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Performance Evaluation of Chemometric Prediction Models—Key Components of Wheat Grain

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The present study was aimed to evaluate the accuracy of using near-infrared spectroscopy (NIRS) for predicting protein, moisture, starch and ash content values of wheat. The physiochemical properties of wheat were predicted using twelve prediction models of preprocessing coupled with regression tools. The performance measure of SVM aided with extended multiplicative scatter correction gave confident prediction results of protein, moisture, ash and starch content with R^2 values of 0.989, 0.987, 0.976, 0.998 and RMSECV values of 0.263, 0.285793, 0.369 and 0.03 respectively. These results indicate the practical applicability of NIRS in wheat grain quality profiling.

Keywords: Wheat, Support vector machine, Quality parameters, Near Infrared Spectrometer

Introduction

Wheat crop adapts to changes in climatic conditions and thus is a preferred cereal crop¹. Wheat and wheat based products are used in everyday consumables and also in the preparation of alcoholic drinks, cattle feed to name a few. The success of final wheat product has a direct dependability on the physiochemical properties of the grain². Traditionally, quality tests for wheat grain and flour have been undertaken by lengthy chemical and rheological methods, which are labor intensive, time-consuming, and requires experienced technicians. NIR technology ascertains speedy, non destructive, qualitative and quantitative analysis to facilitate the millers/buyers and sustain complete confidence in the grain they choose to buy.

NIR Instrumentation

The near infrared spectroscopy is an instrumental technique based on the absorption rate of the infrared radiation reflected or transmitted by the sample, due to the vibration of hydrogen bonds (CH, NH, OH). The wheat grains are of complex composition with numerous hydrogen bonds whose vibration contributes to overlapped spectral bands. Assignment of a specific spectral wavelength to a chemical constituent is not simple through direct relationship which is where chemometrics comes to the rescue³.

NIRS is a secondary analytical method which needs to be backed up with reference data. Prediction ability is based on the relationship modeled between the measured spectral data and the constituent of interest. Chemometric techniques, such as Partial least squares (PLS) regression, Support vector machine(SVM), Multiple linear regression(MLR) and other such calibration model building methods can be used to identify and quantify the key wheat constituents. The NIR instrument used to collect spectral data of whole grain samples was FOSS XDSTM NIR Analyzer, which has a wide wavelength range (450–2,498 nm) including the visible spectrum. The characteristic absorption peaks were marked at 990nm, 1440nm, 1940nm, 2100nm, 2336nm and absorption bands from 2168 to 2180 nm and 973 to 1020 nm in the NIR spectrum based on intensive literature survey and prior work⁴. The amount of near infrared absorption is related to the presence of a few key components in wheat grains⁵. The aim of this study was to apply chemometric models on NIR spectroscopic data to predict the quantity of protein, moisture, starch and ash which are a few vital features that depicts a characteristic profile of wheat⁶. The quantitative analysis of these parameters can further be used to segregate different varieties of wheat.

Wheat Quality Analysis

Moisture content is a key stricture of any grain⁷ which affects its sale weight, quality and ability to be

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stored long-term. The ability of wheat flour to be processed into different food merchandise is largely determined by the protein content⁸. Mature wheat grains contain 8% to 20% proteins. About 75-85% of the total grain proteins is composed of glutenins and gliadins. In common wheat, these parameters contributes to elasticity during the dough formation⁹. Starch content in wheat grain contains 66–76% carbohydrates¹⁰. It is not merely an important part of our nutrition but also dictates the textural properties of many food products, wherein baked items can be mentioned in specific¹¹. The measure of ash content relates to the grade of flour¹². Ash content indicates milling performance by indirectly revealing the amount of bran contamination in flour. Ash in flour can affect color, imparting a darker color to finished products.

Materials and methods

20 varieties of wheat grain (Triticum aestivum) cultivars like pbw17, ankurkedar, pbw502, sonalika, to name a few that were grown in different locations of India for a period of two years were collected and stored. Severely wrinkled, discolored, and split grains were manually removed from the sample. Five sample sets from each variety were chosen as carefully selected and representative units, which have in turn been quality profiled using analytical reference standards. A total of 100 spectra were collected from the sample set. The procedure flow chart for the measurement of key constituents of wheat grain is illustrated in Figure1. Prediction models were developed for each quality parameter using PLS, MLR and SNV. Statistics used to determine the ability of NIR to predict specific parameters were the R^2 ; coefficient of determination¹³ and RMSECV; the Root mean squared error of crossvalidation which is a statistical measure of the proximity of the data to the fitted regression line.

Partial least square Regression (PLS)

After the first step of preprocessing the spectral signal X, the newly built test data matrix was employed in the PLS calculation using the expression¹⁴

$$y = Xb + e = \sum_{i=1}^{A} (t_i p_i^T) * b + e \qquad \dots (1)$$

Where y is the predicted response, b is the regression coefficient, t_i and p_i are the scores and loadings vectors used for training the model, A is the number of Latent variables which indicates the length of test data and e is the residual vector.

Multiple Linear Regression (MLR)

Linear regression with more than one variable is believed to enhance the estimation of new predictor variable. For the model performance with multiple variables, the cost function to fit the best straight line in the data was calculated from the expression

$$J(\theta_0, \theta_1, \dots, \theta_n) = \frac{1}{2m} \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)})^2 \quad \dots (2)$$

Where m is the amount of features, h is predictive value and y is real world value.

The algorithms that minimized the cost function were gradient descent and normal equations¹⁵.

Support Vector Machine (SVM):

The basic idea behind SVM is to construct a hyper plane that lies in close proximity to as many of the data points as possible¹⁶. The hyperplane (decision boundary) can be expressed as

$$wT x + b = 0$$
 ... (3)



Fig. 1 — The flow diagram of calibration model development

	RMSECV of						R^2 of						
Prediction model	Moisture	Protein		tarch	Ash		Moisture	Pro	Protein		Ash		
SG 1st der with SVM	0.258	0.244	0.369		0.032		0.989	0.989		0.976	76 0.999		
SG 1st der with MLR	0.524	0.302	0.402		0.097		0.961	0.986		0.977	0.998		
SG 1st der with PLS	0.299	0.303	0.393		0.097		0.988	0.	985	0.978	C	0.998	
SNV with SVM	0.229	0.243	0.367		0.030		0.989	0.989 0.9		0.976		0.999	
SNV with MLR	1.302	0.511	0.369		0.096		0.781	0.781 0.9		0.980	0.998		
SNV with PLS	0.290	0.414	0.404		0.095		0.987	0.977		0.978		.998	
MSC with SVM	0.260	0.265	0.369		0.029		0.989	0.988		0.976 (.999	
MSC with MLR	0.531	0.318	0.455		0.096		0.961	0.982		0.967	67 0.998		
MSC with PLS	0.336	0.347	0.426		0.096		0.982	0.977		0.969	0.998		
EMSC with SVM	0.263	0.286	0.369		0.030		0.989	0.987		0.975	0.999		
EMSC with MLR	0.341	0.315	0.402		0.097		0.977	0.985		0.976	0.998		
EMSC with PLS	0.289	0.298	0.405		0.097		0.988	0.987		0.977	0.999		
Table 2 — RMSECV	and \mathbf{R}^2 of the	different	predict	ion mod	els for th	e four co	nstituents	under co	nsiderati	on namely	Moisture,	Protein,	
Starch and Ash													
Preprocess +	SG +	SG +	SG +	SNV +	SNV+	SNV +	MSC +	MSC +	MSC+	EMSC +	EMSC +	EMSC +	
Prediction	SVM	MLR	PLS	SVM	MLR	PLS	SVM	MLR	PLS	SVM	MLR	PLS	
Constituents Chemica	ıl result	result Predicted result											
Protein 1	3 12.85	12.87	12.89	12.61	12.59	12.65	12.85	12.93	12.99	12.62	12.73	12.8	
Carbohydrate 70.	18 71.05	70.79	70.84	71.05	70.72	70.68	71.06	71.05	71.05	71.05	70.9	70.87	
Moisture 10	.4 10.28	9.76	10.39	10.3	10.54	10.47	10.27	9.72	10.2	10.27	10.23	10.33	
Ash 1 (57 1 59	16	1 59	1 63	16	1 59	1 58	16	1 59	1 56	16	1 59	

Table1 — The validation result of the wheat grain variety: UP 262 under 12 prediction model combinations of preprocessing and regression model

Where x is the input vector, w and b are coefficients. Wheat spectrum is not a perfectly linear separable data, thus mapping it into a hyperplane was formulated as

$$\min_{\mathbf{w},\mathbf{b},\mathbf{\epsilon}} \frac{\mathbf{w}^{\mathrm{T}}\mathbf{w}}{2} + C\sum_{i} \varepsilon_{i}, \qquad \dots (4)$$

While $yi(wT xi + b) \ge 1-\varepsilon_i$ and $\varepsilon_i \ge 0$ determines the trade-off between the flatness of the function and the tolerable signal error, ε - the slack variables introduced as $\varepsilon_i \ge 0$ for each xi. The prediction results produced with SVM on NIR signals were greatly reassuring.

Results and Discussion

Savitzky-Golay first-derivative (SG), Standard normal transformation (SNV), Multiplicative scattering correction (MSC), extended multiplicative signal correction (EMSC) were the different preprocessing methods applied on the raw spectrum. These preprocesses were combined with different prediction models like SVM, PLS and MLR. Accuracy, defined as the conformance of a measured value and the true value, in the case of the NIRS method, it is expressed as a measure of nearness of the NIRS predicted value with the reference (wet chemistry method). SG, SNV, MSC and EMSC were the different preprocessing methods applied on the spectrum. These preprocesses were combined with different prediction models like SVM. PLS and MLR. Twelve different models from these combinations were validated. The validation result of the wheat grain variety: UP 262 under all the twelve prediction model combinations is displayed in the Table 1. 60 spectral signals of the calibration model were worked with all the 12 prediction models and the Model performance was defined in terms of the multivariate coefficient of determination (\mathbf{R}^2) , the standard error of prediction between the measured and modeled values as listed in Table 2. All the preprocessing and regression model combinations performed confidently well with the RMSECV values in the range of 0.091 to 1.3 and R^2 coefficient values in the range of 0.78 to 0.998. EMSC provided the best prediction ability while SVM displayed an improved performance on nonlinear data, thus an overall enhanced prediction capability was observed when EMSC was combined with SVM. The success is presumably due to the ability of spectral modeling to separate chemical light- absorbance and physical light scatter effects



Fig. 2 — The prediction of moisture, protein, starch and ash content using SVM and EMSC.

effectively. Figure 2 shows the plot of reference value against the predicted value of all the 20 wheat varieties.

Conclusion

Chemometric methods were applied in the current study to extract information from the spectra for the total Moisture, Protein, Starch and Ash by analyzing the influence of absorption at particular wavelength of the wheat grain samples. Protein rich varieties are more suitable for baking products, high desirable constituents of wheat grain can be chosen as breeding cultivars to produce higher nutritional values; such are the outcomes of the quantitative analysis of the grain composition. Calibration development and evaluation were performed using MLR, PLS and SVM regression techniques wherein the R² and RMSECV values can be fine tuned with a larger sample number. Also many more constituents of the wheat grain can be predicted to provide a complete profiling of the wheat sample. Through the convincing results obtained from the quantitative analysis the corresponding spectral wavelength fingerprinting can be a confidently marked for future work.

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