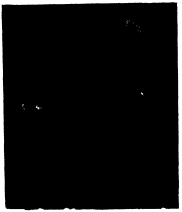
IDENTIFICATION OF A MALE-STERILE GENE IN SORGHUM

The development of random mating populations in sorghums was made possible with the availability of genetic and cytoplasmic-genetic male-sterility systems. So far, ten genetic male-steriles have been reported but only a few of them are free from undesirable character associations. An efficient and easily identifiable male-sterility source is very important for effective random mating and recurrent selection

This note describes a genetic male-sterility system in sorghum and reports its inheritance.

MATERIALS AND METHODS

Three male-sterile plants were observed in the sorghum line IS 104 in 1974 rabs season. The anthers were very small, thu, white and rudimentary (Fig 1) and there were no traces of poller in them. There was no female-sterility. This sorghum line had several desirable characteristics such as short stature, early maturity, bold and white grains (but with persistent sub-coat). No differences were noticed between male-sterile and fertile plants.



Male-sterile (left) and fertile anthers

Die male-sterile plant was crossed with bulk pollen from all the fertile plants in the family Seventyfive plants were grown in F_1 in 1975 summer. In 1975 kharif, an F_3 population of 688 plants was grown and male-sterile plants were identified at bloom +illy male-sterile plants were crossed with pollen from separate fertile plants (plant-to-plant sibbing) Ten F_1 fertile plants were grown as F_3 families along with 50 sibs in 1975 rabi In each segregating family, steries and fertiles were counted at flowering

RESULTS AND DISCUSSION

All F₁ plants were fertile indicating that the malemility is not due to cytoplasmic factors but genetic in nature. The results obtained from $F_{\rm g1},\,F_{\rm g}$ and sibs are presented in Table 1. In the $F_{\rm g}$ generation

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'Inheritance of male-sterility in sorghum							
Segregation in	F,	X	Fa fami	lies	X	Sibs	X1
517 (516)* F	0	006	6 (6 • 67)	Sg 0	• 201	35 (33 0·2	·33) Sg 250
171 (172) S			4 (3·33)	NSg		15 (16-	67)NSg
* Figures	IN	pa	rentheses	are	expe	ected v	alues.

F = Fertile; S = Sterile,

Sg = Showing segregation,

NSg = Not showing segregation.

171 male-steriles appeared out of a total of 688 plants which exactly fit into 3 1 ratio. The ten F_8 families derived from individual F_8 fertile plants had six families (60%) segregating in 3.1 ratio as against expected value of 6 67 (66.67%) on the basis that the ratio of fertile homozygous plants to the heterozygous fertile plants in F_8 is 1 2. These results are confirmed by the segregation pattern observed in families derived from plant-to-plant subbing. Out of 50 families 35 (70%) segregated into 1.1 ratio and 15 (30%) families did not segregate. These results clearly show that the male-sterility reported here is inherited as a single gene recessive

Morphological features of this male-sterility appear to be very distinct and superior to ms_3 and ms_7 geres, and it is much more easily recognised in the field Comparatively the anthers are very small, thin and show complete sterility as against ms_8 gere which is often showing only partial sterility now. Geretic studies are in progress to determine whether it is different from them. A gene symbol will be proposed after confirmation. The stability of this gene in a wide range of genetic and cytoplasmic backgrounds is being tested

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