Inheritance of Dhurrin Content in Mature Sorghum Leaves

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Inheritance of Dhurrin Content in Mature Sorghum Leaves

H. J. Gorz, F. A. Haskins, and K. P. Vogel

ABSTRACT

Seedlings of both KS8 and N32 sorghum [Sorghum bicolor (L.) Moench] were high in dhurrin [p-hydroxy-(S)-mandelonitrile-β-D-glucoside] and thus in hydrocyanic acid potential (HCN-p), but the HCN-p of mature leaves from field-grown plants of KS8 was only about one-tenth as high as that of N32. A study of the inheritance of this large difference between KS8 and N32 revealed that a single major gene pair was responsible. There were no obvious maternal effects and F1's were generally intermediate in HCN-p level between the two parents, indicating that neither high nor low HCN-p was completely dominant. HCN-p level was influenced by genetic background since mean values of the low and intermediate HCN-p classes increased slightly as the proportion of genetic background ascribable to N32 was increased. A survey of 102 additional sorghum lines revealed that the gene for low HCN-p carried by KS8 does not appear to occur widely because KS8 had the lowest HCN-p of all entries assayed.

Additional index words: Cyanogenesis, Genetics, Hydrocyanic acid, p-Hydroxybenzaldehyde, Prussic acid, Sorghum bicolor (L.) Moench, Spectrophotometric assay.

The biosynthesis of the cyanogenic glucoside, dhurrin [p-hydroxy-(S)-mandelonitrile-β-D-glucoside], in sorghum [Sorghum bicolor (L.) Moench] has been studied extensively, and much has been learned about the process (1). However, studies of the inheritance of dhurrin content in this species have been much less conclusive. Nass (10) reviewed published reports on the inheritance of cyanogenesis in sorghum, various Lotus species, and Trifolium repens L., and concluded that the situation in sorghum was more complex than that in other species. There was general agreement that the hydrocyanic acid potential (HCN-p) of sorghum leaves was a heritable trait, but the reports failed to agree on such matters as dominance of low or high HCN-p and the number of genes involved. Lack of agreement may have been due in part to differences among the various studies with respect to lines and cultivars used, conditions of growth and sampling, and analytical procedures.

Krauss (7) concluded that HCN-p in sorghum was governed by four gene pairs with additive effects and without dominance. His work was based on crosses among four sorghum cultivars differing in HCN-p. More recent reports have included information about the HCN-p of hybrids and their parents in forage sorghum (13, 15) and also in sudangrass and sudangrass-sorghum combinations (14), but these studies were not designed specifically to investigate the inheritance of HCN-p.

Recently Haskins et al. (4) reported that the two sorghum lines, KS8 and N32, both had high HCN-p levels as seedlings, but when upper leaves from field-grown plants were compared in mid-August, the HCN-p of KS8 was only about one-tenth as high as that of N32. The existence of this large difference suggested that these lines might be useful in a study of the inheritance of HCN-p in mature sorghum leaves.

MATERIALS AND METHODS

Plant Material

The A (cytoplasmic male-sterile) and B (maintainer) lines of KS8 (11) and N32 (12) sorghum were used as parents in the initial crosses for this study. Two crosses, AKS8 X BN32 and AN32 X BK8S, were made during the summer of 1982, and the male-sterile F1's were each backcrossed to both BK8S and BN32 in 1983. Similarly, reciprocal crosses of BK8S and BN32 were made in 1982 following hand-emasculation of florets of the female parents, and the two fertile F1's produced were self-pollinated in 1983 to obtain F2 seed. Additional crosses of AKS8 X BN32 and AN32 X BK8S also were made in 1983 to produce more F2 seed.

Pla...
wiped free of dust, midribs were removed and discarded, and the remaining tissue was cut into pieces 1 to 2 cm² in area. The pieces were thoroughly mixed, and a 2.5-g portion from each leaf was weighed and dried for 2 h at 75°C as previously described (4). The dried samples were weighed, transferred to coin envelopes, and held in a laboratory freezer at −18°C until they were ground. A small Wiley mill fitted with a 1-mm screen was used for grinding the tissue. Ground samples were placed in small plastic vials and stored at −18°C until November, 1984, when they were extracted for HCN-p determination.

**Extraction and Assay of Dried Tissue**

Samples were extracted and assayed in 10 groups, with each of the five replications being divided into two groups; each of which included parental, F₁, backcross, and F₂ entries. A 50-mg portion of each dried and ground leaf sample was weighed into a 20 × 150-mm test tube, 10 mL of distilled water was added, and tubes were shaken briskly on a Gyrotory shaker for 2 h at room temperature. The tissue residue was removed by filtration (Whatman no. 1 filter paper) to provide extracts for assay.

Each extract was assayed as follows: A 1-mL portion was diluted to 10 mL with 0.1 M NaOH. This diluted solution was allowed to stand at room temperature for at least 3 h to hydrolyze any dhurrin present in the extract (9). Cyanide present in the basic solution was measured by the colorimetric procedure of Lambert et al. (8) as modified in the laboratory of Dr. E. E. Conn, Univ. of California, Davis (1982, personal communication). To a 1-mL portion of the hydrolyzed extract in a 20 × 150-mm test tube, 0.5 mL of 1 M acetic acid was added, followed by 5 mL of a solution consisting of 2.5 g of succinimide (Sigma Chemical Company) and 0.25 g of N-chlorosuccinimide (Sigma) in 1 L of water. A 1-mL portion of a solution consisting of 6.0 g of barbituric acid (Sigma), 30 mL of reagent grade pyridine, and 70 mL of water was added. The tube was shaken vigorously and allowed to stand at room temperature for precisely 20 min when absorbance of the solution at 580 nm was read with a Spectronic 20 colorimeter. Extracts were assayed in sets of about 18, with each set including a blank (similar to the F₁) in HCN-p levels; backcrosses to KS8, e) backcrosses of reciprocal F₁'s to BS8, f) backcrosses from reciprocal F₁'s.

A total of 480 seedlings were transplanted from the growth chambers, and 455 survived for sampling as field-grown plants. All data, for seedlings as well as older plants, were based on these 455 plants. All seedling HCN-p values were high (Table 1). Leaves of week-old sorghum shoots typically have a dry matter content of about 11%. On this basis, the HCN-p values of first seedling leaves ranged from about 750 mg kg⁻¹ to 10 000 mg kg⁻¹ dry matter. The HCN-p of K88 was about 1.5 times as high as that of N32 at the seedling stage. At head emergence, however, upper leaves of K88 plants were only about 0.03 as high in HCN-p as comparable N32 leaves (Table 1). The ability to synthesize and/or retain dhurrin decreased for both K88 and N32 between the seedling stage and head emergence, but this decrease was much more pronounced for K88.

The HCN-p values for backcross and F₂ generations were relatively more variable for leaves at head emergence than for seedling leaves, as shown by the magnitudes of means and standard errors in Table 1. Cursory examination of the backcross data suggested that unlike the values for seedling leaves, values for leaves at head emergence had a bimodal distribution. The backcrosses to K88 yielded progeny that were either low (similar to K88) or intermediate (similar to the F₁) in HCN-p; backcrosses to N32 yielded either intermediate or high (similar to N32 progeny. The F₂ distribution had peaks corresponding to low, intermediate, and high HCN-p levels. These observations led to the hypothesis that a single major gene pair accounted for the difference between K88 and N32 with respect to the HCN-p of upper leaves at head emergence. The HCN-p of the F₁ (323 mg kg⁻¹) clearly indicated, however, that neither the high HCN-p of N32 nor the low value of K88 was completely dominant.

To test the single gene hypothesis, it was necessary to classify backcross and F₂ plants as low, intermediate, or high in HCN-p. The F₁ plants would all be heterozygous for the putative gene, and the F₂ mean

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**Table 1. Hydrocyanic acid potential (HCN-p) of first leaves of sorghum seedlings and of mature sorghum leaves at head emergence for the K88 and N32 parents and for the F₁, backcross (BC), and F₂ generations.**

<table>
<thead>
<tr>
<th>Entry</th>
<th>n</th>
<th>First seedling leaves</th>
<th>Mature leaves at head emergence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>HCN-p mg kg⁻¹ fresh wt.</td>
<td>HCN-p mg kg⁻¹ dry wt.</td>
</tr>
<tr>
<td>K88</td>
<td>17</td>
<td>1085 ± 49</td>
<td>24 ± 3</td>
</tr>
<tr>
<td>N32</td>
<td>19</td>
<td>838 ± 21</td>
<td>858 ± 39</td>
</tr>
<tr>
<td>F₁</td>
<td>38</td>
<td>959 ± 28</td>
<td>323 ± 12</td>
</tr>
<tr>
<td>BC to K88</td>
<td>99</td>
<td>996 ± 18</td>
<td>174 ± 17</td>
</tr>
<tr>
<td>BC to N32</td>
<td>94</td>
<td>837 ± 16</td>
<td>713 ± 37</td>
</tr>
<tr>
<td>F₂</td>
<td>188</td>
<td>856 ± 14</td>
<td>387 ± 24</td>
</tr>
<tr>
<td>Total</td>
<td>455</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

† Mean ± SE.
Table 2. Backcross and F1 segregations, levels of hydrocyanic acid potential (HCN-p), and chi-square tests for goodness of fit to indicated genetic ratios. Data are based on mature sorghum leaves sampled at early head emergence.

<table>
<thead>
<tr>
<th>Progeny</th>
<th>Low (mg kg⁻¹)</th>
<th>Intermediate (mg kg⁻¹)</th>
<th>High (mg kg⁻¹)</th>
<th>Ratio</th>
<th>x²</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>BC to KS8</td>
<td>n</td>
<td>46</td>
<td>58</td>
<td>124</td>
<td>x²</td>
<td>P</td>
</tr>
<tr>
<td>Mean ± SE</td>
<td>n</td>
<td>35 ± 3</td>
<td>295 ± 20</td>
<td>1L:1H</td>
<td>0.49</td>
<td>0.48</td>
</tr>
<tr>
<td>BC to N32</td>
<td>n</td>
<td>40</td>
<td>54</td>
<td>94</td>
<td>x²</td>
<td>P</td>
</tr>
<tr>
<td>Mean ± SE</td>
<td>n</td>
<td>387 ± 18</td>
<td>904 ± 37</td>
<td>1L:1H</td>
<td>2.09</td>
<td>0.15</td>
</tr>
<tr>
<td>F1</td>
<td>n</td>
<td>44</td>
<td>94</td>
<td>50</td>
<td>1L:2:1H</td>
<td>0.38</td>
</tr>
<tr>
<td>Mean ± SE</td>
<td>n</td>
<td>46 ± 3</td>
<td>307 ± 13</td>
<td>838 ± 31</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

† HCN-p class limits: Low (L) < 92 mg kg⁻¹; Intermediate (I) 92 to 514 mg kg⁻¹; High (H) > 514 mg kg⁻¹ (dry wt. basis).
‡ mg kg⁻¹ dry wt.

(323 mg kg⁻¹) ± 3 standard deviation units (3 × 77) was used to establish the needed class limits (the probability of an individual F1 (heterozygous) value falling within the F1 mean ± 3 standard deviation units is 0.9973 [6, p. 431]). Thus, plants that assayed from 92 to 554 mg kg⁻¹ were considered to be intermediate (heterozygous), those that assayed higher than 554 were considered to be homozygous high, and those that assayed below 92 were considered to be homozygous low. This classification resulted in the segregations shown in Table 2. Chi-square values were calculated to test the goodness of fit of the various segregations to the ratios shown in the table. Satisfactory fits to the ratios were indicated in each case; therefore, the hypothesis that a single gene pair had a major influence on the difference in HCN-p between KS8 and N32 was supported. The means in Tables 1 and 2 suggest, however, that genetic background also was involved in determining HCN-p levels. Thus, mean HCN-p values for the Low classes were in the order KS8 < backcross to KS8 < F2. Also the Intermediate classes were in the order backcross to KS8 < F1, or F2 < backcross to N32. Not all of the indicated differences were statistically significant, but the trends suggested that in the Low and Intermediate classes, as the proportion of genetic background attributable to N32 increased, HCN-p values increased.

As noted in Materials and Methods, the 46 extracts comprising one of the 10 groups of samples representing a wide range in HCN-p were assayed spectrophotometrically as well as colorimetrically. Mean HCN-p values were 310 mg kg⁻¹ for the spectrophotometric procedure and 371 mg kg⁻¹ for the colorimetric assay. The relationship between the two sets of values was highly consistent, as indicated by the correlation coefficient which exceeded 0.99. Similar results were observed previously (4).

A survey of 104 sorghum lines and cultivars (KS8, N32, and 102 other entries) was conducted. Upper leaves were harvested, dried, ground, extracted, and assayed colorimetrically as in the KS8 × N32 study. The KS8 line was lowest in HCN-p of all the entries, thus, the gene for low HCN-p carried by this line does not appear to occur widely in the entries tested.

The current study and previous work (4) have shown that upper leaves of KS8 were much lower in HCN-p than those of N32. Fall-harvested KS8 tillers also were considerably lower in HCN-p than were comparable tillers of N32 (5). A future study will be designed to determine at what point in plant development these large differences between the lines can be detected, and the extent to which the differences persist throughout the growing season. If both mainstem and tiller leaves of KS8 prove to be consistently low in HCN-p, KS8 would appear to be a promising line for use in breeding low-cyanide sorghums which would provide less dangerous forage for livestock.

ACKNOWLEDGMENTS

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REFERENCES