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values and yields similar results to those obtained by the DBE procedure, it can be concluded that the DBE procedure is a rapid and reliable method for estimating protein content.

The regression equations obtained by the DBE and TAA methods are presented in Table I. The correlation coefficient was found to be 0.996 and the regression equation obtained by the TAA method had a standard error of estimate of 0.29%.

TABLE I
Correlation Coefficients and Standard Errors of Estimate of Protein Content Determined by DBE and TAA Methods

Method	Correlation Coefficient	Standard Error of Estimate (%)
DBE	0.996 ^a	0.29
TAA	0.986 ^b	0.33
DBE-TAA	0.996 ^c	0.29
DBE-TAA	0.996 ^d	0.29

^aDetermined by linear regression analysis.
^bDetermined by linear regression analysis.

^cDetermined by least squares regression analysis.
^dDetermined by least squares regression analysis.

^eDBE, Cereal Protein Estimator; TAA, total amino acid analysis.

The results obtained by both the procedures were compared.

RESULTS AND DISCUSSION

Protein Estimation

A highly significant correlation coefficient ($r = 0.997^{\circ}$) was obtained between the DBE procedure and total amino-acid analysis. Protein content in cereals ranged from 7.1 to 19.1%. A regression equation derived from 120 samples was calculated as follows:

$$\text{DBE} = 0.958X + 0.159 \quad (1)$$

In this equation, X is the total amino-acid analysis value expressed as mg/g, and DBE is the protein content expressed as mg/g. The regression equation obtained by the DBE procedure had a standard error of estimate of 1.13% and a correlation coefficient of 0.995. The results obtained by the DBE and TAA procedures were compared as follows:

$$\text{DBE} = 0.958X + 0.159 \quad (2)$$

In this equation, X is the total amino-acid analysis value expressed as mg/g, and DBE is the protein content expressed as mg/g. The regression equation obtained by the DBE procedure had a standard error of estimate of 1.13% and a correlation coefficient of 0.995. The results obtained by the DBE and TAA procedures were compared as follows:

$$\text{DBE} = 0.958X + 0.159 \quad (3)$$

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$$\text{DBE} = 0.958X + 0.159 \quad (5)$$

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$$\text{DBE} = 0.958X + 0.159 \quad (6)$$

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$$\text{DBE} = 0.958X + 0.159 \quad (7)$$

In this equation, X is the total amino-acid analysis value expressed as mg/g, and DBE is the protein content expressed as mg/g. The regression equation obtained by the DBE procedure had a standard error of estimate of 1.13% and a correlation coefficient of 0.995. The results obtained by the DBE and TAA procedures were compared as follows:

$$\text{DBE} = 0.958X + 0.159 \quad (8)$$

In this equation, X is the total amino-acid analysis value expressed as mg/g, and DBE is the protein content expressed as mg/g. The regression equation obtained by the DBE procedure had a standard error of estimate of 1.13% and a correlation coefficient of 0.995. The results obtained by the DBE and TAA procedures were compared as follows:

$$\text{DBE} = 0.958X + 0.159 \quad (9)$$

ASTRACT

Sorghum (Sorghum bicolor (L.) Moench) samples were analyzed for protein content by two different methods. The DBE method was determined by a colorimetric method, while the TAA method was determined by a micro-Kjeldahl method. Both methods provided similar results. Correlation coefficients between DBE and TAA were found to be 0.996 for the DBE-TAA procedure, 0.997 for the DBE-TAA procedure, 0.996 for the DBE-TAA procedure, 0.996 for the DBE-TAA procedure, and 0.996 for the DBE-TAA procedure. The results obtained by the DBE method were found to be 9.6% lower than those obtained by the TAA method.

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was obtained that had 0.17 as standard error of estimate. This equation was used to predict the lysine values in the 42 unknown experimental samples. The results of observed and estimated lysine values were compared (Table II). The mean error percentage for the various groups ranged between -2.51 and +4.27% indicating that the DBC method in combination with protein content in samples could give a reliable estimate of lysine content in a sorghum sample. A paired t test did not show any significant difference between the two means. Preliminary results obtained with nine samples that varied in their tannin content (catechin equivalents) from 1 to 2.4% showed that tannin content might not influence the estimated lysine values.

When the results of the 58 samples and the 42 samples were combined, the correlation coefficient between UIR/P and lysine values was observed to be 0.933**. The relationship between these two parameters is given in Fig. 1. A regression equation $Y = 0.644 \times -0.082$ with a standard error of estimate of 0.16 was obtained for the 100 samples. This equation has been used in our laboratory for predicting the lysine values in routine screening of several thousand sorghum samples every year.

The biuret procedure of Johnson and Crane (1971) was checked carefully for the estimation of protein in sorghum in our laboratory. We modified the procedure using 50 mg of sample and reducing the quantities of all other chemicals proportionately so that the reaction could be performed in screw-capped test tubes for the analyses of large numbers of samples. A highly significant correlation was obtained between the biuret procedure and MKJ values. However, the errors associated with the biuret procedure when used for routine screening were large and unpredictable and hence we could not rely on this procedure in our laboratory (Jambunathan 1977). When the UIR values obtained for the 100 sorghum samples used in the present study were compared with the micro Kjeldahl protein values, a positive and significant correlation ($r = 0.782^{**}$) was obtained indicating the possibility of using the DBC procedure for the estimation of protein in sorghum

samples. However DBC values would overestimate the protein content of high-lysine samples. Therefore DBC might be more accurate for the estimation of protein for each class.

Our earlier dye binding analyses were performed on samples containing 80 mg of protein and this procedure was found to be satisfactory in predicting the lysine content (unpublished data). Unfortunately, however DBC analysis could not be performed unless the results of protein analyses were available for these samples as constant protein levels were necessary. As we could analyse about 160 samples per day by the DBC procedure and only about 100–120 samples by the TAA method, we could not proceed with the DBC analysis. Therefore, we decided to use the procedure described in this article. However, our experiments showed that the UIR reading obtained at constant protein could be related to the UIR reading obtained on 1 g of sample if the latter values are calculated and expressed on a constant protein basis (unpublished data). Also a negative and significant correlation ($r = -0.743^{**}$) was obtained between the UIR values obtained at one protein level and the MKJ protein value indicating the possible existence of a negative relationship between protein and lysine content if

TABLE I
Deviation of Technicon Auto Analyzer (TAA) Protein from Micro Kjeldahl (MKJ) Protein Values

Protein Class (MKJ)	No. of Samples	Estimated MKJ Protein (% from TAA Values)		
		Determined MKJ Protein (%) (mean)	TAA Values (mean) ^a	Error (%)
6.0–6.9	1	6.40	6.41	0.5
7.0–7.9	1	7.40	7.10	-1.4
8.0–8.9	7	8.57	8.70	1.5
9.0–9.9	21	9.45	9.53	0.8
10.0–10.9	21	10.42	10.59	1.6
11.0–11.9	14	11.39	11.47	0.7
12.0–12.9	20	12.40	12.16	-0.1
13.0–13.9	15	13.56	13.63	0.5
14.0–14.9	9	14.57	14.53	-0.1
15.0–15.9	5	15.34	15.30	-0.1
16.0–16.9	4	16.11	15.78	-1.4

^aEstimated MKJ protein (%) = 0.985 (TAA) + 0.159

TABLE II
Correlation Between Dye-Binding Capacity (DBC) and Lysine Content

Protein Group	Protein (%)	No. of Samples	UIR ^a (% T)	UIR/P ^b	Lysine (g/100 g P)	UIR/P ^b and Lysine	r Between
Low	7.3–10.7	16	22.36	2.66–1.50	5.9–2.38	0.768 ^c	
Medium	10.8–14.7	17	31.63	2.46–4.36	14.2–2.76	0.919 ^c	
High	15.0–18.2	25	35.72	2.12–4.57	14.4–2.98	0.951 ^c	
All	7.3–18.2	58	22.72	2.32–4.57	13.4–2.98	0.927 ^c	

^aUIR = Udy Instrument reading P = percent of protein

^bSignificant at $P = 0.01$

^c $y = 0.662 \times -0.144$

TABLE III
A Comparison of Determined and Estimated Lysine Values (g/100 g P)

Lysine Level	No. of Samples	Mean Determined Lysine ^a (%)	Mean Estimated Lysine ^b (%)	Error (%) of AAA ^c Values
1.0–1.9	2	1.14	1.15	0.75
1.4–1.79	6	1.64	1.71	4.27
1.8–2.19	15	1.99	1.94	-2.51
2.2–2.59	12	2.39	2.39	
2.6–2.99	7	2.69	2.68	-0.37

^aAmino acid analyzer values

^bEstimated as percent lysine = $0.662 \times (\text{UIR/P}) - 0.144$

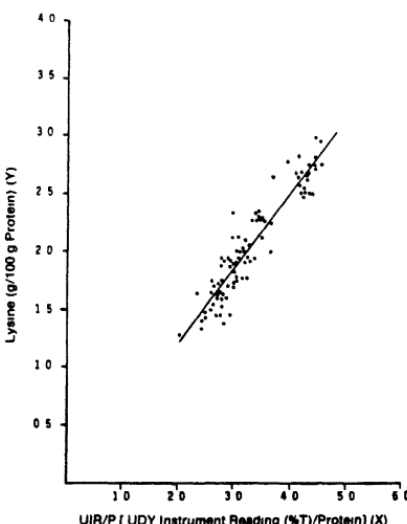


Fig. 1 Relationship between UIR/P and lysine content for sorghum
 $Y = 0.644x - 0.082$ ($n = 100$, $r = 0.933^{**}$)

sorghum. However, conclusions cannot be drawn from this observation as this was not the objective of the study and also because of contrasting reports made about this relationship (Hulse et al 1980).

We observed that the results of protein and lysine analyses based on TAA and DBC methods agree fairly well with the MKJ and amino acid analyzer values, respectively. Considering the time and effort required to determine protein and lysine values, we recommend these two rapid procedures in any large breeding program in which attempts are being made to improve the protein content and protein quality of sorghum. However, it is important to test and standardize these two procedures independently in any laboratory, using sorghums containing a wide range of protein contents, before general use in a breeding program. Also, a bulk check sample should be used to monitor the day-to-day variations, if any, in the analytical results. We have analyzed over 30,000 sorghum samples using these two rapid procedures. This procedure was also successfully used to monitor the progenies arising out of crosses between high-lysine and normal sorghums (Riley 1980).

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