is local method, which has been widely used in near-infrared spectroscopy analysis because of its advantages in terms of the simplicity of the model constructed and the ability to cope with nonlinearities [12–14]. The essential idea of local method is to develop specific calibration subsets spectrally similar to each query sample whose properties are to be predicted and build a calibration model on the selected relevant samples for the query sample.

The key issue in local learning is how the similarity criterion should be constructed. In general, the most commonly used similarity checks in NIR spectroscopy are the Euclidean distance (ED) [15], the Mahalanobis distance (MD) [16], and spectral angle mapper (SAM) [17] distance on the spectral space or principal component space [18]. Moreover,
the computation of the principal components-Mahalanobis (PC-M) [19] distance has become the standard procedure for NIR distance measurements. However, the samples are usually multivariate and influenced by several compositional attributes, which are expressed as highly overlapped and non-specific NIR absorption or reflectance [20]. For this reason, the samples that are very close in X space are frequently not close (or similar) in terms of Y space. Therefore, traditional similarity criterion uses only X information to select relevant samples, which may result in a waste of Y information and inaccurate sample selection [21]. In order to overcome such problem, some supervised or semisupervised methods have been developed, which take account of information of both X and Y. Recently, another reliable similarity estimation called supervised locality preserving projection (SLPP) [22] has been successfully used in local approach to enhance the similarity measurement accuracy [21, 23].

As a simple linear approximation technique, local method could address the nonlinearity through the locally linear assumption that the samples that are most close in the property space are very similar in terms of spectra space. Thus, the SLPP is used in this study, which not only selects most similar sample considering both spectra and property information but also reduces the computational load.

The rationale behind this approach is based on the assumption that the samples that are most close in the property space are very similar in terms of spectra space. Given a set of N-dimensional calibration spectral data \( X_C = \{x_1, x_2, \ldots, x_n\} \) in \( \mathbb{R}^N \) with corresponding property set \( Y_C = \{y_1, y_2, \ldots, y_n\} \), N-dimensional prediction spectral data \( X_P = \{x_1, x_2, \ldots, x_m\} \) \( \in \mathbb{R}^N \) with corresponding property set \( Y_P = \{\hat{y}_1, \hat{y}_2, \ldots, \hat{y}_m\} \). Here \( X_C \) is \( n \times N \) matrix containing N spectral responses of \( n \) samples; \( X_P \) is \( m \times N \) prediction matrix containing \( N \) spectral data response of \( m \) samples; \( Y_C \) is \( n \times l \) matrix; and \( m \times l \) matrix \( \hat{Y}_P \) is the predicted properties of prediction set with global PLS regression, \( l \) being the number of components. Let spectra set \( X_{(n+m)} \) \( \in \mathbb{R}^{(n+m) \times N} = \{X_C \cup X_P\} \). The algorithmic procedure for the calibration subset selection is stated as follows:

1. Constructing a neighborhood graph as follows:
   - K Nearest Neighbors. The ith and the jth samples are connected by an edge if the jth sample is among K nearest neighbors of the ith sample or the jth sample is among K nearest neighbors of the ith sample. Here the distance between the samples is calculated in the property space.

2. Computing the Weights. \( W \) is a sparse symmetric \((n+m) \times (n+m)\) matrix with \( W_{ij} \) having the weight of the edge joining the ith and the jth samples, and \( W_{ii} = 0 \) if there is no such edge, and \( W_{ij} = 1 \) if and only if the ith and the jth samples are connected by an edge.

3. Finding the Basis Vectors of the Subspace. This step aims at finding a transformation matrix \( A_{N \times d} \) to project spectra set X to a low-dimensional set \( Z_{(n+m) \times d} = \{z_1, z_2, \ldots, z_{n+m}\} \) in \( \mathbb{R}^d \) \( (d \ll N) \). Suppose that there exists a linear transformation \( Z = Xa^T \), where a is the basis vector. The basis vector a is computed by solving the following minimization problem:

\[
\text{Min } aX^TWXa^T
\]

Subject to \( aX^TXa^T = 1 \).
It is observed that the minimization problem can be calculated by solving the following generalized eigenvalue problem:

\[ X^T L X a^T = \lambda X^T D X a^T, \quad (2) \]

where \( D \) is a diagonal matrix whose entries are column sums of \( W \) and \( D_{ii} = \sum_j W_{ij}; L = D-W \) is the Laplacian matrix. Let the column vectors \( a_1^T, \ldots, a_d^T \) be the solutions to (2) which are ordered according to their eigenvalues, \( \lambda_1 < \cdots < \lambda_d \). Thus, transformation matrix and low-dimensional matrix can be calculated as follows: \( A = [a_1^T, \ldots, a_d^T] \), \( Z_C = X_C A \), and \( Z_Y = X_Y A \).

Finally, for each query sample, the most similar sample will be selected according to the ED in the low-dimensional space \( Z_C \).

2.2. Partial Least Squares Regression. The PLS regression has been extensively employed to obtain a quantitative model for prediction of analytes on spectral data. In this paper, the PLS is used in the procedure of local selection and the establishment of regression model. In addition, the critical step is the determination of the number of factors for achieving the best prediction. Generally, the optimum PLS factors can be determined by minimizing the prediction error of cross-validation groups. However, the cross-validation is time-consuming and does not consider the information from the query sample. For these reasons it would not be a desirable choice to local method. In this paper, the selection of calibration subset is based on the prediction error with global PLS, which are expressed as highly linear between samples in subset. Consequently, the PLS factor would have little effect on the performance of calibration models. Here the PLS factor fixed at a constant value by minimizing the root mean squared error of prediction (RMSEP) of validation dataset and the experimental verification is presented in Section 4.1.

The performance of the final calibration models was evaluated in terms of the RMSEP, Residual Predictive Deviation (RPD), and correlation coefficient (\( R^2 \)) in prediction set. It should be noted that RPD is the ratio between the standard deviation of the reference data and RMSEP of prediction.

2.3. Other Algorithms. In order to compare the predictive performance of our local errors method with other local approaches, the following similarity criterions were used: ED, angle distance, PC-M distance, and supervised or semisupervised methods. A brief description of these criterions is given as follows.

2.3.1. Euclidean Distance. The ED between query sample \( x_q \) and calibration sample \( x_i \) is given by

\[ d_E(x_q, x_i) = \sqrt{(x_q - x_i)^T (x_q - x_i)}, \quad (3) \]

where \( T \) is the transpose operation, \( x_q \in X_P \), and \( x_i \in X_C \).

2.3.2. Angle Distance. In this method, the Cosine is used to evaluate the similarity between query data \( x_q \) and \( x_i \) in the spectral space, and the angle distance is defined as

\[ d_{\text{angle}}(x_q, x_i) = \frac{x_q \cdot x_i}{\sqrt{(x_q \cdot x_q)(x_i \cdot x_i)}}. \quad (4) \]

2.3.3. Mahalanobis Distance in the Principal Component. The PC-M distance is obtained through computing the Mahalanobis distance (MD) between samples in spectral principal component space. In this case, the appropriate number of principal components is determined by minimizing the root mean square of the compositional differences between property \( Y \) and predicted value \( \hat{Y} \) of samples in calibration set. The Mahalanobis distance between query sample \( x_q \) and calibration sample \( x_i \) is defined as

\[ d_M(x_q, x_i) = \sqrt{(x_{q\text{PC}} - x_{i\text{PC}})^T V^{-1} (x_{q\text{PC}} - x_{i\text{PC}})} \]

where \( V \) is the covariance matrix in spectral principal component space of calibration set and \( x_{q\text{PC}} \) and \( x_{i\text{PC}} \) are spectra of query sample and calibration sample in spectral principal component space. For details, see [20].

2.3.4. Supervised or Semisupervised Methods

(1) Euclidean Distance Based on Both Spectra and Property. In this method, both spectra \( X \) and property \( Y \) are used to evaluate the similarity between query data \( x_q \) and \( x_i \) in the calibration dataset, and the similarity value \( d_{XY} \) is defined:

\[ d_{XY}(x_q, x_i) = d_E(x_q, x_i) \cdot (1 - \gamma) + d_E(\hat{Y}_q, \hat{Y}_i) \cdot \gamma, \quad (6) \]

where \( \gamma \) is a weight parameter to balance the importance of spectra \( X \) and property \( Y \), \( 0 \leq \gamma \leq 1 \), and \( d_{XY} \) is the Euclidean distance between \( x_q \) and \( x_i \).

(2) Euclidean Distance in the Low-Dimensional Space. Here, the similarity measurement considers both \( X \) and \( Y \) information and utilizes SLPP technique to select relevant samples. For details, see [21].

In summary, the detailed implementation of local errors regression is described as follows.

Step 1. Compute the predicted properties of prediction set \( \hat{Y}_P \) and predicted properties of calibration set \( \hat{Y}_C \) with global PLS regression.

Step 2. For each query sample, select the most relevant sample from calibration set with SLPP.

Step 3. Predefine parameters such as PLS factors and error ranges by minimizing the RMSEP of validation dataset. Here PLS factors are set to 2, 3, 4, or 5 and error range is specified as 1.5. A much detailed description will be given in Section 4.1.

Step 4. For each query sample \( y_q^d \) calibration subset is selected based on predicted error similarity criteria, and the
### Table 1: Descriptive statistics for the calibration set, validation set, and prediction set.

| Sample sets     | Number | Range          | Mean  | Standard deviations |
|-----------------|--------|----------------|-------|---------------------|
| Calibration     | 460    | 154.3–237.7    | 188.4 | 15.8                |
| Validation      | 40     | 168.2–219.5    | 194.8 | 12.4                |
| Prediction      | 155    | 151.6–239.1    | 192.9 | 22.0                |

### 3. Materials and Experiment

The dataset was obtained from the web of http://software.eigenvector.com/Data/tablets/index.html. The spectra were recorded by Instrument I of Foss NIR Systems 6500 Spectrometer. It contains 655 transmittance NIR spectra of pharmaceutical tablets, measured between 600 and 1898 nm with a 2 nm sampling step. The initial dataset was split into a calibration set of 460 samples and prediction set of 155 samples and validation set of 40 samples. Table 1 showed the statistical results of pharmaceutical tablets. All sets have similar averages and standard deviations which indicate that all the datasets can be used to represent the main variability of pharmaceutical tablets. A microcomputer (Lenovo) with an Intel Core 2 processor was used for all the calculations. All the algorithms were implemented using MATLAB 2012b. The raw NIR spectra of pharmaceutical tablets were presented in Figure 1.

### 4. Results and Discussion

#### 4.1. Parameter Setting

The determination of the PLS factors and error ranges is an essential step in this study, which decides the accuracy of the model. Here these parameters were determined with the most similar sample for each query sample. Therefore, in Step 2, parameters $K$ and $d$ are set as 50 and 20, respectively. Such a selection is based on the authors’ experience without further discussion.

#### Step 5. Build PLS calibration model on the selected relevant samples and predict each query sample.

It should be noted that the number of nearest neighbors $K$ and dimension of transformation matrix $d$ had little effect on the selection of the most similar sample for each query sample. Therefore, in Step 2, parameters $K$ and $d$ are set as 50 and 20, respectively. Such a selection is based on the authors’ experience without further discussion.

#### 4.2. Size of Calibration Subset

The final results of the local errors strategy for each query sample in prediction set are summarized in Table 2. It can be seen that each number of selected samples in calibration subset is smaller than 210, which is less than the half size of calibration set. Moreover, 78.7% sizes of calibration subsets are between 13 and 50. That means that the computational complexity of the prediction model can be reduced a lot compared with global method. It is noted that here the minimum size of the subset is set...
Figure 2: Variation of RMSEP with the PLS factors for validation set. RMSEP = root mean squared error of prediction. PLS = partial least squares.

Figure 3: The probability of insufficient selected samples with the PLS factors and error ranges for the validation set. × = insufficient selected samples for building calibration model. PLS = partial least squares.
to 13 so as to avoid insufficient selected samples for building calibration model. When the number of selected samples is not reaching 13, the calibration subset has been made up of 50 samples which are the spectral similarity to query sample.

4.3. Comparison. Figure 4 is the scatter plot that shows a correlation between reference values and NIR prediction values in the prediction set using global PLS method and proposed method. According to the scatter plots, the predicted values associated with local errors regression best match the reference ones. It can thus be concluded that the proposed method maintains good precision compared with global PLS method. Furthermore, the method proposed here was compared with other local methods.

The performance of local errors regression, global PLS, and other local methods with different similarity criterions was listed in Table 3. In each local-PLS model, two parameters have to be optimized: size of calibration subset and PLS factors. In this study, the number of calibration subset samples included was evaluated at 30, 50, 100, 150, 200, and 250 samples and checked so as to find which value for this parameter is better. The optimum number of PLS factors was identified by minimizing the prediction error of cross-validation groups. Best prediction results from local-PLS approaches with different similarity criterions are listed in the table. On the contrary, the PLS factors used in the proposed method, as discussed in the previous section, was fixed at either 2 or 3 or 4 or 5.

Figure 4: Predicted versus reference values of prediction set for the local errors regression and global method.

| Method                      | Similarity criterion | RMSEP | $R^2$ | RPD | Size of subset | Parameters | Time consumption (s) |
|-----------------------------|----------------------|-------|------|-----|----------------|------------|---------------------|
| Global                      | —                    | 4.30  | 0.96 | 5.11| 460            | —          | 4.4                 |
| ED                          |                      | 4.18  | 0.96 | 5.26| 150            | —          | 175.4               |
| Cosine                      |                      | 4.25  | 0.96 | 5.17| 150            | —          | 179.1               |
| Other local methods         |                      |       |      |     |                |            |                     |
| PC-M                        |                      | 4.21  | 0.96 | 5.22| 50             | PC factors = 10 | 53.5               |
| $X + Y + ED$                |                      | 4.24  | 0.96 | 5.18| 100            | $y = 0.8$  | 123.1               |
| $X + Y + SLPP$              |                      | 4.27  | 0.96 | 5.15| 200            | $y = 0.8, d = 20$ | 233.9               |
| Local errors regression     | Errors + ED          | 3.21  | 0.98 | 6.85| 13–205         | $d = 20$   | 9.5                 |

ED: Euclidean distance; PC-M: Principal components-Mahalanobis distance; $X + Y + ED$: Euclidean distance considering both spectra $X$ and property $Y$; $X + Y + SLPP$: Euclidean distance in the low-dimensional space obtained with supervised locality preserving projection method; errors + ED: Euclidean distance between predicted errors; RMSEP: root mean squared error of prediction; $R^2$: correlation coefficient in prediction set; RPD: residual prediction deviation; PC factors: Principal component factors; symbol $y$: a trade-off parameter to balance the importance of spectra $X$ and property $Y$; $d$: dimension of transformation matrix; and s: second.
In terms of RMSEP, with the same spectra data the prediction performance of the local methods is better than that of global method, showing lower RMSEP values. Moreover, the RMSEP obtained with the proposed method is the lowest compared to the best values obtained with other local methods. Furthermore, Table 3 also shows that the difference of $R^2$ with different methods is not significant and all the values of $R^2$ are approximated to 1. The RPD is proportional to the RMSEP and with a similar variation tendency. Another important issue is the prediction speed. In this context all the local methods were slower than global method, showing higher time consumption. As shown in Table 3, global method requires around 4.4 s to calculate all the samples in prediction set and local methods are much more time-consuming, from 53.5 s to 233.9 s depending on subset size. The reason is that the local methods will select different calibration subset and perform new calibration models specifically to different query samples. Thus, in the proposed method, instead of cross-validation, the fixed PLS factor strategy is adopted, which resulted in a noticeable improvement of calculation speed, showing a lower time consumption about 9.5 s.

5. Conclusion

In this paper, a new local method for nonlinear spectra is proposed. The main contributions include a new similarity measurement utilizing ED of predicted errors, a selection method extended from SLPP aiming to search the most relevant sample for each query sample and fixed PLS factors prediction model without cross-validation. Also, a comparison has been carried out in this study, to compare the predictive performance of the proposed method, global PLS, and other local strategies with different similarity criterions such as ED, Cosine, PC-M, $X + Y + ED$, and $X + Y + SLPP$ by using the pharmaceutical tablets dataset. The prediction results have demonstrated that the proposed method could bring higher calibration performances and lower time consumption.

Competing Interests

The authors declare that they have no competing interests.

Acknowledgments

The authors would like to thank the Beijing Municipal Natural Science Foundation (KZ2001010772032), National Defense Basic Scientific Research Program of China (J132012C001), and Program for Changjiang Scholars and Innovative Research Team in University, PCSIRT (IRT1212), for their financial support.

References

[1] H. Ebrahimi-Najafabadi, R. Leardi, P. Oliveri, M. Chiara Casolino, M. Jalali-Heravi, and S. Lanteri, “Detection of addition of barley to coffee using near infrared spectroscopy and chemometric techniques,” Talanta, vol. 99, pp. 175–179, 2012.

[2] Y. Roggo, P. Chalus, L. Maurer, C. Lema-Martinez, A. Edmond, and N. Jent, “A review of near infrared spectroscopy and chemometrics in pharmaceutical technologies,” Journal of Pharmaceutical and Biomedical Analysis, vol. 44, no. 3, pp. 683–700, 2007.

[3] H. Nakagawa, T. Tajima, M. Kano et al., “Evaluation of infrared-reflection absorption spectroscopy measurement and locally weighted partial least-squares for rapid analysis of residual drug substances in cleaning processes,” Analytical Chemistry, vol. 84, no. 8, pp. 3820–3826, 2012.

[4] H. Tran, P. Salgado, E. Tillard, P. Dardenne, X. T. Nguyen, and P. Lecomte, “‘Global’ and ‘local’ predictions of dairy diet nutritional quality using near infrared reflectance spectroscopy,” Journal of Dairy Science, vol. 93, no. 10, pp. 4961–4975, 2010.

[5] F. Estienne, L. Pasti, V. Centner et al., “A comparison of multivariate calibration techniques applied to experimental NIR data sets: part II. Predictive ability under extrapolation conditions,” Chemometrics and Intelligent Laboratory Systems, vol. 58, no. 2, pp. 195–211, 2001.

[6] M. Praisler and S. Ciochina, “Global clustering quality coefficient assessing the efficiency of PCA Class Assignment,” Journal of Analytical Methods in Chemistry, vol. 2014, Article ID 342497, 12 pages, 2014.

[7] S. E. Barnes, M. G. Sibley, H. G. M. Edwards, and P. D. Coates, “Process monitoring of polymer melts using in-line spectroscopy,” Transactions of the Institute of Measurement and Control, vol. 29, no. 5, pp. 453–465, 2007.

[8] M. Clavaud, Y. Roggo, R. Von Daeniken, A. Liebler, and J.-O. Schwabe, “Chemometrics and in-line near infrared spectroscopic monitoring of a biopharmaceutical Chinese hamster ovary cell culture: prediction of multiple cultivation variables,” Talanta, vol. 111, no. 3, pp. 28–38, 2013.

[9] P. Roestorstorf and J. Fohlman, “Proposal for a common nomenclature for sequence ions in mass spectra of peptides,” Biomedical Mass Spectrometry, vol. 11, no. 11, p. 601, 1984.

[10] J. A. K. Suykens, T. V. Gestel, J. D. Brabanter, B. D. Moor, and J. Vandewalle, “Least-squares support vector machines,” Euphytica, vol. 2, no. 2, pp. 1599–1604, 2002.

[11] D. Pérez-Marín, A. Garrido-Varo, and J. E. Guerrero, “Nonlinear regression methods in NIRS quantitative analysis,” Talanta, vol. 72, no. 1, pp. 28–42, 2007.

[12] D. Pérez-Marín, A. Garrido-Varo, and J. E. Guerrero, “Implementation of LOCAL algorithm with near-infrared spectroscopy for compliance assurance in compound feedingstuffs,” Applied Spectroscopy, vol. 59, no. 1, pp. 69–77, 2005.

[13] D. Pérez-Marín, T. Fearn, J. E. Guerrero, and A. Garrido-Varo, “Improving NIRS predictions of ingredient composition in compound feedingstuffs using Bayesian non-parametric calibrations,” Chemometrics and Intelligent Laboratory Systems, vol. 110, no. 1, pp. 108–112, 2012.

[14] J. S. Shenk, P. Berzaghi, and M. O. Westerhaus, “Investigation of a LOCAL calibration procedure for near infrared instruments,” Journal of Near Infrared Spectroscopy, vol. 5, no. 4, pp. 223–232, 1997.

[15] S. Kim, R. Okajima, M. Kano, and S. Hasebe, “Development of soft-sensor using locally weighted PLS with adaptive similarity measure,” Chemometrics and Intelligent Laboratory Systems, vol. 124, pp. 43–49, 2013.

[16] E. Zamora-Rojas, A. Garrido-Varo, F. Van den Berg, J. E. Guerrero-Ginel, and D. C. Pérez-Marin, “Evaluation of a new local modelling approach for large and heterogeneous NIRS.
data sets,” *Chemometrics and Intelligent Laboratory Systems*, vol. 101, no. 2, pp. 87–94, 2010.

[17] C. Cheng and M.-S. Chiu, “A new data-based methodology for nonlinear process modeling,” *Chemical Engineering Science*, vol. 59, no. 13, pp. 2801–2810, 2004.

[18] F. Gogé, R. Joffre, C. Jolivet, I. Ross, and L. Ranjard, “Optimization criteria in sample selection step of local regression for quantitative analysis of large soil NIRS database,” *Chemometrics and Intelligent Laboratory Systems*, vol. 110, no. 1, pp. 168–176, 2012.

[19] L. Ramirez-Lopez, T. Behrens, K. Schmidt, R. A. V. Rossel, J. A. M. Dematté, and T. Scholten, “Distance and similarity-search metrics for use with soil vis-NIR spectra,” *Geoderma*, vol. 199, pp. 43–53, 2013.

[20] L. Ramirez-Lopez, T. Behrens, K. Schmidt, A. Stevens, J. A. M. Dematté, and T. Scholten, “The spectrum-based learner: a new local approach for modeling soil vis-NIR spectra of complex datasets,” *Geoderma*, vol. 195-196, pp. 268–279, 2013.

[21] K. Chen, J. Ji, H. Wang, Y. Liu, and Z. Song, “Adaptive local kernel-based learning for soft sensor modeling of nonlinear processes,” *Chemical Engineering Research and Design*, vol. 89, no. 10, pp. 2117–2124, 2011.

[22] F. Dornaika and A. Assoum, “Enhanced and parameterless locality preserving projections for face recognition,” *Neurocomputing*, vol. 99, pp. 448–457, 2013.

[23] K. He, H. Cheng, W. Du, and F. Qian, “Online updating of NIR model and its industrial application via adaptive wavelength selection and local regression strategy,” *Chemometrics and Intelligent Laboratory Systems*, vol. 134, no. 8, pp. 79–88, 2014.