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RFLP analysis of mitochondrial DNA from cytoplasmic male-sterile lines of pearl millet

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Abstract Mitochondrial DNA (mtDNA) from 13 cytoplasmic male-sterile (cms) lines from diverse sources were characterized by Southern blot hybridization to pearl millet and maize mtDNA probes. Hybridization patterns of mtDNA digested with *Pst*I, *Bam*HI, *Sma*I or *Xho*I and probed with 13.6-, 10.9-, 9.7- or 4.7-kb pearl millet mtDNA clones revealed similarities among the cms lines 5141 A and ICMA 1 (classified as the S-A1 type of cytoplasm based on fertility restoration patterns), PMC 30A and ICMA 2. The remaining cms lines formed a distinct group, within which three subgroups were evident. Among the maize mitochondrial gene clones used, the *cox*I probe revealed two distinct groups of cytoplasmic similar to the pearl millet mtDNA clones. The *atp*9 probe differentiated the cms line 81 A4, derived from *P. glaucum* subsp. *monodii*, while the *cox*II gene probe did not detect any polymorphism among the cms lines studied. MtDNA digested with *Bam*HI, *Pst*I or *Xho*I and hybridized to the *atp*6 probe revealed distinct differences among the cms lines. The maize *atp*6 gene clone identified four distinct cytoplasmic groups and four subgroups within a main group. The mtDNA fragments hybridized to the *atp*6 gene probe with differing intensities, suggesting the presence of more than one copy of the gene in different stoichiometries. Rearrangements involving the *cox*I and/or *rrn*18-*rrn*5 genes (mapped within the pearl millet clones) probably resulted in the S-A1 type of sterility. Rearrangements involving the *atp*6 gene (probably resulting in chimeric form) may be

responsible for male sterility in other cms lines of pearl millet.

Key words RFLP · Mitochondrial DNA · Cytoplasmic male sterility · Pearl millet

Introduction

Cytoplasmic male sterility (cms) is the inability of plants to produce viable pollen while being female fertile. The trait is maternally inherited, indicating that the factors responsible are present on the organellar genome. Cumulative evidence indicates that variations in the mitochondrial genome are associated with male sterility (Hanson and Conde 1985).

Pearl millet is an important cereal and forage crop of the semi-arid tropics and is widely grown in India. The development of cytoplasmic male-sterile lines by pearl millet breeders has facilitated the production of commercial hybrids, thereby leading to increased yields. The introduction of the 'HBI' hybrid, obtained by using cms line Tift 23 A, led to the doubling of pearl millet production between 1965 and 1970 (Burton 1983). However, the extensive use of a single cytoplasm results in genetic vulnerability to prevailing pathogens, as became evident in maize where the use of cms-T cytoplasm led to a Southern leaf blight epidemic caused by *Bipolaris maydis* in the early 1970s, thereby necessitating the search for new sources of cytoplasmic. The conventional method of classifying cytoplasmic on the basis of fertility restoration patterns is cumbersome and time consuming. Restriction fragment length polymorphism (RFLP) of mitochondrial DNA (mtDNA) provides a suitable tool to assess the heterogeneity in the male-sterile cytoplasmic. This technique has been successfully used for the classification of male-sterile cytoplasmic in maize (Borck and Walbot 1982), sorghum (Bailey-Serres et al. 1986) and sugar beet (Weihe et al. 1991).

Fertility restoration patterns distinguished the S-A1, S-A2 and S-A3 cytoplasmic in pearl millet (Burton and

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Athwal 1967). A stable male-sterile cytoplasm was subsequently isolated from *P. glaucum* subsp. *monodii* (Maire) by Hanna (1987). The S-A1 to S-A4 cytoplasmic lines were differentiated following hybridizations of *Bam*HI-digested mtDNA to the maize *atp6* gene probe (Smith and Chowdhury 1989). Several stable cms lines from diverse source have been identified in pearl millet and are currently being used in a breeding program at the International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), India. The study presented here was undertaken to define RFLPs in the mtDNA using 13 cms lines of pearl millet following Southern blot hybridizations to pearl millet and maize mtDNA clones.

Materials and methods

Plant material

The pearl millet accessions used in the study were 5141 A, developed by repeated backcrossing of a downy mildew-resistant donor 1587 to Tift 23 A (Pokhriyal et al. 1976); ICMA 1(81 A), derived by inducing mutations in Tift 23 DB (Andrews and Anand Kumar 1982); ICMA 2 (843 A); L67 A, a cms line described by Burton and Athwal (1967); DSA 59-1, DSA 105, DSA 118, DSA 134 and DSA 144-1, germ plasm accessions from Ghana; and PMC 23 and PMC 30, from Botswana. The male-sterile (A) and their inbred maintainer (B) lines from Ghana and Botswana were identified by Appa Rao et al. (1989) 81 A4 and ICMA 88001 were derived from *Pennisetum glaucum* subspecies *monodii* (Maire) Brunken (Hanna 1987) and *P. violaceum* (Lam.) L. Rich. (Marchais and Pernes 1985), respectively. All of the male-sterile lines were from the International Crop Research Institute for the Semi-Arid Tropics (ICRISAT) Hyderabad, India.

Clones used

The pearl millet mtDNA clones consisted of four *Pst*I fragments, cloned and characterized from the S-A1 system of sterility (Smith and Chowdhury 1991). The maize clones were F₁-F₀ ATPase subunit 6 (*atp6*; Dewey et al. 1985a) and subunit 9 (*atp9*; Dewey et al. 1985b), kindly supplied by CS Leving's III, Genetics Department, North Carolina State University, Raleigh, N. C., USA and cytochrome c oxidase subunit I (*coxI*; Isaac et al. 1985) and subunit II (*coxII*; Fox and Leaver 1981) a gift of CJ Leaver, Department of Plant Sciences, University of Oxford, U.K.

MtDNA extraction, digestion and Southern transfer

Mitochondrial DNA from pearl millet was purified according to the procedure described by Smith et al. (1987). It was then treated with restriction enzymes *Bam*HI, *Pst*I, *Sma*I and *Xho*I according to the supplier's instructions in the presence of RNaseI (15 µg/ml). The reaction was terminated by the addition of BPB solution (25% sucrose, 0.1% Bromophenol blue and 20 mM EDTA). MtDNA fragments were separated by electrophoresis in a 0.8% agarose horizontal slab gel (5 mm thick) at 2 V/cm for 16 h in TBE buffer (89 mM Tris, 89 mM boric acid and 2 mM EDTA, pH 8.3). *Hind*III-generated Lambda DNA fragments were used as standard molecular weight markers. The gels were stained in 0.5 µg/ml ethidium bromide for 40 min and destained for 5 min in deionized water. The gels were viewed on a UV transilluminator and photographed using Kodak Wratten Nos. 2B and 23A filters with type 667 Polaroid film. The Southern blot transfers of DNA fragments onto nylon membrane (Genescreen, Dupont) were performed using the VacuGene blotting apparatus (Pharmacia) according to the manufacturer's protocol.

The blots were washed in 3 × SSC (1 × SSC = 0.15 M NaCl, 0.015 M sodium citrate, pH 7.0) air-dried and baked at 80 °C under vacuum for 1 h.

Probe labelling and Southern hybridization

Purified inserts from various mtDNA clones were labelled with alpha [³²P]-deoxyctidine 5'-triphosphate ([³²P]-dCTP) using the random-primed DNA labelling kit from Boehringer Mannheim and applying the method of Feinberg and Vogelstein (1983). Filter hybridizations were conducted according to Smith et al. (1989). Hybridized nylon filters were washed twice in 3 × SSC containing 0.1% SDS at 65 °C for 15 min each, and autoradiography was conducted at -70 °C for various exposure times using X-ray films and intensifying screens.

Cluster analysis

The fragment sizes were determined using standard molecular weight markers. Similarity index matrices were generated based on the proportion of common restriction fragments between two lanes (Nei 1987) using the formula

$$F = \frac{2M_{xy}}{M_x + M_y}$$

where *F* is the similarity index; *M_x*, the number of fragments in one accession; *M_y*, the number of fragments in a second accession; and *M_{xy}*, the number of fragments common to both accessions.

The value of 1.0 would mean that the patterns in the two accessions are identical. The data are represented in the form of matrices of order *n* × *n* where '*n*' is the number of accessions. The genetic analysis was based on the expression of similarity of objects and respective groups by the agglomerative method of hierarchical clustering technique, which proceeds by a series of successive fusions of the '*n*' objects into clusters. Initially, each object is considered to be a separate member of a cluster. The two objects having the maximum similarity index are then grouped together, and proximities between each of the remaining objects and the two membered cluster is calculated according to the single linkage method. A computer program written in high-level language 'C' was used to construct the dendrograms.

Results

MtDNA restriction patterns

Restriction patterns of mtDNA purified from soft stem internodes of field-grown plants of the cms line PMC30A and its maintainer PMC 30B restricted with *Pst*I, *Bam*HI or *Hind*III showed differences between A and B lines (Fig. 1) The *Pst*I restriction pattern revealed differences in two fragments in each of the A and B lines. The *Bam*HI digestion yielded only a single fragment difference, while *Hind*III showed a difference of two fragments in A and one fragment in B line.

The restriction patterns of etiolated seedlings of some of the A and B lines were complex, and the low-molecular-weight fragments were not clearly resolved. The quality of mtDNA varied among different preparations depending on the genotype and source of tissue used for extraction. In general, mtDNA obtained from soft stem tissues showed better restriction patterns than that obtained from etiolated seedlings.

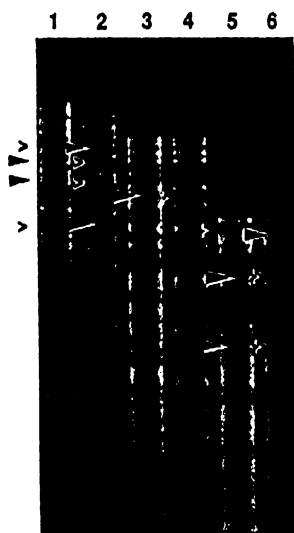
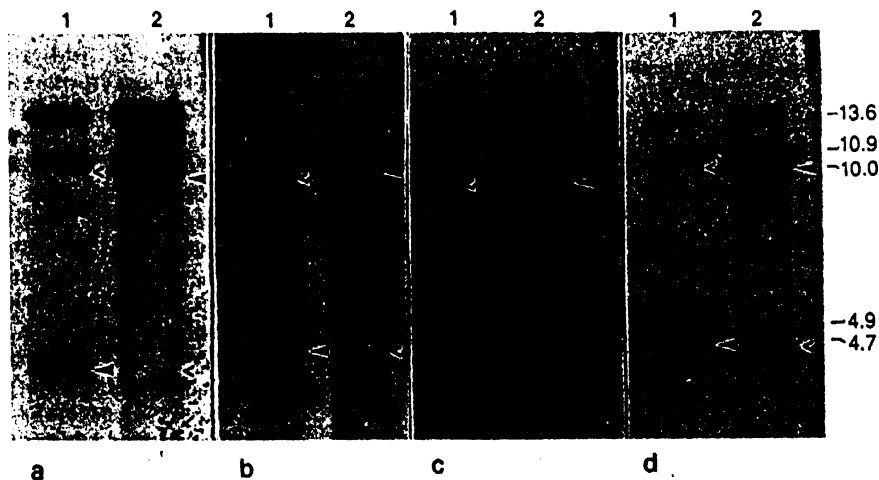


Fig. 1 Restriction fragment patterns of *Pst*I-(lanes 1,2), *Bam*HI-(lanes 3,4) or *Hind*III-(lanes 5,6) digested mtDNA from PMC 30A (lanes 1, 3, 5) and PMC 30B (lanes 2, 4, 6). Note the presence (▶)/absence (>) of fragments

Hybridizations of mtDNA with homologous clones

Hybridization patterns of mtDNA digested with *Pst*I, *Bam*HI, *Sma*I or *Xho*I and hybridized to any one of the four pearl millet clones distinguished one group of cytoplasm consisting of cms lines 5141, PMC 30, ICMA 1 and ICMA 2 (Group I) from the remaining cytoplasm consisting of cms lines L 67, PMC 23, DSA 59-1, DSA 105, DSA 118, DSA 134 and DSA 144-1 (Group II). The cms lines of Group I were characterized by the presence of a 4.7-kb *Pst*I fragment instead of the 10-kb fragment present in the Group II when hybridized to 13.6-, 10.9- and 4.7-kb clones (Fig. 2a, b, d). Additional fragments hybridizing to each of these clones were common to all cytoplasm. The 9.7-kb clone, however,

Fig. 2 Southern hybridization of *Pst*I-digested mtDNA from ICMA 1 (lane 1, representative of group I cytoplasm) and ICMB 1 (lane 2, representative of group II cytoplasm) hybridized to the pearl millet clones. a 13.6 kb, b 10.9 kb, c 9.7 kb, d 4.7 kb. Arrows indicate presence (▶)/absence (>) of fragments



hybridized to the 10-kb *Pst*I fragment in Group II cms lines, but the 4.7-kb fragment hybridizing to the other three pearl millet clones was absent in the Group I cytoplasm (Fig. 2c).

MtDNA digested with *Bam*HI and probed with any one of the four clones revealed a 6-kb fragment in the Group II cytoplasm (Fig. 3), whereas the Group I cytoplasm (represented by 5141 A in Fig. 3, lane 3) was characterized by a relatively higher intensity of a 4.9-kb fragment, suggesting additional fragments of this size. The maintainer lines of Group I and II showed the 6-kb fragment.

A distinct 7.5-kb *Sma*I fragment hybridizing to all four clones was observed in Group I cms lines (represented by 5141 A and PMC 30A), whereas the remaining cms lines showed a 6-kb fragment instead (Fig. 4). Various other fragments hybridizing to the four clones were common to all the cytoplasm.

Southern blot hybridization of the four clones to the *Xho*I digests of mtDNA revealed a 20-kb fragment in Group I cms lines (represented by 5141 A and PMC 30A) instead of the 7-kb fragment present in the remaining cytoplasm (Fig. 5). All of the cytoplasm had several common fragments hybridizing to each of these clones.

Hybridizations of mtDNA with heterologous clones

The maize *atp9* gene probe distinguished cms line 81 A4 from the remaining cytoplasm by hybridizing to a 7.6-kb *Xho*I fragment (Fig. 6 lane 9) not present in the other cms lines as well as to the 4.6- and 5.4-kb fragments present in all of the cytoplasm. The maize *cox*I probe distinguished Group I cytoplasm from the remaining cms lines (similar to the four pearl millet clones). This probe produced a distinct pattern with mtDNA from ICMA 88001 (Fig. 7, lane 6). The maize *cox*II gene probe did not show polymorphism among the various cms lines (data not shown).

Fig. 3 Autoradiogram of 13.6-kb pearl millet clone hybridized to *Bam*HI-digested mtDNA from cms lines. Lanes 1 and 2 PMC 23 A and B, 3 and 4 5141 A and B, 5 and 6 DSA 105 A and B, 7 and 8 DSA 134 A and B, 9 and 10 L 67 A and B, 11 and 12 DSA 59-1 A and B. Note the absence (>) of a 6-kb fragment and the increased intensity of the 4.9-kb fragment in 5141 A (lane 3)

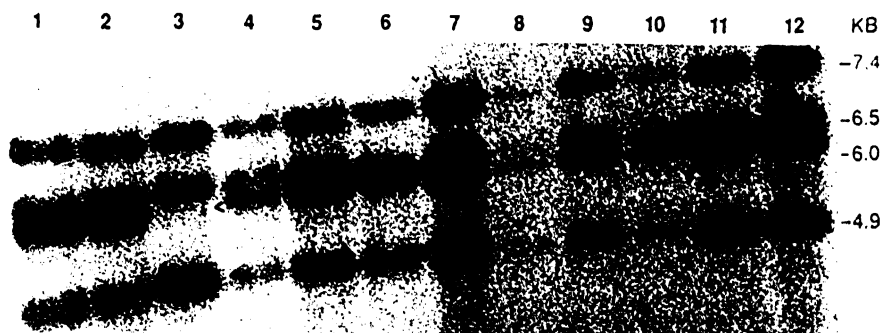
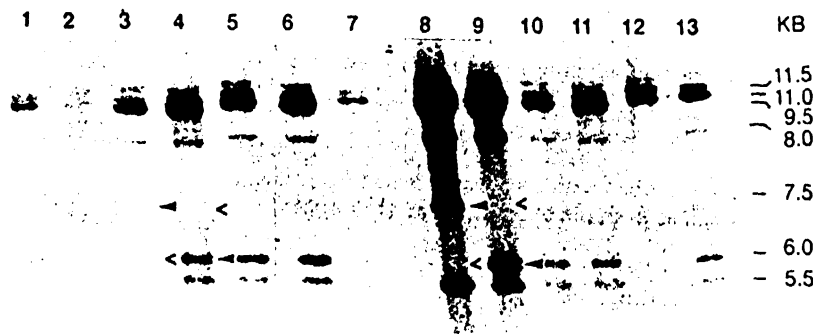


Fig. 4 Hybridization patterns of the 13.6-kb pearl millet clone with *Sma*I digests of mtDNA from cms and their maintainer lines. Lanes 1 and 2 PMC 23 A and B, 3 and 4 5141 A and B, 5 DSA 134 A, 6 and 7 DSA 105 A and B, 8 and 9 PMC 30 A and B, 10 and 11 DSA 118 A and B, 12 and 13 DSA 144-1 A and B. Note the presence (▶) of a 7.5-kb fragment and the absence (>) of a 6-kb fragment in 5141 A and PMC 30A (lanes 3 and 8, respectively)



Further polymorphisms were detected among the cms lines following the hybridization of *Pst*I-, *Bam*HI- or *Xho*I-digested mtDNA with the maize *atp6* gene probe. The probe revealed similar hybridization patterns for mtDNA from Group I cms lines when digested with any one of the three restriction enzymes, as represented in Fig. 8, which shows *Bam*HI-restricted mtDNA from various cms lines hybridized to the *atp6* probe. The hybridization patterns of mtDNA from cms lines L 67, PMC 23, DSA 118 and DSA 134 digested with either

Fig. 5 Southern hybridization of the 4.7-kb pearl millet clone to *Xho*I-digested mtDNA from cms and their maintainer lines. Lanes 1 and 2 PMC 23 A and B, and 3 and 4 5141 A and B, 5 DSA 134 A 6 and 7 DSA 105 A and B, 8 and 9 PMC 30 A and B, 10 and 11 DSA 118 A and B, 12 and 13 DSA 144-1 A and B. Note the presence (▶) of a 20-kb fragment and absence (>) of a 7-kb fragment in 5141 A and PMC 30A (lanes 3 and 9, respectively)

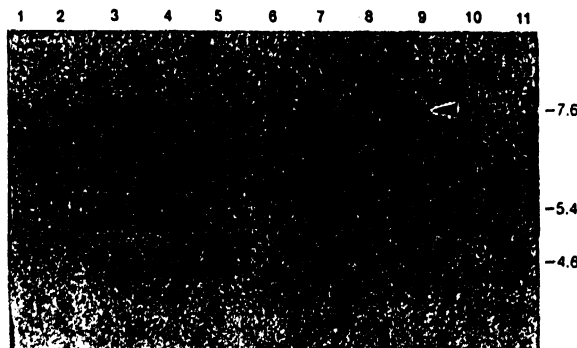
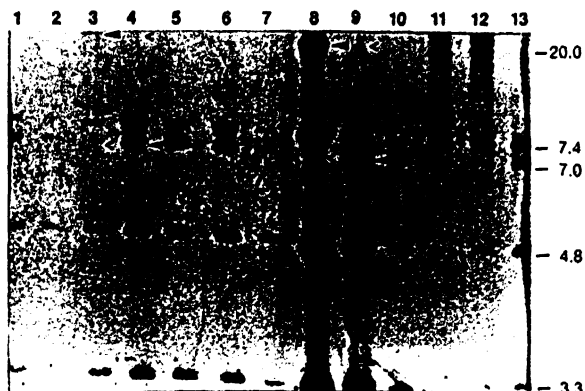


Fig. 6 Southern hybridization of the maize *atp9* gene clone to *Xho*I-digested mtDNA from cms lines. Lane 1 PMC 30A, 2 L 67 A, 3 PMC 23A, 4 DSA 59-1A, 5 DSA 105A, 6 DSA 118A, 7 DSA 134A, 8 DSA 144-1A, 9 81 A4, 10 ICMA 88001, 11 ICMA2. Arrow (▶) indicates the presence of a fragment

*Pst*I or *Bam*HI and probed with the *atp6* clone were similar, whereas cms line DSA 144-1A showed some variation, as represented in Southern blots of *Pst*I-digested mtDNA (Fig. 10a, lane 5). The digests of mtDNA with *Xho*I showed identical hybridization patterns between cms lines L 67 and PMC 23, cms lines DSA 105 and DSA 118, and cms lines DSA 59-1 and DSA 134, whereas DSA 144-1 showed a pattern distinct from all of the above cms lines (Fig. 9, lane 8). A prominent 3.6-kb *Xho*I fragment present in most cytoplasms was absent in DSA 144-1A and 81 A4 (Fig. 9, lanes 8 and 9, respectively). The hybridization patterns of mtDNA from cms lines 88001 and 81 A4 digested with either *Pst*I, *Bam*HI

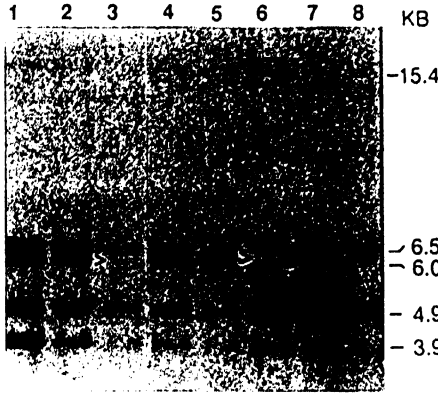


Fig. 7 Autoradiogram of maize *cox1* gene clone hybridized to *Bam*HI-digested mtDNA from cms lines. Lane 1 ICMA 1, 2 ICMB 1, 3 PMC 30A, 4 PMC 23A, 5 L 67A, 6 ICMA 88001, 7 81 A4, 8 ICMB 2. Arrows (>) indicate the absence of fragments

or *Xho*I and probed with *atp6* were distinct from each other as well as from the other cms lines (Fig. 10b).

Analysis of dendrograms

The dendrogram constructed on the basis of similarity indices among various cms pairs following hybridizations of *Pst*I-, *Bam*HI-, *Sma*I- or *Xho*I-digested mtDNA with four pearl millet clones revealed two distinct

Fig. 8 Autoradiogram of the maize *atp6* gene clone hybridized to *Bam*HI-digested mtDNA from cms lines. Lane 1 ICMA 1, 2 ICMA 2, 3 5141A, 4 PMC 30A, 5 L 67A, 6 PMC 23A, 7 DSA 59-1A, 8 DSA 105A, 9 DSA 118A, 10 DSA 134A. M Molecular markers (λ DNA digested with *Hind*III)

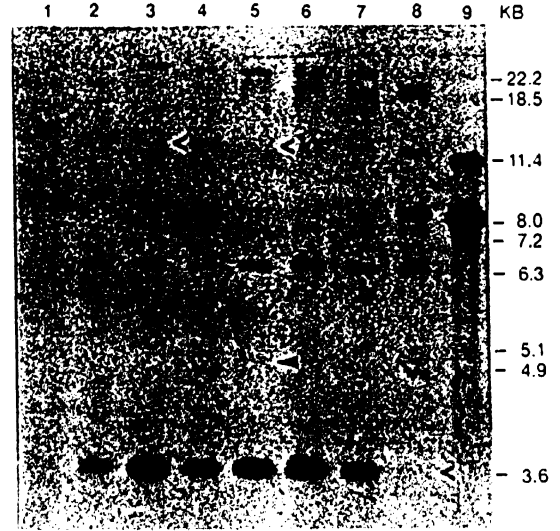
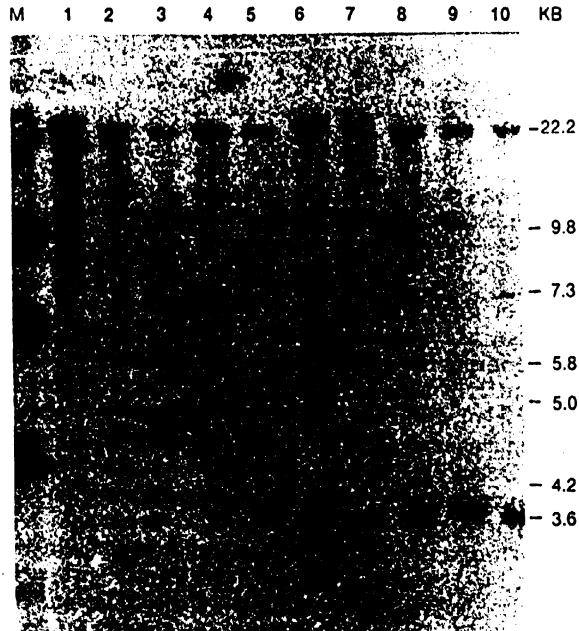
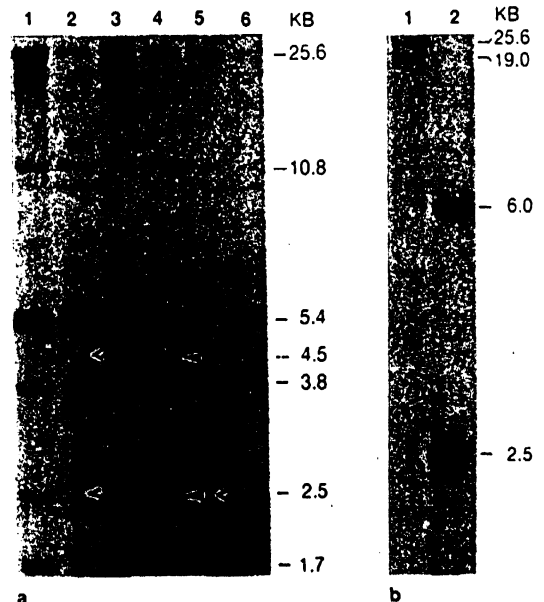


Fig. 9 Southern hybridization of the maize *atp6* gene clone to *Xho*I-digested mtDNA from cms lines. Lane 1 PMC 30A, 2 L 67A, 3 PMC 23A, 4 DSA 59-1A, 5 DSA 105A, 6 DSA 118A, 7 DSA 134A, 8 DSA 144-1A, 9 81 A4. Arrows indicate the presence (\blacktriangleright)/absence (\triangleright) of fragments

groups with $F = 0.8$ as the significance limit (Fig. 11). The male-sterile line 5141 A, ICMA 1, ICMA 2 and PMC 30A formed a distinct cluster, while the remaining 9 cms lines formed a second cluster widely separated from the first cluster. Three subgroups were evident

Fig. 10 a, b Southern hybridization of the maize *atp6* gene clone to *Pst*I-digested mtDNA from cms lines. a Lane 1 ICMA 1, 2 PMC 30 A, 3 L 67A, 4 PMC 23A, 5 DSA 144-1A, 6 ICMA-2. b Lane 1 ICMA 88001, 2 81 A4



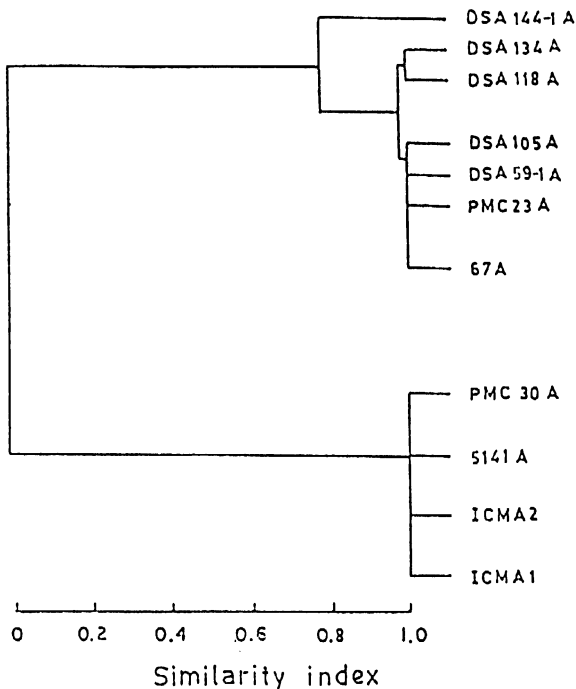
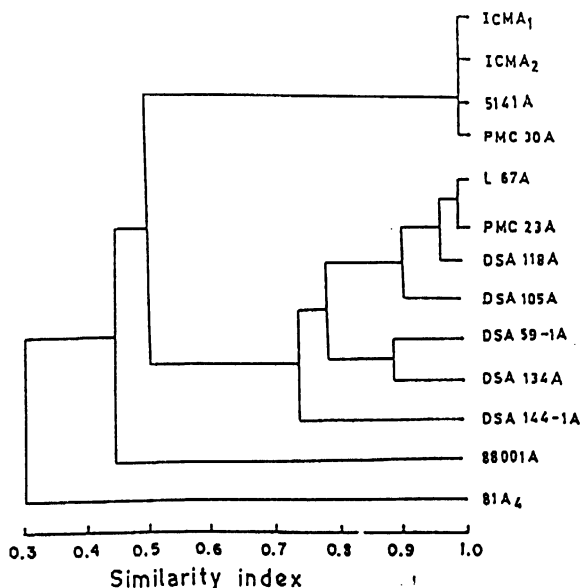


Fig. 11 Dendrogram of cms lines of pearl millet based on homologous mtDNA clones and four restriction enzymes. The dendrogram was constructed on the basis of similarity indices among the various cms lines following hybridization of *Pst*I-, *Bam*HI-, *Sma*I- or *Xho*I-digested mtDNA with the four pearl millet clones as described in the Materials and methods

Fig. 12 Dendrogram of cms lines of pearl millet based on maize *atp6* gene clone and three restriction enzymes. The dendrogram was constructed on the basis of similarity indices among the various cms lines following hybridization of *Pst*I-, *Bam*HI- or *Xho*I-digested mtDNA with the maize *atp6* clone as described in the Materials and methods



within the second cluster: male-sterile lines L 67A, PMC 23A, DSA 59-1A and DSA 105A form a close cluster, differing slightly from DSA 118A and DSA 134A, which were clustered together; DSA 144-1A showed a 20% variation from the others.

The dendrogram based on the similarity indices generated from hybridization patterns of mtDNA digested with *Pst*I, *Bam*HI or *Xho*I and probed with the maize *atp6* clone revealed four groups (Fig. 12) at a level of significance equivalent to the first dendrogram ($F = 0.75$). PMC 30A, 5141A, ICMA 1 and ICMA 2 formed a distinct cluster similar to that seen with the homologous clones. Four subgroups were evident within the second group of cytoplasm. Cytoplasmic male-sterile line 81 A4 and ICMA 88001 were distinct from each other as well as from all the other cms lines.

Discussion

Detection of polymorphism based on restriction patterns

The method of Smith et al. (1987), involving the use of low viscosity, high ionic strength, saline buffers (Bookjans et al. 1984), combined with a stringent washing of the mitochondria with a high salt buffer, potassium acetate precipitation to remove polysaccharides and lipids (Dellaporta et al. 1983) and the precipitation of the nucleic acids with isopropanol-ammonium acetate consistently yielded mtDNA that could be digested with restriction enzymes. However, the yield and purity of the mtDNA varied among the different preparations depending on the genotype and source of tissue used for extraction. In general, mtDNAs obtained from soft stem tissues showed better restriction patterns than those from etiolated seedlings. Restriction patterns have been reported to provide a reliable method to characterize male-sterile cytoplasm in pearl millet (Smith and Chowdhury 1989), whereas under our conditions, the patterns were not consistently clear across the accessions. Therefore, Southern hybridizations using mtDNA specific clones were used to study polymorphisms in various cms lines.

Detection of polymorphism using homologous clones

Hybridization patterns of *Pst*I-, *Bam*HI-, *Sma*I- or *Xho*I-digested mtDNA when probed with any one of the four pearl millet mtDNA clones distinguished Group I cytoplasm consisting of 5141 A, ICMA 1, ICMA 2 and PMC 30A from the Group II male-sterile lines, L 67A, PMC 23A, DSA 59-1, DSA 105A, DSA 118A, DSA 134A and DSA 144-1A as well as their maintainer (fertile) B-lines. On the basis of fertility restoration studies, cms lines 5141 A (Pokhriyal et al. 1976) derived from Tift 23A cytoplasm and ICMA 1 (Andrews and

Anand Kumar 1982) have been earlier classified as one group, designated S-A1 (Smith et al. 1987). According to the present data, cms lines PMC 30A (derived from Botswana accessions) and ICMA 2 could also be grouped as S-A1 cytoplasm. The four pearl millet mtDNA clones used as probes were *Pst*I fragments associated with reversion to fertility in the S-A1 type of cytoplasm (Smith et al. 1987). These clones revealed two sets of homologies following double digestions with various restriction enzymes (Smith and Chowdhury 1991): the 9.7- and 13.6-kb clones shared one set of homology, while the 13.6-, 10.9- and 4.7-kb clones with the *rrn18-rrn5* and *coxI* genes mapped on them showed a second set of homology. The S-A1 cytoplasm was characterized by the presence of a 4.7-kb *Pst*I fragment and an additional 4.9-kb *Bam*HI, 7.5-kb *Sma*I and 20-kb *Xho*I fragment instead of the 10-kb *Pst*I, 6-kb *Bam*HI, 6-kb *Sma*I and 7-kb *Xho*I fragments present in the remaining cytoplasm; the other fragments hybridizing to the four clones were common to all cytoplasm. The 4.7-kb *Pst*I fragment to which the *rrn18-rrn5* and *coxI* genes are mapped is present only in the S-A1 cytoplasm; this corresponds to the 10-kb *Pst*I fragment in the remaining cms lines and also in their maintainer (fertile) cytoplasm. This suggests that rearrangements in the *rrn18-rrn5* and/or *coxI* genes are probably responsible for S-A1 type of sterility, as has been suggested by Smith et al. (1991).

Detection of polymorphism using heterologous clones

Hybridization patterns using maize mtDNA gene probes revealed further differences among the cms lines of pearl millet. The *atp9* probe distinguished cms line 81 A4, derived from *P. glaucum* subsp. *monodii* and previously classified as the S-Am cytoplasm (Smith and Chowdhury 1989). The maize *coxI* probe distinguished the S-A1 type of cytoplasm similar to the pearl millet clones.

The maize *atp6* clone distinguished a wide range of cytoplasm. The four main groups of cytoplasm revealed by the *atp6* probe agreed in general with the classification arrived at by breeders using fertility restoration studies. PMC 30A, 5141 A, ICMA 1 and ICMA 2 formed a distinct cluster, whereas L 67A, previously classified as S-A3 cytoplasm (Burton and Athwal 1967) along with PMC 23A, DSA 118A, DSA 105A, DSA 59-1A, DSA 134A and DSA 144-1A (identified from Botswana and Ghana accessions; Appa Rao et al. 1989) formed another cluster within which further variations were evident. Most of these accessions are stable cms lines possessing several desirable agronomic traits. This variation among the cytoplasm could be of great value in the production of hybrids in different cytoplasmic backgrounds, thus providing greater variability. A similar variation was reported by Weihe et al. (1991) among eight cms lines of *Beta vulgaris* derived from the same source of cytoplasm (Owen's cytoplasm) collected from

various research stations, following hybridization of mtDNA with the maize *coxII* gene probe.

In pearl millet, 81 A4 and ICMA 88001 were distinct from each other as well as from all of the other cms lines. 81 A4 showed patterns similar to the S-Am group of cytoplasm reported by Smith and Chowdhury (1989). ICMA 88001, however, was distinct from all of the cms lines being derived from *P. violaceum* (Lam.) L. Rich (Marchais and Pernes 1985).

Additional fragments in mtDNA of cms lines other than the S-A1 type (Fig. 8) hybridized to the maize *atp6* gene clone with varying intensities, indicating the presence of more copies of this gene with different stoichiometries. This suggests rearrangements in the mitochondrial genome at the *atp6* region that probably lead to the formation of chimeric genes.

The present study revealed polymorphisms in the mtDNA from cms lines and their maintainer lines of pearl millet. Homologous clones could identify one group of cytoplasm, whereas differences among the remaining cytoplasm could only be detected using heterologous probes. The enzyme-probe combinations which differentiated various cytoplasm in the present study would aid in the identification of specific clones. Selection of clones based on the enzyme-probe combinations specific for a particular type of cytoplasm or unique transcripts or translational products would enable us to identify genes responsible for sterility.

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