Original Research Article

CMS Line in Carrot

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A B S T R A C T

Male sterility in vegetables is commercially utilized to develop their hybrids due to cost effective and easy mechanism to produce large-scale hybrid seeds commercially. Different types of male sterility, and three-line and two-line hybrid breeding approaches involving Cytoplasmic male sterility (CMS) and environmental sensitive genic male sterile (GMS) lines, are used for hybrid seed production. In carrot which produces many but small sized flowers, making hand-emasculuation tedious and time consuming. Identification of male sterile and fertile counterpart of CMS lines is also not possible at early seedling stage by using test cross methods. Marker assisted selection for male sterile and fertile (maintainer) genes greatly enhance selection of A and B plants in breeding program. So, molecular markers that can distinguish CMS status and fertile plants at an early seedling stage will be valuable tool in carrot breeding including CMS based hybrid seed programs in hybrid seed purity evaluation, selection and management. Keeping these in views, the present study was undertaken to characterize different sources of cytoplasmic male sterile lines and their maintainers. By using available molecular markers closely linked to sterility gene to distinguish our male sterile and fertile lines.

Keywords
Hybrid carrot seed, Cytoplasmic male sterility, Maintainer, Molecular Markers, PCR.

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Introduction

Cytoplasmic male sterility (CMS) is a maternally inherited trait that has been correlated with changes of the mitochondrial genome or with incompatibility between nuclear and mitochondrial genomes. Evaluation of cytoplasm that induces CMS in carrot for use in breeding programs. Several induce ‘brown anther’ (Sa) male sterility in which the anthers are formed. Sa CMS cytoplasms have been isolated from several open pollinated varieties as well as from wild populations (Welch & Grimball, 1947; Banga et al., 1964; Ronfort et al., 1995).
Other CMS type in carrot is ‘petaloid’ (Sp), in which the stamens are replaced by petals or bract-like structures. This type of cytoplasm found in feral populations of (Daucus carota) ssp. carota from North America and Sweden. Petaloid male-sterile plants were found among wild carrots from USA, Canada, Sweden and Germany. The American sources of sterility were subsequently introduced into the cultivated germplasm allowing production of the modern hybrid varieties (reviewed in Bach 2000). Stamen petaloidy in cultivated carrots was also induced upon cytoplasm transfer from D. c. maritimus (Nothnagel et al., 1997).

CMS, caused by mutations in the mitochondrial genome (Jing et al., 2015), is found in higher plants and increases heterosis and improve genetic resources. Mitochondrial markers can be used to differentiate between the various types of CMS sources (Zhao et al., 2010; Wang et al., 2012; Zhang et al., 2013). In protoplast fusion (Yarrow et al., 1990) and distant hybridization (Jing et al., 2015) phenomenon of cytoplasmic male sterility (CMS; Kaul, 1988) a thesaurus contribution of extra-nuclear genes in flower formation. CMS plants are unable to produce fertile pollen and hence are male sterile. The maternal inheritance of the CMS indicates that the genetic information of either the mitochondria or the plastids participates in certain steps of flower formation. Mitochondrial genes have been recognized as being implicated in all well-investigated CMS systems.

In many cases of CMS, rearrangements in the mitochondrial genome were found to produce unique chimeric genes, and often function of genes encoding subunits of the mitochondrial ATP synthase is impaired, suggesting that reduced mitochondrial respiration and/or ATP production lead to male sterility (reviewed by Budar and Pelletier, 2001; Mackenzie and McIntosh, 1999; Schnable and Wise, 1998). The mitochondrial dysfunction of CMS plants can be either maintained or concealed by specific nuclear gene functions leading, in the latter case, to restored male fertility. Genetic and molecular mechanisms of restoration vary among the different CMS systems (reviewed by Schnable and Wise, 1998). Elimination of CMS-related mitochondrial DNA segments (Janska et al., 1998), restoration of effective editing of specific mitochondrial mRNAs (Howad and Kempken, 1997) as well as post-transcriptional decrease of aberrant mitochondrial gene products (e.g. Gagliardi and Leaver, 1999; Menassa et al., 1999) and compensatory effects at the metabolic level (Cui et al., 1996; Liu et al., 2001) have been described. Tissue-specific differences in mitochondrial gene expression and in energy load are suggested explanations for the specific effects of mitochondrial dysfunction on male flower organs and pollen formation, respectively (e.g. Bergman et al., 2000; Conley and Hanson, 1995; Ducos et al., 2001). Therefore, the mitochondria have to impede with the function of certain, yet unrevealed nuclear genes during flower formation. Nuclear genome responsible for flower formation in plants, cytoplasmic male sterility indicates (CMS; Kaul, 1988) a participation of extra-nuclear genes in flower formation. Inability of CMS plants produce fertile pollen and hence are male sterile. The maternal inheritance of the CMS trait indicates that the genetic information of either the mitochondria or the plastids participates in certain steps of flower formation. Mitochondrial genes have been identified as being involved in all well-investigated CMS systems. In many cases of CMS, rearrangements in the mitochondrial genome were found to create unique chimeric genes, and often function of genes encoding subunits of the mitochondrial ATP synthase is impaired, suggesting that reduced mitochondrial respiration and/or ATP
production lead to male sterility (reviewed by Budar and Pelletier, 2001; Mackenzie and McIntosh, 1999; Schnable and Wise, 1998). The mitochondrial dysfunction of CMS plants can be either maintained or censored by specific nuclear gene functions leading, in the latter case, to restored male fertility. Genetic and molecular mechanisms of restoration vary among the different CMS systems (reviewed by Schnable and Wise, 1998). Elimination of CMS-related mitochondrial DNA segments (Janska et al., 1998), restoration of effective editing of specific mitochondrial mRNAs (Howad and Kempken, 1997) as well as post-transcriptional decrease of aberrant mitochondrial gene products (e.g. Gagliardi and Leaver, 1999; Menassa et al., 1999) and compensatory effects at the metabolic level (Cui et al., 1996; Liu et al., 2001) have been described.

Tissue-specific differences in mitochondrial gene expression and in energy demands are recommended explanations for the specific effects of mitochondrial dysfunction on male flower organs and pollen formation, respectively (e.g. Bergman et al., 2000; Conley and Hanson, 1995; Ducos et al., 2001). Consequently, the mitochondria have to impede with the function of certain, yet unknown nuclear genes during flower formation. Impairment in CMS plants in late stages of pollen formation without affecting female organs or general flower architecture. But rarely, like in the ‘petaloid’ CMS system of Daucus carota (De) (Kitagawa et al., 1994; Linke et al., 1999; Nothnagel et al., 2000), is the whole composition of flowers altered, frequently showing homeotic-like organ replacements. Similar floral malformations occur in alloplasmic lines of Nicotiana (Farbos et al., 2001; Kofer et al., 1991) and wheat (Murai et al., 2002) and in cybrids between Nicotiana and Hyoscyamus (Zubko et al., 2001). Exhibition of altered morphologically CMS flowers of petals and stamens (Eisa and Wallace, 1969; Straub, 1971). Petals have sepaloid features (Kitagawa et al., 1994) and stamens are converted into either petals (‘petaloid’-type) or carpel-like structures (‘carpeloid’-type), respectively. Characterisation of petaloid CMS system of carrot at the genetic level (Banga et al., 1964; Börner et al., 1995; Thompson, 1961; Welch and Grimball, 1947; Wolyn and Chahal, 1998).

A single dominant gene (M) and two homozygous recessive alleles (ll, tt) are anticipated to maintain male sterility. Heterozygosity of the dominant allele (Mm) and a dominant allele L or T can re-establish partial fertility at high temperatures (Börner et al., 1995 and references therein). Molecular analysis of the mitochondrial DNA and mRNA exposed differences between fertile and male-sterile plants including rearrangements of genes as potential causes for the CMS phenotypes (Börner et al., 1995; Nakajima et al., 2001; Scheike et al., 1992; Szklarczyk et al., 2000).

Investigation of nuclear mutants of Antirrhinum and Arabidopsis reveals similar homeotic organ alterations affecting two neighbouring flower whorls (Bowman et al., 1989; Schwarz-Sommer et al., 1990). Observation of CMS phenotypes in carrot lines containing cytoplasm from Daucus carota subspecies (Linke et al., 1999) but these cytoplasms have not yet been integrated in to cultivated carrots for hybrid breeding. Brown anther CMS was the first type used for developing hybrid carrots, but the majority of current hybrid carrot varieties utilize petaloid cytoplasms derived from two different sources of Sp cytoplasm, designated ‘Cornell’ (SpC; Thompson, 1961) and ‘Wisconsin Wild’ (SpW; Morelock et al., 1996). The SpC and SpW cytoplasms were found in Massachusetts and Wisconsin, respectively. Evaluation of SpC, SpW and
Canadian Sp cytoplasms with the cytoplasms of maintainer lines (N cytoplasms) using mitochondrial restriction length fragment polymorphisms (RFLPs) exposed that the Sp cytoplasms were very associated to each other and clearly distinct from N cytoplasms, which were found to be substantially more erratic (Bowes & Wolyn, 1998; Pingitore et al., 1989; Scheike et al., 1992; Steinborn et al., 1992). A chloroplast RFLP marker has been acknowledged among N cytoplasms of cultivated carrot (Vivek et al., 1999), but chloroplast markers that differentiate N from Sa and Sp cytoplasms have not been found (Scheike et al., 1992; Vivek & Simon, 1999).

Molecular markers that can envisage the CMS status at an early developmental stage will be essential tools in carrot breeding and seed making programs, as well as for basic studies of male sterility. Recently, six mitochondria-specific Sequence Tagged Site (STS) primer pairs derived from Randomly Amplified Polymorphic DNA (RAPD) markers were reported to distinguish SpC and SpW cytoplasms from a collection of three male fertile inbred carrot lines and five open pollinated varieties (Nakajima et al., 1999).

However, only two of these primer pairs amplified fragments that were constantly linked with either Sp or N cytoplasms. Because of the high degree of disparity surrounded by mitochondrial genomes of fertile carrots and the energetic nature of the mitochondrial genome of higher plants in general, several markers may be requisite to establish whether an uncharacterized cytoplasm confers petaloidy.

In addition, a range of mitochondria-specific markers will be useful to make out the mitochondrial genomes of putative cybrids consequential from protoplast fusions, and to illustrate a wide range of wild and cultivated carrot germplasm.

Carrots instead of petaloid type of CMS are not exceptional among plants. Similar abnormalities have been reported for a few other genera like Nicotiana, Brassica and Plantago. Most cases of CMS in tobacco resulted from interspecific hybridization and recurrent backcrossing placing Nicotiana tabacum nuclear genome in the perspective of cytoplasm from some wild Nicotiana species (Kaul 1988; Bonnett et al., 1991).

Mutations in definite nuclear genes may also condition the conversion of stamens into petals. Agamous—floral homeotic mutant of Arabidopsis—and plena—its equivalent from Antirrhinum—are by far the best studied instances of this phenomenon (Causer et al., 2010).

According to the ABC(DE) model of floral morphogenesis (Ferrario et al., 2004), both mutants correspond to impairment of the so-called C function and its replacement with activity of genes representing A function. Joint actions of the A, B and E functions indicate petal distinctiveness. Phenotypic similarities between petaloid CMS and nuclear homeotic mutations propose that the cytoplasm, by an as yet unknown mechanism, influences the expression of transcription factors controlling floral patterning (Linke et al., 2003).

It seems that cytoplasmic factors could act either through impairment of the C function or by enhancing the functions engaged in petal formation. In favor of the latter possibility, the combined ectopic expression of AP3, PI (both are B function genes) and AP1 (A function) as well as AP3, PI together with SEP3 (E function) converts vegetative leaves into petaloid organs (Honma and Goto 2001).
Basic steps in the use of cms

A major role played by three in the use of CMS system. female parent represented as A line with good floral and agronomic traits. It is mostly a line of preferred genetic background (genetically Srff) where s stands for sterile cytoplasm and rf for a recessive male sterility gene. To maintain A line, an isogenic line with the genetic background of A line known as B line (maintainer), is required. To employ this system for hybrid seed production, the third line commonly known as C line (male parent) is there which also acts as the paramount specific combiner with A-line. Genetically this could be N/S-Rf/- (fertility restoration) or Nrff since the economic product in cauliflower is not the exact seed. To carry out additional hybridization work protection of A, B and C line is the then important step. For hybrid seed creation, female to male ratios generally favored are 2:1; 3:1 or 4:1 depending on varietal characters.

Mechanism of cms

The occurrence of male sterility in wild species is revealed by the coexistence of hermaphrodite and female (male sterile) plants in natural populations (gynodioecy). Most of the studies on such populations have led to the emergence of theoretical models based on the premise that CMS is a result of interaction between nuclear and cytoplasmic genomes. Gynodioecy is supposed to appear and be maintained in populations either due to female advantage or counter selection of nuclear fertility restorers (Budar et al., 2003).

The genomic conflict theory (Cosmides and Tooby, 1981) provides for the interaction of cytoplasmic determinants (that prevent pollen production) and nuclear restorers (that restore fertility). The modes of inheritance of cytoplasmic (uniparental) and nuclear (biparental) factors generate a situation where selective interests are conflicting, resulting in differential resource allocation for pollen and seed production. Most of the CMS’s were a result of an alloplasmic state, wherein an impaired mitochondrial activity occurred due to a conflict at genomic, proteomic or transcriptomic levels (Kaul, 1988).

Studies on the genetic and physiological factors underlying male sterility have revealed that significant differences exist between normal and sterile anthers (Nair, 1993). The tapetum, the innermost cell layer surrounding pollen grain, is a physiologically active site and nourishes the developing gametes. However, it degenerates later on and is not found as an organized tissue in mature anthers. The tapetum and other surrounding cell layers suffer early degeneration in sterile pollen as compared to fertile ones. An important question was: how does a mitochondrial dysfunction at the genetic and physiological level specifically alter pollen development (Levings, 1993).

Since mitochondria harbor genes governing vital biochemical pathways such as the electron transport system and encode enzyme systems and structural RNA’s, how do such mutations specifically block pollen development, as mutations to any of them could be lethal? One explanation could be that mitochondrial genes are expressed differently during anther development. Such differential gene expression may occur either by the interaction of CMS genes with some unknown factors in target tissue leading to localized developmental changes or interference of CMS gene products in the physiological machinery resulting in reduced respiration and ATP synthesis (Levings, 1993).

Wallace (1989) suggested that minor impairments in mitochondrial function might
selectively affect degeneration of specific cell types. Such deviations in expression were observed in all cell and tissue types and have also been reported in case of human cardiac dysrhythmia (Wallace, 1988). Lee and Warmke (1979) observed a 10-fold increase in mitochondria/cell count in the tapetum during anther development. Such amplification indicates greater demand for energy.

Even though male sterility is strongly associated with gene dysfunction in mitochondria, chloroplasts have emerged as a more than ideal choice for engineering male sterility through the transgenic approach. Such an approach offers a number of advantages such as high levels of transgene expression, multi-gene cassette engineering, transgene containment viz., maternal inheritance, lack of gene silencing, position effect due to sitespecific transgene integration, and lack of pleiotropic effects (Daniell, 2005).

Moreover, such engineering strategy can enable the transfer of male sterility systems in certain recalcitrant crops such as cotton and soybean (Grevich and Daniell, 2005). It also overcomes the disadvantages of nuclear transformation, wherein the transformants segregate for fertility and sterility (Ruiz and Daniell, 2005).

**Plant mitochondrial mutations**

Variation in the mitochondrial genome of plant and animal of variable sizes to undergo recombination (Arrieta-Montiel and Mackenzie, 2010). This recombination activity in plants can subdivide the genome to a highly redundant, multi-partite structure, with different submolecules maintain at varying levels. Sites with the potential to undergo recombination can be numerous within the mitochondrial genome, with 47 repeat pairs functionally identified in the *Arabidopsis* mitochondrial genome (Arrieta-Montiel et al., 2009, Davila et al., 2011). This type of recombination provides an opportunity for accelerate genome evolution and capacity for extensive genomic variation.

Variation of different forms are particularly prevalent in the mitochondrial genomes of plants, sequence chimeras from non-homologous end joining, and express intermolecular genomic copy number changes within the mitochondrial population, formed by recombination and termed substoichiometric shifting (SSS). Mutually phenomena are key components of CMS systems.

Identification of all CMS mitochondrial mutations comprise of sequence chimeras, generally fusing identifiable mitochondrial gene sequences with segments of unknown origin (Schnable and Wise, 1998, Hanson and Bentolila, 2004). No two CMS mutations have been the same. Rearrangement of these sequences give rise to new gene products that usually show some degree of hydrophobicity and their interaction with the mitochondrial membrane. Various CMS sequences contain segments of genes involved in the ATPase complex simply that energy-associated alterations might participate in the CMS phenotype.

CMS mutations differ not only in their sequence origins, but often in their phenotype, from alterations in microspore development (Datta et al., 2002), to callose dissolution (Abad et al., 1995), to premature tapetal breakdown patterns (González-Melendi et al., 2008).

Whether these phenotypic differences are the consequence of distinct mechanisms of action or simply a reflection of variation in spatial and temporal expression patterns of the sterility lesion is not known.
Fertility restoration

suppression of male sterile phenotype by nuclear fertility restorer genes. Identification of valuable genes provide insight into the nature of mitochondrial gene regulation (Schmitz-Linneweber and Small, 2008). majority of cloned genes are fertility restorer genes them have been found to contain pentatricopeptide repeats (PPR) within the protein. participation of PPR proteins in mRNA splicing, editing, and translation initiation processes (Delannoy et al., 2007). this family of genes functions in both mitochondria and plastids and is, numbering over 400 in Arabidopsis (Small and Peeters, 2000). even if essential for the proper processing and translation of native organellar transcripts, PPR-mediated processing can serve to inactivate a rearranged, CMS-associated gene transcript.

Interestingly, regions of a genome may find PPR protein-coding genes organized in large duplicated gene clusters, representing fertility restorer loci (Barr and Fishman, 2010). association of these genes within plant genomes suggests that they are subject to extreme evolutionary selection, somewhat similar in pattern to disease-resistance genes (Geddy and Brown, 2007).

In naturally occurring CMS hybrid inappropriateness systems, where the male sterility phenotype can arise by particular hybridizations, evidence from PPR protein gene organization and maintenance across angiosperms recommend co-evolution of these nuclear and cytoplasmic factors to influence cross compatibility.

Two other types of nuclear fertility restorers have also been reported. The first is the R2 restorer of maize CMS-T cytoplasm, which encodes an aldehyde dehydrogenase that is recommended to play a detoxification role (Cui et al., 1996, Liu et al., 2001). a different type of nuclear restorer was reported in the common bean Fr gene, which effects thorough stoichiometric changes in mitochondrial genome organization and results in remarkable decrease of the mitochondrial male sterility sequence, pvs-orf239, to near imperceptible levels (Mackenzie and Chase, 1990). This latter fertility restoring mechanism is non-reversible in subsequent generations; therefore, it does not fall under the formal definition of a fertility restorer gene. However, such nuclear-directed influences on mitochondrial genome organization likely play an vital role in controlling mitochondrial genetic variation in natural plant populations.

Steadiness of the CMS trait

An fundamental feature of the CMS mutation for use in crop breeding is its stability, with little or no spontaneous reversion to fertility. degeneration to fertility is linked with SSS activity (Arrieta-Montiel and Mackenzie, 2010).

Recombination-mediated genomic changing can result in rapid copy number decrease of the DNA molecule on which the CMS mutation resides, departure the remainder of the mitochondrial genome essentially unchanged (Jansa et al., 1998, Bellaoui et al., 1998).

In this manner, the CMS mutation is silenced and the plant displays a male fertile phenotype. Reversion frequency in most CMS systems that have been studied appears to be influenced by nuclear genetic background (Mackenzie and McIntosh, 1999).

Therefore, a breeder wishing to employ a CMS system must test the strength of the system on a range of germplasm in the program.
Male sterility: Metabolic and evolutionary implications

Observation of CMS in over 150 plant species (Kaul, 1988), signifying that the competence to convert to male sterility is established in plants. In CMS common bean (Arrieta-Montiel et al., 2001) and Brassica spp. (Murayama et al., 2004, Giancola et al., 2007), fertile plants from wild populations contain the male sterility gene at substoichiometric levels.

These observations imply that it is feasible to interconvert male fertile and sterile types under particular, not yet defined conditions, identification of two nuclear genes that control the mitochondrial SSS process, MSH1 (Abdelnoor et al., 2003) and RECA3 (Shedge et al., 2007). Disruption or RNAi suppression of the nuclear gene MSH1 results in a marked increase in the frequency of mitochondrial recombination (Sandhu et al., 2007) and, interestingly, results in a CMS phenotype in six different plant species tested to date (Shedge et al., 2010; Sandhu and Mackenzie, unpublished).

This raises the intriguing possibility that CMS induction in nature can occur under conditions that suppress MSH1 expression. Interestingly, transcription of MSH1 is markedly downregulated under environmental stresses, including heat, drought, cold, and high light (Zimmermann et al., 2004, Shedge et al., 2010).

Occurrence of male sterility by numerous mechanisms in plants, the phenotype is apparently adaptive within a plant population under certain circumstances. Male sterility has been linked with increases in seed production and vigor as the consequences of out-crossing (Charlesworth, 2002). The occurrence of gynodioecy, or the mixture within a plant population of both hermaphrodites and females (male steriles), has been expected to occur in over 7% of all plant species (McCauley and Olson, 2008).

In various of these cases, the male sterility appears to be cytoplasmic, likely a outcome of the dynamic nature of the mitochondrial genome. Important questions enduring in this field turn around the nature of environmental and/or cellular cues that generate interconversion of male fertile and male sterile phenotypes in natural populations and in agricultural field conditions.

Organelle metabolism influences pollen development

There is support to propose that organelle metabolism, and overall cellular metabolic state, influences pollen development. Mutation of the plastidial glyceraldehyde-3-phosphate dehydrogenase, in the form of a gapcp1 gapcp2 double mutant, renders the plant male sterile (Muñoz-Bertomeu et al., 2010). Alterations in pollen mutant displays unusual collapsed shapes due to the tapetum and possibly changes in carbohydrate pools and/or trehalose level. Similarly, interruption of cytosolic phosphoglucomutase prevents both male and female gametogenesis, signifying that proper carbohydrate allotment is critical at these stages (Egli et al., 2010).

Temperature has a major communicate with on pollen development and viability. Cold temperature stress can hamper pollen development, and male-gametophyte-specific classes of transcription factors appear to influence pollen response to adverse temperature effects (Zou et al., 2010).

The programming of tapetum for an apoptosis-like development (Wang et al.,
expressing BoCysP1 showed swelling and vacuolation, perhaps intervention with the normal pattern of programmed cell death.

Realization of male sterility in agricultural systems

Genetic engineering attempts of male sterility, often useful in suppressing or disrupting normal pollen development patterns, have generally suffered from their infeasibility to scale up for agricultural implementation. An important motivation for identifying an effective genetic system for male sterility is the opportunity to cultivate hybrids. An F1 hybrid provides more uniform crop performance in the field and, more importantly, the significant advantage of heterosis.

Heterosis, or hybrid vigor, refers to the ability of the F1 hybrid to outperform either of the parents used in the cross. Consequently, development of hybrids has, in the case of many crops, accounted for significant gains in overall crop yield. For example, hybrid rice in China can be credited with reducing the total amount of land planted to rice from 36 Mha in 1975 to 30 Mha in 2000, and increasing total production from 128 to 189 million tons, a yield increase of up to 6.2 tons/ha (http://www.fao.org/rice2004).

Several different genetic models proposed to report for heterosis, including dominance, overdominance, and pseudo-overdominance, all appearing to have some merit (Chen, 2010). Suggestion has been given on that epigenetic factors, influenced by circadian-controlled regulatory mechanisms, may underlie much of the heterosis phenomenon.

However, what has not been investigated sufficiently is the influence of maternal effects on hybrid vigor. Often reciprocal
differences in cross-compatibility and vigor outcomes will be observed in a cross. There are several cases in CMS plant crossed to its sterility maintainer line will create a more vital progeny than self-pollination of the identical maintainer line (S. Mackenzie, unpublished). In these crosses, the two hybridizing plants are almost indistinguishable in nuclear genotype, implying that the vigor may result from cytoplasmic factors. In fact, very little research has been conducted to inspect the possible authority of mitochondria or plastids, the two essential generators of ATP, on heterosis.

It is appealing to consider that organellar metabolic factors, together with epigenetic signals, may integrate to enhance energy in the hybrid. Organellar changes arising as a outcome of mutation or inhibition of MSH1 in Arabidopsis and other crop species give rise to remarkable changes in growth patterns and transcript profiles, suggesting that organellar functions can directly persuade several developmental pathways in plants (Shedge et al., 2007, Shedge et al., 2010, Sandhu et al., 2007; Sandhu and Mackenzie, unpublished).

The natural system of cytoplasmic male sterility remains the most generally utilized agriculturally. CMS has been effectively implemented in a number of crops, including rice, sorghum, millet, onion, sugar beet, and carrot, but restrictions exist with this genetic system as well. Following the Southern Corn Leaf Blight of 1970, when a huge area of the Midwestern United States was planted with CMS-T type maize that proved vulnerable to the T-toxin of Bipolaris maydis (Ullstrup, 1970), maize CMS is no longer used to such a great extent.

What was erudite from that episode was the need for multiple distinct, stable sources of CMS for any given crop to avoid overplanting to a single cytoplasm type. Furthermore, a useful source of CMS has not yet been identified in many crops in which hybrid vigor is well renowned. With the recent demonstration that CMS can be transgenically induced (Sandhu et al., 2007), the opportunity may now exist for detection of additional CMS lines. apparently, this type of cytoplasmic genetic manipulation can be carried out in various species and in several distinct genotypes. If MSH1 modulation occurs naturally in plant populations, and the CMS arising in these manipulations also occurs in nature, we can imagine that fertility restoration mechanisms are also relatively widespread.

The accessibility of essential information on the genetic and physiological pathways variable stamen development provides well-documented opportunities to persuade these processes transgenically. More practical strategies for introduction of male sterility traits to large-scale agricultural production are needed now.

**Advantages of cms over other systems**

In carrot, F1 hybrids are valuable due to uniform maturity, high early and total yield, better curd quality with respect to curd compression and colour as well as for resistance to insect-pests, diseases and adverse weather conditions. An efficient, reliable and established method of F1 seed production without infectivity by self-fertilized seeds from each parent is vital.

Manual emasculation and pollination method in carrot is not useful at commercial level due to incompatible flower size and structure. So far hybrids in carrot have been developed using self-incompatibility (SI) system. Several reports concerning
self-incompatibility in Indian carrot which revealed that inbreds_LINES of maturity group I have strongest self-incompatibility followed by maturity group II and group III. In the case of the snowball cauliflower group, self-incompatibility is either very weak or not present at all. Moreover, SI system poses a risk of occurrence of selfing in hybrid seeds besides problems of maintenance, multiplication of SI lines through tedious bud pollination or CO2 and NaCl spray and breaking down of SI under different environmental conditions like high temperature and drought. Even in case of genic male sterility (GMS), roguing of fertile individuals from the female line at the time of anthesis poses a main problem.

To overcome such problems, CMS system offers a better alternative. Efficacy of CMS-based hybrids to exploit heterosis has been established by. In recent years, important heterosis has also been reported in hybrids developed using CMS system for yield, yield linked and quality traits.

There are three types of male sterility in carrot:
- Brown anther (ba) (Welch and Grimbal, 1947)
- Petaloid (pt) (Thompson, 1962)
- Gummifer (Nothnagel, 1992)

**Petaloid male sterility**

Petaloid sterility is commercially used for hybrid seed production in the world. According to Morelock (1974), the pt type of male sterility is due to interaction between “Sp cytoplasm” and two independent dominant genes (M1 and M2). This is manifested as the replacement of stamens with petals (white petaloidy) or both stamens and petals with green bract like structures (green petaloidy) (Kitagawa, *et al.*, 1994). It is stable across a wide range of environments through flowering and seed production. CMS system has been established for the first time in Asiatic carrot germplasm in India at IARI.

In India, at IARI, petaloid CGMS was transferred to nantes type and crossed it with the indigenous variety “Pusa yamagadi” to develop hybrid Pusa nayanjyoti in 2009. Cytoplasmic male sterility system has been established for the first time in tropical carrot and tropical carrot hybrid “Pusa Vasudha” has been developed which is the first from any public sector institution.

**Mechanisms of male sterility**

**Cytological changes**

Breakdown in the process of microsporogenesis can occur at a pre-or postmeiotic stage. The abnormalities can involve aberration during the process of meiosis in the formation of tetrads, during the release of tetrad (the dissolution of callose), at the vacuolate microspore stage or at mature or near-mature pollen stage.

**Biochemical changes**

Male sterility is accompanied by certain biochemical changes such as qualitative and
quantitative changes in amino acids, protein, and enzymes in developing anther. It has been found to be associated with reduced levels of proline, leucine, isoleucine, phenylalanine and valine and increased levels of glycine, arginine and aspartic acids (Kaul, 1988). Level of proline has been found to be particularly affected. Kakihara et al., (1988) reported that mature male sterile anthers contain one-eighth amount of proline in comparison to the fertile anthers. Anthers of male sterile plants contain lower soluble protein content and fewer polypeptide bands. Some polypeptides synthesized in normal stamens were absent in mutant stamens. The activity of enzyme callase which is required for breakdown of callose that surrounds PMCs and the tetrad is decreased in male sterile anthers.

Low activity callase leads to premature or delayed release of meiocytes and microspore, resulting in male sterility. Male sterility is also accompanied by decreased activity of isozyme esterase which plays an important role in hydrolysis of sporopollenin polymer that is required for pollen formation.

Decreased activity of esterase in male sterile plants has been observed in tomato (Bhadula and Sawhney, 1987) and radish (Zhou and Zhang, 1994). Compared to fertile lines, there is reduced activity of amylase in male sterile anthers and it corresponds with high starch content and reduced levels of soluble sugars.

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