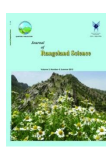


Contents available at ISC and SID

Journal homepage: www.rangeland.ir



Full Length Article:

Potential of Near-Infrared Reflectance Spectroscopy (NIRS) to Predict Nutrient Composition of *Bromus tomentellus*

Hossein Arzani^A, Anvar Sour^B, Javad Motamedi^C

Manuscript Received: 04/11/2012 Manuscript Accepted: 12/12/2012

Abstract. Determination of forage quality of available species is one of the fundamental factors for the management of rangelands. Near-Infrared Reflectance Spectroscopy (NIRS) was used to analysis the Nitrogen (N), Acid Detergent Fiber (ADF), Dry Matter Digestibility (DMD) and Metabolizable Energy (ME) content of three phenological stages (vegetative, flowering and seeding) of Bromus tomentellus samples in grazing pastures of Iran. The sample set consisted of 40 samples for calibration and 23 samples for validation was used to prediction N, ADF, DMD and ME, separately. The samples were measured by reflectance NIR in a 950-1650 nm range. Calibration models between chemical data and NIR spectra were produced using the method of Partial Least Squares (PLS). The coefficients of determination (R2) and standard error of cross validation (SECV) were 0.94 (SECV: 0.208%), 0.98 (SECV: 1.76%), 0.98 (SECV: 1.97%), and 0.97 (SECV: 0.34%) for N, ADF, DMD and ME, respectively. The results obtained from this study indicated that NIRS had a potentiality to be used in predict the N, ADF, and the estimated DMD and ME of forage samples content.

Key words: Range management, Animal nutrition, N, ADF, DMD, ME, *Bromus tomentellus*, Near-Iinfrared Reflectance Spectroscopy (NIRS).

^AProfessor, Faculty of Natural Resources, University of Tehran, Iran.

^BM.Sc. Student of Range Management, From Natural Resources Faculty University of Tehran, Iran.

⁽Corresponding Author). Email: anvarsour@ut.ac.ir

^CAssistant Professor of Department of Natural Resources, University of Urmia, Iran.

Introduction

One of the main objectives of range management is livestock production, which depends on great extent on the nutritive value of available forage (Stoddart et al., 1975). Ganskopp & Bohnert (2003) proposed that livestock and wildlife managers must be aware of the nutritional dynamics of forages to satisfactory growth sustain reproduction of their animals, and assure fair value of the pasture. Valentine (1990) believed that the nutrient balance of animals, whether grazing or penfed is dependent upon four basic factors including the animal's nutrient requirements, nutrient content of the feedstuffs consumed and the amount consumed. Knowledge of generalized nutrient trends in the forage plants available to grazing animals will assist in achieving their most timely utilization and help predict nutrients decencies, and suggest supply maintain Knowledge of nutritional quality of the forage for maintaining animal health requires forage quality analysis and monitoring for proper feed rationing development (Calderon et al., 2009). The Bromus tomentellus is a stable species with cold season grazing value and cluster biological form. It is a palatable species which is consumed by all classes of livestock, particularly sheep. Forage greatly in will vary chemical nutritional composition and value Traditionally (Murray. 1986). chemical analyses have been used to characterize forages, and to predict their value. These nutritive are consuming, costly and in some cases hazardous chemicals are involved (Kokaly & Clark, 1999; Graeff et al., 2001; Li et al., 2006). Forage analysis with NIRS was first reported in 1976 (Norris et al., 1976). Near Infrared Reflectance (NIR) has become widely recognized as a valuable tool in the accurate determination of the chemical composition of a wide range of forages

(Murray, 1993; Shenk & Westerhaus, 1994). NIRS technology is based on the fact that the major organic chemical components of a sample have near infrared absorption properties in the region 700-2500 nm allowing the instantaneous prediction of the nutritive value of feeds and forages (Garrido, 1997). This technique requires consistent sample handling and a calibration method based in multivariate analysis, which converts spectral (NIR absorption) laboratory reference information (Alomar et al., 1999; Deaville & Flinn, 2000; Cozzolino et al., 2000, 2003; Roberts et al., 2004). Several authors have tested NIR to estimate Forage Nutrient Content (Starks et al., 2004; Andrés et al., 2005; Scholtz et al., 2009; Charehsaz et al., 2010).

The objective of this study was to assess the potential of the NIRS technique to predict the N, ADF, DMD and ME contents of *Bromus tomentellus* species.

Materials and Methods

A total of 63 samples of *Bromus* tomentellus were collected at 5 localities (sites) grazing pasture of Iran [West Azarbaijan, East Azarbaijan, Ardabil, Zanjan and Isfahan]. Samples were collected from three phenological stages (vegetative, flowering and seeding stages) with three replications. The range of mean annual precipitation in above regions is 326.2-590 mm and its climates located in cold semi arid with moderate summers and cold winters. The soil texture of these regions was combination of moderate to semi deep and was sandy and clay-loam texture.

The data of chemical analysis provided by Arzani *et al.* (2011) was used to compare NIR results with laboratory methods [Nitrogen was measured using the Kjeldhal technique (AOAC, 1995), Acid Detergent Fiber (ADF) was measured using the procedure described by Van Soest (1963). Dry matter digestibility was estimated using the

formula

DMD% = 83.58-(0.824ADF% + 2.626N%) suggested by Oddy *et al.* (1983) and metabolizable energy was also predicted using the equation ME = [0.17DMD% - 2] suggested by SCA (1990)].

Samples were ground to pass through a 2 mm sieve size and 5 grams of each sample were scanned by NIRS. The scanning range from 950-1650 nm (DA 7200 Perten instruments, Sweden) and the spectra were recorded as log (1/R) at 2 nm intervals. Before scanning the samples pre-dried at 60°C overnight in an oven to standardize moisture conditions. Samples were scanned twice in duplicate repacking.

Spectral data was exported into the Unscrambler (CAMO AS, version 9.5, Norway) software for multivariate analysis. Principal Component Analysis (PCA) was performed before Partial Least Squares (PLS) regression models were developed. PCA is a mathematical procedure for resolving sets of data into orthogonal components whose linear combinations approximate the original data to any desired degree of accuracy (Naes et al., 2002). Since the PLS method is a soft-modelling method (Wold et al., 2001; Naes et al., 2002), therefore calibrations models developed using PLS regression. The optimum number of terms in the PLS calibration models were determined by full cross validation and defined by the PRESS (prediction residual error sum of squares) function in order to avoid over fitting of the models (Naes et al., 2002). resulting calibration equations between the chemical reference values

and the NIRS data were evaluated based on the coefficient of determination in calibration (R²_{cal}) and the Standard Error of Cross Validation (SECV). Another measure of the models is the Residual Prediction Deviation (RPD) which is the ratio of Standard Deviation (SD) to the standard error of cross validation (SECV). This is particularly useful in comparing prediction the between alternative models (Lomborg et al., 2009). An RPD value greater than three is considered adequate analytical purposes in most of the NIRS applications for agricultural products (Williams, 2001; Fearn, 2002), whereas a value of 2.5 for the RPD may be regarded as a lower limit for robust NIRS calibrations in quantitative analysis (Williams, 2001). The NIRS calibration models were tested using the validation set. The standard error of prediction (SEP), bias and slope were used to evaluate the calibrations developed.

Results and discussion

The descriptive statistics (mean, range and standard deviation) of the chemical parameters in the calibration validation sets are showed in (Table 1). In both calibrations and validation sets a wide range in variation in chemical composition was observed due to the different stages (phenological or harvest times) of Bromus tomentellus samples (vegetative, flowering and seeding) collected. This variation or range in chemical composition was considered adequate to test the feasibility of developing NIR calibrations for the chemical parameters analysed.

Table 1. Descriptive statistics of <i>Bro</i>	<i>us tomentellus</i> samples	used to develop the NIRS
calibration (% DM basis)		

Variable	N	Mean	SD	Range			
Calibration samples (40)							
N%	40	1.36	0.87	0.36-3.88			
ADF%	40	47.34	7.86	30.4-61.84			
DMD%	40	48.13	8.58	33.49-68.72			
ME%	40	6.19	1.45	3.69-9.68			
Validation samples (2	3)						
N%	23	1.58	1.00	0.37-4.17			
ADF%	23	42.38	6.72	33.07-58.08			
DMD%	23	52.81	7.72	38.29-67.12			
ME%	23	6.98	1.31	4.24-9.41			

N: number of samples; N: Nitrogen; ADF: Acid Detergent Fiber; DMD: Dry Matter Digestibility; ME: Metabolizable Energy; SD: Standard Deviation

Fig. 1 shows the score plot of the first two principal components of the *Bromus tomentellus* samples analysed NIRS. The first PC accounts obtained the maximum possible variance of the data (Osborne *et al.*, 1993). The second PC is orthogonal to the first PC, uncorrelated with it must explained the largest part of the residual

variance unexplained by the first component (Ventura & Papini, 1997). The first two principal components (PCs) explain 91% of the variation in the spectra of the samples analysed (PC1 = 83% and PC2 = 8%). The separation observed between samples is related to the time of sampling harvest.

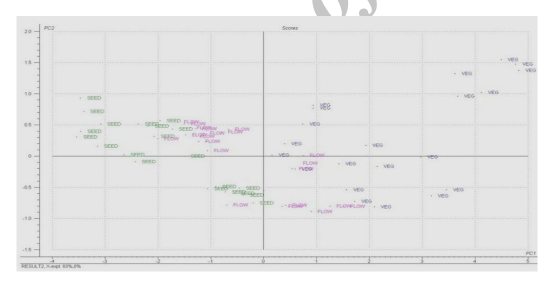


Fig. 1. Score plot of the two principal components in *Bromus tomentellus* marked according to phenological stages (Vegetative=VEG, Flowering=FLOW and Seeding=SEED)

Tables 2 and 3 show the calibration and validation statistics for each of the chemical constituents analysed. The R² and SECV were for DMD 0.98 (SECV: 1.97 %), for N 0.94 (SECV: 0.208 %), for ADF 0.98 (SECV: 1.76 %) and for ME 0.97 (SECV: 0.34 %). According to the report by Stchur *et al.* (2002), if the SEC and SECV values approximately be

similar values, then the calibration model is probably a valid one. In this study according to the Table (2), the SEC and SECV values obtained for the parameters analysed were the same values, approximately. Williams (2003) stated that a value for R² between 0.66 and 0.81 indicates approximate quantitative predictions, whereas, a value for R²

between 0.82 and 0.90 reveals good prediction. Calibration models having a value for R² above 0.91 are considered to be excellent. In our research the values of R² for all parameters analysed were above 0.91. The RPD for the NIRS calibration for the evaluated parameters demonstrated how well the calibration models performed in predicting the reference data. If a product shows a narrow range in composition, or if the error in estimation is large compared with the spread (as SD) in composition, then the regression method finds increasing difficulty in finding stable NIRS calibrations. Sinnaeve et al. (1994) reported that a ratio >3.0 is indicative of a calibration which should perform well for quantitative purposes whilst a ratio >2.5 and <3.0 indicates a calibration that can be used for screening purposes. A value <2.5 indicates that a calibration is of only limited value. The RPD values obtained in calibration for the chemical parameters analysed were 4.35, 4.46, 4.35 and 4.26 for N, ADF, DMD and ME respectively. The RPD values indicated that the PLS calibrations developed can be used on routine analysis. Similar results were reported by other authors when forage samples were analysed using NIRS (Shenk Westerhaus. 1993; Muray, 1993: Garrido, 1997; Alomar et al., 1999; Deaville & Flinn, 2000; Roberts et al., 2004).

Table 2. Near infrared reflectance calibration statistics for whole *Bromus tomentellus* samples variables

Variable	N	Mean	SEC	SECV	R^2	1-VR	RPD
N%	40	1.33	0.19	0.20	0.94	0.75	4.35
ADF%	40	43.75	1.49	1.76	0.98	0.95	4.46
DMD%	40	51.00	1.44	1.97	0.98	0.95	4.35
ME%	40	6.80	0.3	0.34	0.97	0.95	4.26

N: number of samples in calibration; SD: Standard Deviation; SEC: Standard Error of Calibration; SECV: Standard Error of Cross Validation; R²: Coefficient of Determination for Calibration; 1-VR: Coefficient of Determination for Cross Validation; RPD: SD/SECV

Table 3 shows the NIRS validation statistics. The R² and SEP were for DMD 0.96 (SEP: 2.48), for N 0.93 (SEP: 0.36), for ADF 0.97 (SEP: 2.45) and for ME 0.97 (SEP: 0.55). The predictive accuracy for the NIR models was

considered intermediate as judged by the RPD values obtained. The RPD values obtained in validation for the chemical parameters analysed were 2.71, 2.73, 3.45 and 2.38 for N, ADF, DMD and ME respectively.

Table 3. Validation statistics for whole *Bromus tomentellus* samples variables

Variable	N	SEP	Bias	R^2	Slope	Offset	RPD
N%	23	0.36	-0.001	0.93	0.88	0.13	2.71
ADF%	23	2.45	-0.015	0.97	0.95	2.24	2.73
DMD%	23	2.48	-0.017	0.96	0.93	3.23	3.45
ME%	23	0.55	-0.002	0.97	0.94	0.36	2.38

N: number of samples in validation; SEP: Standard Error of Prediction; Bias: average between reference and NIRS values; Slope: Slope of reference vs. NIRS; Offset = the point where a regression line crosses the ordinate (y-axis); RPD: Standard Deviation/SEP

The results from this study suggested that Bromus tomentellus samples might be analysed by NIRS spectroscopy in order to determine N, DMD, ME and ADF. However, the prediction accuracy obtained (RPD values in validation) is desirable for less than analytical purposes. Differences in the calibration statistic were observed when samples were split into calibration and validation sets. Differences in the prediction performance of the NIRS method (see Table 3) developed imply that the calibration models might be sensitive to the range of sample types (harvest or phenological stages) used to develop calibration models. Therefore, samples from more years or harvest need to be included in the calibration data in order to increase the robustness of the NIRS models for routine analysis. Further work will be carried out in order to assess the robustness of the NIRS calibrations models and to incorporate more chemical parameters.

Acknowledgments

We thanks to the Forest, Rangelands and Watershed Management Organization of Iran that supported the research project.

References

- Alomar, d., Fuchslocher, R. and Stockebrands, J., 1999. Effect of ovenor freeze-drying on chemical composition and NIR spectra of pasture silage. *Jour. Animal Feed Science and Technology*, **80:** 309-319.
- Andrés, S., Javier Giráldez, F., López, S., Mantecón, A. R. and Calleja, A., 2005. Nutritive evaluation of herbage from permanent meadows by near-infrared reflectance spectroscopy: 1. Prediction of chemical composition and in vitro digestibility. *Jour. the Science of Food and Agriculture*, **85:** 1564-1571.
- AOAC, 1995. Official methods of analysis (15th ed). Association of

- Official Analytical Chemists. Inc. Arlington, USA Volume 2, 600 p.
- Arzani, H., Motamedi, J. and Zare Chahouki, M. A., 2011. Forage quality of Iranian rangeland species. Forest, Rangeland and watershed management organization and university of Tehran, 234 p. (In Persian).
- Calderon, F. J., Vigil, M. F., Reeves, J. B. and Poss, D. J., 2009. Mid-infrared and near-infrared calibrations for nutritional parameters of triticale (Triticosecale) and pea (*Pisum sativum*). *Jour. Agricultural food chemistry*, **57:** 5136-5142.
- Charehsaz, N., Jafari, A. A., Arzani, H., and Azarnivand. H., 2010. Evaluation of the changes in the water soluble carbohydrate percentage in three species *Bromus tomentellus*, Agropyron intermedium and *Dactylis glomerata* in three phenological stages. *Jour. Rangeland*, **4(1)**: 121-129. (In Persian).
- Cozzolino, D., Fassio, A. and Gimenez, A., 2000. The use of near infrared reflectance spectroscopy (NIRS) to predict the composition of whole maize plants. *Jour. Science Food Agriculture*, **81:** 142-146.
- Cozzolino, D., Smyth, H. E. and Gishen, M., 2003. Feasibility study on the use of visible and near infrared spectroscopy together with chemometrics to discriminate between commercial white wines of different varietal origins. *Jour. Agricultural and Food. Chemistry*, **51**: 7703-7708.
- Deaville, G. D. and Flinn, P. C., 2000. Near infrared spectroscopy: an alternative approach for the estimation of forage quality and voluntary intake. In: Forage Evaluation in Ruminant Nutrition (Eds D. I. Givens, E. Owen, R. F. E. Oxford & H. M. Omed), CABI publishing, Wallingford, UK. 301-320 p.

- Fearn, T., 2002. Assessing calibrations: SEP, RPD, RER and R2. NIR News, **13:** 12-14.
- Ganskopp, D. and Bohnert, D., 2003. Mineral concentration dynamics among seven northern great basin grasses. *Jour. Range Management*, **56:** 174-178.
- Garrido, A., 1997. Current and future applications of NIRS technology. *Jour. Options Mediterranean's, Series Cahiers*, **26:** 87-92.
- Graeff, S., Steffens, D. and Schubert, S., 2001. Use of reflectance measurements for the early detection of N, P, Mg, and Fe deficiencies in corn (*Zea mays L.*). *Jour. Plant Nutrition and Soil Science*, **164:** 445-450.
- Kokaly, R. F. and Clark, R. N., 1999. Spectroscopic determination of leaf biochemistry using band-depth analysis of absorption features and stepwise multiple linear regression. *Jour. Remote sensing environment*, **67:** 267-287.
- Li, B., Liew, O. W. and Asundi, A. K., 2006. Pre-visual detection of iron and phosphorus deficiency by transformed reflectance spectra. J. Photochemi. Photobio. B: *Jour. Biology*, **85:**131-139.
- Lomborg, C. J., Holm-Nielsen, J. B., Oleskowicz-Popiel, P. and Esbensen, K. H., 2009. Near infrared and acoustic chemometrics monitoring of volatile fatty acids and dry matter during codigestion of manure and maize silage. *Jour. Bioresour. Technol*, **100**: 1711–1719.
- Murray, I., 1986. Near infrared reflectance analysis of forages. In Haresign, W., Cole, D. J. A. (Eds) Recent Advances in Animal Nutrition. Studies in the Agricultural and Food Sciences. Butterworths, UK. 141-156 p.
- Murray, I. 1993. Forage analysis by Near Infrared Spectroscopy. In: Davies, A., Baker, R. D., Grant, S. A., Laidlaw, A. S. (Eds), Sward Management

- Handbook. 285-312p. British grassland society, UK.
- Naes, T., Isaksson, T., Fearn, T. and Davies, T., 2002. A user-friendly Guide to Multivariate Calibration and Classification; NIR Publications. Chichester, UK. 420 p.
- Norris, K. H., Barnes, R. F., Moore, J. E. and Shenk, J. S., 1976. Predicting forage quality by Infrared Reflectance Spectroscopy. *Jour. Animal Science*, **43:** 889–897.
- Oddy, V. H., Robards, G. E. and Low, S. G., 1983. Prediction of in vivo dry matter digestibility from the fiber nitrogen content of feed. In: Robards, G. E., Packham, R. G (Eds.), feed information animal production. Common wealth agricultural bureaux. farnahm royal, UK. 395-398 p.
- Osborne, B. G., Merten, B., Thomson, M. and Fearn, T., 1993. The authentication of Basmati rice using near infrared spectroscopy. *Jour. Near Infrared Spectroscopy*, **1:** 77-83.
- Roberts, C. A., Stuth, J. W. and Flinn, P., 2004. Analysis of forages and feedstuffs. In: Roberts, C, A. Workman, and J. B. Reeves (eds.) Journal of Near infra-spectroscopy in agriculture. Agron. Mongor. 321. ASA, CSSA, and SSSA, Madison, Wisconsin, USA. 231-269 P.
- SCA (Standing Committee on Agriculture). 1990. Feeding standards for Australian livestock ruminants, CSIRO, Australia.
- Scholtz, G. D. J., Merwe, H. J. V. and Tylutki. T. P., 2009. Prediction of chemical composition of South African *Medicago sativa* L. hay from a near infrared reflectance spectroscopy spectrally structured sample population. *Jour. South African Animal Science*, **39**: 183-187.
- Shenk, J. S. and Westerhaus, M. O., 1993. Analysis of Agriculture and Food

- Products by Near Infrared Reflectance Spectroscopy, Infrasoft International Software (ISI), silver spring, MD, USA. 26 p.
- Shenk, J. S. and Westerhaus, M. O. 1994. The application of Near Infrared reflectance Spectroscopy (NIRS) to forage analysis. In: Fahey GC Jr (ed) Forage quality evaluation and utilization. American Society of Agronomy, Crop Science Society of America, Soil Science Society of America, Madison, Wisconsin.
- Sinnaeve, G., Dardenne, P., Agneessens, R., and Biston, R., 1994. The use of near infrared spectroscopy for the analysis of fresh grass silage. *Jour. Near Infrared Spectroscopy*, **2:** 79–84.
- Starks, P. J., S. W. Coleman, and W. A. Phillips. 2004. Determination of Forage Chemical Composition Using Remote Sensing. *Jour. Range Management*, **57**: 635-640.
- Stchur, P., Cleveland, D., Zhou, J. and Michel, R. G., 2002. A review of recent applications of near infrared spectroscopy, and of the characteristics of a novel PBS CCD array-based near-infrared spectrometer. *Jour. Applied Spectroscopy Reviews*, **37(4)**: 383–428.
- Stoddart, L. A., Smith, A. D. and Box, T. W., 1975. Range management. Third Ed, Mc Grow hill Book Company. New York. 532 p.
- Valentine, J. F., 1990. Grazing management. Academic press Inc., San Diego. 528 p.
- Van Soest, P. J. 1963. Use of detergents in the analysis of fibrous feeds. II. A rapid method for the determination of fiber and lignin. *Jour. Association Official Agriculture Chemistry*, **46:** 829-835.
- Ventura, C. and Papini, M., 1997. Analysis of the scattering properties of granular materials. *Jour. Near Infrared Spectroscopy*, **5:** 123-133.

- Williams, P. C., 2001. Implementation of near infrared technology. In: Near Infrared Technology in the Agricultural and Food Industries (second edition), Williams, P. C. & Norris, K. (eds). American Association of Cereal Chemist St. Paul, Minnesota, USA. 1455-169 p.
- Williams, P. C., 2003. Near-infrared Technology–Getting the Best out of Light. PDK Grain. Nanaimo, Canada. 132 p.
- Wold, S., Sjostrom, M. and Erikssonn, L., 2001. PLS-regression: a basic tool of chemometrics. *Jour. Chemometrics and Intelligent Laboratory Systems*, 58: 109–130.