Complete mitochondrial genome sequencing and identification of candidate genes responsible for C5-type cytoplasmic male sterility in cabbage (B. oleracea var. capitata)

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Cytoplasmic male sterility (CMS) is widely used in cruciferous vegetables hybrid breeding. The C5-type CMS cabbage line exhibits stable male sterility and offers great value for cabbage breeding. However, the underlying CMS mechanism remains unclear. Here, the complete mitochondrial genome was sequenced and assembled for this line. The genome size was 221,862 bp. Mitochondrial genome comparison showed that the mitochondrial genome was likely generated by recombination with a nap-type CMS B. napus strain. Sixty-seven unknown-function open reading frames (ORFs) were identified. Seven orfs, orf114a, orf123a, orf188a, orf222a, orf261a, orf286a, and orf322a, were specifically identified in this genome. The presence of these candidate CMS genes decreased ATPase activity and ATP content by affecting the transcript levels of energy metabolism-related genes and F1F0-ATP synthase assembly. Among them, orf188a, orf222a, orf261a, orf286a, and orf322a possessed a transmembrane structure, and orf188a was cotranscribed with rps7 and trnM. orf222a was partially homologous to atp8 and coexpressed with nad5. orf261a and orf322a were cotranscribed with cox1 and atp9, respectively. Additionally, orf114a was cotranscribed with atp8. Yeast two-hybrid assays showed that the ORF222a protein interacts with a B. oleracea ATP17 homolog (Bo7g114140) during F0-type ATP synthase assembly, reducing the quantity and activity of assembled F1F0-ATP synthase. Cytological sections showed that premature separation of the tapetum from the connective tissue and delayed tapetal programmed cell death (PCD) might be the immediate causes of CMS in C5-type CMS cabbage lines. Our results may help uncover the molecular mechanism of C5-type CMS in B. oleracea from the perspectives of the whole mitochondrial genome and cytology of anther development.
Introduction

Cabbage (Brassica oleracea var. capitata) is one of the most important vegetables and is grown on five continents around the world (Chiang et al., 1993). In the last few years, approximately 70 million tonnes of cabbage have been produced per year worldwide; according to the Food and Agriculture Organization of the United Nations (FAO), China is the largest cabbage producer in the world, producing approximately 34 million tonnes in 2020. Heterosis has been widely applied in cereal crop and vegetable production (Yu et al., 2021). Cytoplasmic male sterility (CMS) is an extranuclear maternally transmitted trait that produces either aborted or infertile pollen grains (Dong et al., 2013). CMS lines, along with maintainer and restorer lines, are highly valuable resources for F1 hybrid seed production, which has been widely developed and applied to all Brassicaceae crops (Yamagishi and Bhat, 2014; Yang et al., 2021).

In Brassica species, a variety of CMS types have been independently reported, such as Ogu, nap, Nig, tour, Shaan2A, pol, Nsa, hau, inap, and oxal. Among them, Ogu-CMS was first identified in Raphanus sativus (Ogura, 1967); Nig CMS comes from Brassica nigra (Pearson, 1972); nap CMS, Shaan2A CMS and pol CMS are derived from Brassica napus (Shiga and Baba, 1971; Li, 1980; Fu, 1981); Nsa CMS is an alloplasmic male sterility system derived from somatic hybridization between Brassica napus and Sinapis arvensis (Hu et al., 2002); inap CMS was obtained from somatic hybridization between Brassica napus and Isatis indigotica by recurrent backcrossing (Kang et al., 2017); and tour CMS, hau CMS and oxal CMS originated from Brassica juncea (Rawat and Anand, 1979; Wan et al., 2008; Heng et al., 2018). The Nig, tour, pol and Ogu CMSs were transferred into B. oleracea by protoplast fusion or the interspecific cross; however, the cybrids exhibited poor agronomic performance except for Ogu-CMS system (Pearson, 1972; Bannerot H et al., 1974; Yarrow et al., 1990; Cardi and Earle, 1997). To date, the Ogu-CMS system has been used as the major CMS system for F1 seed production in B. oleracea crops because of its stable sterility and lack of adverse effects on plant growth (Dey et al., 2013). However, the broad use of a single CMS results in a high degree of cytoplasmic genetic uniformity, which may lead to genetically vulnerable plants (Yamagishi and Terachi, 2017). Therefore, it is urgent to develop more CMS systems for cabbage breeding.

Thus far, many CMS-related genes have been identified and reported in Brassica species. Most open reading frame (ORF)-encoded proteins in CMS lines disrupt energy metabolism, which disrupts the $F_1F_0$-ATP synthase subunit. The molecular structure of the $F_1F_0$-ATP synthase complex from Saccharomyces cerevisiae has already been built (Srivastava and Luo, 2018). However, the structure of this complex in plants has not been resolved. Plant $F_1F_0$-ATP synthase shares the basic structure of the enzyme complexes described in yeast. The water-soluble complex ($F_0$) component possesses 3 copies each of subunits a (ATP1) and β (ATP2) and one copy each of subunits γ (ATP3), δ (ATP16) and ε (ATP15). The membrane complex ($F_1$) is composed of subunits α (ATP1) and ATP6-2), b (ATP4), 8 (ATP8), c (ATP9), f (ATP17), g (ATP20) and e (ATP21) (Zancani et al., 2020). Subunits a (ATP6), b (ATP8), and c (ATP9) have been extensively investigated in most CMS types. Several previous findings showed that products of candidate CMS-related genes often affect mitochondrial functions via interacting with the other essential nuclear/mitochondria-encoded proteins. For examples, ORFH79 impairs mitochondrial function via interaction with P61, which is a subunit of electron transport chain complex III in the Wild Abortive CMS (CMS-WA) rice line (Luo et al., 2013). The mitochondrial protein WA352 interacts with COX11 to inhibit its function in peroxide metabolism, which induces the premature tapetal PCD and triggers consequent pollen abortion in the Wild Abortive CMS (CMS-WA) rice line (Luo et al., 2013). ORF224 was found to be associated with a respiratory electron transport chain protein (BnaC03g14740D), which affects the development of the downstream gene nad3 and rps12 complex (Sang et al., 2021). The sterility gene orf138, which triggers Ogu-CMS, is cotranscribed with atp8 and trnfM (Tanaka et al., 2012; Zhong et al., 2021). The sterility gene orf224 is responsible for pol CMS, which is cotranscribed with atp6 (Singh and Brown, 1993). orf222, the key gene of nap CMS, is similar to the pol CMS-associated gene orf224 and the cotranscribed genes nad5c and orf139 (Lhomme et al., 1997). orf288, the master gene of hau CMS, is also cotranscribed with the downstream gene atp6 (Heng et al., 2014). The key sterility gene orf346, which controls Nsa CMS, is cotranscribed with the nad3 and rps12 complex (Sang et al., 2021). The sterility gene orf193, which triggers tour CMS (originating from cell fusion between B. napus and B. tournefortii), is cotranscribed with the downstream gene atp9-2 (Dieterich et al., 2003). However, orf263, the causal gene for tour CMS (obtained from sexual hybridization between B. napus and B. tournefortii), is...
cotranscribed with the downstream gene atp6 (Landgren et al., 1996). In total, the expression of specific mitochondrial genes is highly dependent on the species in which the CMS source originated.

In the present study, C5-type CMS was identified as a novel alloplasmic male-sterility system derived from intergeneric hybridization of B. napus and B. oleracea. To provide new insights into the molecular mechanism of this C5-type CMS of cabbage exhibiting anther indehiscence and no pollen dispersal, we sought to identify CMS-associated candidate genes in mitochondria using a next-generation sequencing approach and accordingly analyzed the specific ORFs. The complete mitochondrial genomes of C5-type CMS and its maintainer line were sequenced. To deeply explore the abortive mechanisms of the C5-type CMS in cabbage, we investigated the characteristics of maintainer and C5-type CMS cabbage lines in cytological, physiological, and molecular analyses. To ascertain the responsible gene and the underlying mechanisms for C5-type CMS, we characterized C5-type CMS-associated candidate genes. The mitochondrial genome comparison results indicated that the C5-type CMS original cytoplasm donor material was most likely generated by recombination with the nap-type CMS B. napus strain through intergeneric hybridization during the breeding process. The CMS candidate protein ORF222a interacts with the ATP17 homolog (Bo7g114140) in B. oleracea, impairing the step of F1Fo-ATP synthase assembly, which results in remarkable reductions in the ATP and ATPase activity levels in the anther samples of C5-type CMS cabbage lines compared with those of the maintainer lines. Cytological analysis of anther development between the maintainer and C5-type CMS cabbage lines showed that premature separation of the tapetum from the connective tissue and abnormal degradation of tapetal cells might be the immediate cause of CMS in C5-type CMS cabbage lines.

Materials and methods

Plant materials

The maintainer line was an open-pollinating and early-maturing cabbage variety. The original cytoplasmic donor cabbage material was introduced from B. napus nap-type cytoplasm by intergeneric hybridization. The C5-type CMS cabbage line was obtained from crosses and consecutive backcrosses with B. napus in 2005 by Dr. Jungen Kang from the Beijing Vegetable Research Center of the Beijing Academy of Agriculture and Forestry Sciences (BAAFS). The stability of C5-type CMS was observed for more than 10 years. The mitochondrial DNA of both the C5-type CMS and maintainer lines was sent to Biozeron Company (Shanghai, China) for DNA library construction and sequencing in 2018.

Mitochondrial genome sequencing and assembly

One microgram of purified mitochondrial DNA was fragmented for 300-500 bp paired-end library construction using a TruSeq™ Nano DNA Sample Prep Kit. Sequencing was performed on the Illumina HiSeq 4000 platform (BIOZERON Co., Ltd., Shanghai, China). A DNA library with approximately 15-20 kb SMRTbell libraries was constructed and sequenced on a PacBio Sequel Sequencer (PacBio Inc., Menlo Park, CA, USA). For the cabbage mitochondrial genome assembly, the filtered Illumina HiSeq subreads were preliminarily assembled by ABySS v2.0.2 software (version 1.5) (Jackman et al., 2017). Then, the PacBio Sequel data were aligned by the blasR method for single-molecule sequencing data correction. The corrected PacBio Sequel data and Illumina HiSeq data were combined for the mitochondrial genome framework assembly using SPAdes v3.10.1 software (Antipov et al., 2016). Finally, clean Illumina HiSeq reads were mapped to the assembled mitochondrial genome to verify the accuracy of the sequence. The circular genome maps were drawn using OrganellarGenomeDRAW (version 1.2) (Greiner et al., 2019).

Gene prediction and annotation

Mitochondrial genes were predicted based on a combination of homology-based gene prediction and de novo prediction by GeneWise and AUGUSTUS software. The tRNA and rRNA genes were predicted using tRNAscan-SE (version 2.0) and rRNAmer (version 1.2), respectively (Lowe and Eddy, 1997; Lagesen et al., 2007). The functions of the predicted proteins were annotated based on a BLASTP search against universal databases, such as the National Center for Biotechnology Information (NCBI) database, the Gene Ontology (GO) database (Ashburner et al., 2000), the Evolutionary Genealogy of Genes: Non-supervised Orthologous Groups (eggNOG) database (Jensen et al., 2008), and the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (Kanehisa et al., 2004). The transmembrane domains in each candidate ORF-encoded protein were assessed using TMHMM Server v.2.0 (http://www.cbs.dtu.dk/services/TMHMM/).

ATP content and synthase activity measurement

The ATP content and synthase activity were measured in the anthers of the maintainer and C5-type CMS lines with an ATP content assay kit and a Na`K`-ATP synthase activity assay kit, respectively (Solarbio Co., Ltd., Beijing, China). The procedures were performed according to the manufacturer’s instructions (Solarbio, BC0300 and BC0065).
Y2H assay

The full-length coding sequences (CDSs) of atp4, atp6, atp8, atp9, and two atp17-homologous genes (Bo3g175820 and Bo7g114140) were cloned into pGADT7 vectors; orf188a, orf222a and orf261a were cloned into the pGBKKT7 vector. The eighteen candidate interaction combinations were transformed into the yeast strain Y2HGold. Y2H assays were performed according to the Clontech manual. The transformed yeast cells were inoculated onto synthetic defined medium without leucine and tryptophan (SD/-Leu/-Trp) and incubated at 30°C for 2-3 days. The single transformation colony was then gradient-diluted and inoculated onto SD/-Leu/-Trp/-His-Ade media and cultured for 3–5 days at 30°C to observe strain growth.

Cytological analysis

Flower buds from the maintainer and C5-type CMS lines were collected at six different developmental stages and immediately fixed in formalin-aceto-alcohol (FAA) solution. Paraffin sections were then prepared according to the method of Zhong et al. (2021). Here, anther transverse sections were stained in 1% toluidine blue solution and viewed under a Leica (Tokyo, Japan) optical microscope (M2030). The pistils of the C5-type CMS line exhibited a sterile phenotype due to anther indehiscence and no pollen dispersal. However, the sterile line produced normal pollen grains (Figure 1).

Cloning of CMS-associated candidate ORFs and subunit genes of F1Fo-ATP synthase

Total RNA was isolated from the anthers of maintainer and C5-type CMS lines using an RNA extraction kit (RNAprep Pure Plant Kit, Takara, RR037A, Dalian, China). First-strand cDNA was synthesized using a PrimeScript™ RT Reagent Kit (Takara, RR037A, Dalian, China). Subunits b (atp4), a (atp6), 8 (atp8), and c (atp9) were cloned from cDNA of the C5-type CMS line; however, the subunit f (ATP17) homologs (Bo3g175820 and Bo7g114140) were cloned from cDNA of the maintainer line. Cloning PCR was performed using the primers listed in Supplementary Table S1. The amplified PCR products were detected using gel electrophoresis (1.5% agarose gel).

Alignment and phylogenetic analysis

Geneious Prime software was used to perform multiple sequence alignment between B. napus strain 51218 and our C5-type CMS mitochondrial sequences to obtain consistent sequences. Seven mitochondrial genome sequences of B. napus lines (B. napus strain SW18: AP018473 and AP018474, strain 51218: KP161618, cultivar 088018: MW348924, cultivar NY18: MW001149, and cultivar Westar: AP006444) were obtained from the NCBI database (https://www.ncbi.nlm.nih.gov/genome/browse/). Mitochondrial genome-wide alignments and ML bootstrap analysis with 500 replicates were performed using MEGA 6.0 (Tamura et al., 2013).

Results

Flower morphology of the C5-type CMS line and maintainer lines

The sterile flowers of the C5-type CMS line were significantly smaller than the fertile flowers of the maintainer line. The filaments and anthers of fertile flowers were remarkably longer than those of sterile flowers. There were no differences in the pistils between the C5-type CMS and maintainer lines. The C5-type CMS line exhibited a sterile phenotype due to anther indehiscence and no pollen dispersal. However, the maintainer line produced normal pollen grains (Figure 1).

Mitochondrial genome sequencing and annotation of the C5-type CMS cabbage line

The mitochondrial genomes of the C5-type CMS cabbage line were sequenced with Illumina HiSeq and PacBio Sequel techniques. The Illumina HiSeq sequencing platform generated 6791 Mb of clean data. Then, 47.85 Mb of subread bases were produced for the sample in the PacBio sequencing platform. The mitochondrial genome of the C5-type CMS cabbage line was assembled into a single circular mapping molecule with a size of 221,862 bp (GenBank no. ON960289) (Figure 2). The G+C content of the C5-type CMS mitochondrial genome was 45.19%, which was comparable to that of our previously released mitochondrial genomes from maintainer and Ogura-type CMS cabbage lines. Additionally, the C5-type CMS mitochondrial genome possessed a total of 125 genes, including 99 protein-coding genes (32 known genes and 67 unknown-function ORFs), 3 rRNA genes (5S, 18S, and 26S), and 23 tRNA genes. A comparison between the maintainer and C5-type CMS mitochondrial genomes revealed that most known mitochondrial genes were identical. The C5-type CMS mitochondrial genome comprised three genes encoding the large subunits of ribosomal proteins (rpl2, rpl5, and rpl16), four genes encoding the small subunits of ribosomal proteins (rps4, rps7, rps12, and rps14), nine subunits of NADH dehydrogenase (nad1, nad 2, nad 3, nad4, nad4L, nad5, nad6, nad7, and nad9), three cytochrome oxidase subunits (cox1, cox2, and cox3), one cytochrome b (cob), five subunits of ATP synthase (atp1, atp4, atp6, atp8, and atp9), five cytochrome C
synthesis-related genes (ccmB, ccmC, ccmFN1, ccmFN2, and ccmFC), and one maturase gene (matR).

Collinearity analysis of the mitochondrial genomes

To further analyze the structure and composition of the C5-type CMS cabbage mitochondrial genome, collinearity analysis was performed with the maintainer line. Our sequencing results showed that the mitochondrial genome size of the C5-type CMS line (221,862 bp) was 1,900 bp larger than that of its maintainer line (219,962 bp). In addition, a total of 9 syntenic regions (named blocks 1-9) were identified in the mitochondrial genomes of the maintainer and C5-type CMS cabbage lines on the basis of sequence homology (Figure 3A). These regions ranged from 2,317 to 66,388 bp in length and accounted for 95.85% of the mitochondrial genome sequence (Supplementary Table S2). The corresponding blocks between C5-CMS and its maintainer line had at least 99.22% identity (Supplementary Table S2). Although all 9 syntenic regions shared high sequence similarity, the directions and positions of these syntenic regions differed between the two mitochondrial genomes. The orientation of seven regions was identical, but in the other two (block 2 and block 3), the blocks were inversions. Block 4, block 8, and block 9 were translocations (Figure 3). The syntenic regions were largely discrepant in the two genomic positions, although the genetic sequences were well conserved. Recombination and rearrangement events during the breeding
process were determined to be responsible for the structural differences in the mitochondrial genomes of the same species. Block 4/block 2 and block 3/block 5 in the C5-CMS mitochondrial genome were broken by unique region I and unique region II, which were nonhomologous to the regions in the maintainer mitochondrial genome (Figure 3).

Analysis of the unique regions in the C5-type CMS mitochondrial genome and phylogenetic analysis of mitochondrial genomes

The collinearity analysis results revealed two unique regions in the C5-type CMS mitochondrial genome (Figure 3A). The sizes of unique region I and unique region II were 4,897 bp and 4,141 bp, respectively. The total length of the unique regions was 9,038 bp, accounting for 4.07% of the whole C5-type CMS mitochondrial genome. At the nucleotide level, BLASTn analysis showed that the unique regions shared high similarity with mitochondrial sequences of *B. napus* strain SW18 (AP018473 and AP018474), strain 51218 (KP161618), cultivar 088018 (MW348924), cultivar NY18 (MW001149), cultivar 56366 (KM454975), and cultivar Westar (AP006444) (Figures 4A, B). The results suggested that the C5-type CMS cytoplasm may have been derived from certain *B. napus* CMS haplotypes. Furthermore, phylogenetic analysis was performed using eight complete mitochondrial genomes. Whole-genome-wide alignments and maximum likelihood (ML) bootstrap analysis with 500 replicates were
performed using MEGA 6.0. As shown in Figure 4C, the phylogenetic tree revealed the topological structure of eight selected taxa, which were mainly divided into three clades. The C5-type CMS mitochondrial sequence formed Clade I with *B. napus* strains 51218 and Westar, which are *nap*-type strains of *B. napus*. Clade II comprised the maintainer cabbage line and *B. napus* strains 56366 and NH12A, which are *pol*-type *B. napus*. Another two *nap*-type *B. napus* cultivars, 088018 and NY18, constituted Clade III. Four *nap*-type accessions were clustered into two different clades, indicating that the investigated *nap*-type *B. napus* strains have a polyphyletic maternal origin. To further compare the mitochondrial genomes between C5-type CMS and *B. napus* cultivars 51218 and Westar, dot plot analysis and pairwise alignment were performed with Geneious Prime software. The results showed that the sequence of the C5-type CMS mitochondrial genome was more homologous to that of the *B. napus* cultivar 51218 than to that of Westar. However, there was an inversion located at the position from 80,500 bp to 93,000 bp, and many single-nucleotide polymorphisms (SNPs) were distributed throughout the genome (Figure 5). Taken together, these results show that the C5-type CMS original cytoplasm donor material was most likely generated by intergeneric hybridizations with certain *nap*-type CMS *B. napus* strains during the breeding process.

**Identification of CMS-associated ORFs in the C5-type CMS mitochondrial genome**

To reveal the genes determining the CMS phenomenon, we compared the mitochondrial genomes between the C5-type CMS line and its maintainer lines. Genome rearrangement events may have caused the presence of variation in genome sequence segments. Specific genes were generated by recombination, which is generally responsible for the CMS trait. Seven ORFs that encoded over 100 amino acids, including orf114a, orf123a, orf188a, orf222a, orf261a, orf286a, and orf322a, were specifically identified in the C5-type CMS mitochondrial genome (Table 1). All of these specific ORFs were candidate CMS genes. To further verify whether the proteins encoded by these ORFs were candidate proteins for CMS, the structures of these three proteins were predicted. We found that ORF188a, ORF222a, ORF261a, ORF286a, and ORF322a had transmembrane domains, whereas ORF114a and ORF 123a lacked transmembrane domains (Figure 6).
Organization of mitochondrial genome regions associated with candidate sterility genes for C5-type CMS

A more detailed analysis of the organization of mitochondrial genome regions associated with candidate sterility genes for C5-CMS was carried out. $orf_{188a}$ was located in C5-type CMS unique region I 651 bp downstream of $rps7$, and $orf_{222a}$ was located between C5-CMS block 4 and unique region I (Supplementary Table S3 and Figure 7A); however, $orf_{261a}$ was located 219 bp downstream of $coxl$ between C5-CMS unique region II and block 5 (Supplementary Table S3 and Figure 7A). $orf_{322a}$ was located 311 bp downstream of $atp9$ between block 8 and block 3, and $orf_{114a}$ and $orf_{123a}$ were located 465 bp downstream and 961 bp upstream of $atp8$ within block 1, respectively (Supplementary Table S3, Figures 7B, C). However, $orf_{286a}$ was located 1,324 bp upstream of $atp8$ between block 1 and block 4 of the C5-type CMS mitochondrial genome. Taken together, the findings indicate that the CMS candidate genes $orf_{114a}$, $orf_{188a}$, $orf_{222a}$, $orf_{261a}$, $orf_{286a}$, and $orf_{322a}$ were specifically identified in the C5-type CMS mitochondrial genome. They are identical to the mitochondrial proteins of Brassica napus. $orf_{114a}$ and $orf_{123a}$ were located in C5-type CMS block 1, $orf_{188a}$ was located in C5-type CMS unique region I, $orf_{222a}$ was located between C5-type CMS block 4 and unique region I, $orf_{261a}$ was located between C5-type CMS block 4 and unique region I, $orf_{286a}$ was located between C5-type CMS unique region II and block 5, and $orf_{322a}$ was located between block 8 and block 3.

### TABLE 1 Specific ORFs in the Ogura-CMS mitochondrial genome.

| Specific ORFs | Most similar mitochondrial sequence of another species | Location in the C5-type CMS line |
|---------------|------------------------------------------------------|---------------------------------|
| $orf_{114a}$  | YP_717103.1 hypothetical protein BnapMp004 [Brassica napus] | Block 1 |
| $orf_{123a}$  | AKD00165.1 hypothetical protein [Brassica napus] | Block 1 |
| $orf_{188a}$  | YP_717121.1 hypothetical protein BnapMp023 [Brassica napus] | Unique region I |
| $orf_{222a}$  | YP_717120.1 hypothetical protein BnapMp022 [Brassica napus] | Between block 4 and unique region I |
| $orf_{261a}$  | YP_717164.1 hypothetical protein BnapMp067 [Brassica napus] | Between unique region II and block 5 |
| $orf_{286a}$  | YP_717106.1 hypothetical protein BnapMp007 [Brassica napus] | Between block 1 and block 4 |
| $orf_{322a}$  | YP_717145.1 hypothetical protein BnapMp048 [Brassica napus] | Between block 8 and block 3 |

Seven ORFs, including $orf_{114a}$, $orf_{123a}$, $orf_{188a}$, $orf_{222a}$, $orf_{261a}$, $orf_{286a}$, and $orf_{322a}$, were specifically identified in the C5-type CMS mitochondrial genome. They are identical to the mitochondrial proteins of Brassica napus. $orf_{114a}$ and $orf_{123a}$ were located in C5-type CMS block 1, $orf_{188a}$ was located in C5-type CMS unique region I, $orf_{222a}$ was located between C5-type CMS block 4 and unique region I, $orf_{261a}$ was located between C5-type CMS unique region II and block 5, $orf_{286a}$ was located between block 1 and block 4, and $orf_{322a}$ was located between block 8 and block 3.
FIGURE 6
Transmembrane domain prediction of candidate CMS gene-encoded proteins in the C5-CMS line. The output of the TMHMM server indicated the location and probability associated with the predicted transmembrane domains in the C5-CMS line. (A) ORF188a, (B) ORF222a, (C) ORF261a, (D) ORF286a, and (E) ORF322a.
orf261a, and orf322a are cotranscribed with atp8, rps7, cox1, and atp9, respectively.

**ORF222a affects F1F0-ATP synthase assembly by interacting with an ATP17 homolog (ATP17-2, Bo7g114140) in B. oleracea**

To determine whether candidate CMS-associated ORF-encoded proteins (ORF188a, ORF222a, and ORF261a) were capable of binding to subunits of F1F0-ATP synthase, a yeast two-hybrid (Y2H) assay was performed with ORF-encoded proteins (ORF188a, ORF222a, and ORF261a) and subunits (ATP4, ATP6, ATP8, ATP9, and ATP17) of F-ATP synthase. orf188a, orf222a, and orf261a were cloned into the PGBK7 vector; meanwhile, atp4, atp6, atp8, atp9, and two atp17-homologous genes (atp17-1, Bo3g175820 and atp17-2, Bo7g114140) of F-ATP synthase were cloned into PGADT7 vectors. Eighteen candidate interaction combinations were tested with the Y2H assays. The results showed that only ORF222a interacted with the ATP17 homolog (ATP17-2, Bo7g114140) in B. oleracea (Figure 8 and Supplementary Figure S2). We thus conclude that ORF222a impairs the step of F1F0-ATP synthase assembly by interacting strongly with the ATP17 homolog (ATP17-2, Bo7g114140) in B. oleracea.

**Detection of ATP production in the C5-type CMS cabbage line**

From our whole-mitochondrial genome data, we found that the C5-type CMS-related gene orf188a encodes the atp6-like gene and that orf222a encodes 58 amino acids identical to ATP8 at the N-terminus (Supplementary Figure S1). We also found that orf188a, orf261a, and orf322a are cotranscribed with rps7, cox1, and atp9, respectively. Furthermore, ORF222a interacts with an ATP17 homolog (Bo7g114140) in B. oleracea. Therefore, we wondered whether these events influence the yield of ATP. We measured the ATP content and ATPase activity in anther samples from maintainer and C5-type CMS cabbage lines. We found that the ATP content and ATPase activity in the anther samples of the C5-type CMS cabbage line was remarkably lower than that in the maintainer line (Figure 9). The change in ATP yield may have been caused by altered ATPase activity. In conclusion, CMS-related genes encode partial fragments identical to mitochondrial genes with known functions and are cotranscribed with or interact with these mitochondrial electron
transport chain (mtETC) genes, which might disturb mitochondrial energy metabolic pathways.

Cytological analysis of anther development between the maintainer and C5-type CMS cabbage lines

Our mitochondrial sequence data indicated that the C5-type CMS cabbage line possesses CMS genes from the nap-type CMS system of *B. napus*. However, the mechanism by which male sterility genes induce pollen abortion in cabbage is still uncertain. Thus, comparative cytological analysis of anther development between the maintainer and C5-type CMS cabbage lines was performed to address which developmental stage was affected in the stamens of the C5-type CMS line and to further determine the cause of C5-type CMS in cabbage. In the maintainer line, primordial anther tissue with four microsporangiums was found to differentiate into sporogenous cells (Figure 10A), which differentiated into irregularly shaped microspore mother cells. The tapetal cells, middle layer, endothecium and epidermis differentiated from the surrounding connective tissue (Figure 10B). Tetrads of microspores were generated after meiotic divisions, the tapetal cells began to degrade (marked by the red arrow) (Figure 10C). Free haploid microspores dissociated from the tetrads during the uninucleate microspore stage, and the most of tapetal cells degrade (Figure 10D). The microspores differentiated into mature pollen grains, and the tapetal cells degenerated completely (Figure 10E). Finally, the mature pollen grains were released from the dehiscent anther (Figure 10F).

In the maintainer cabbage line, four locules of anthers developed symmetrically (Figures 10A–F). In contrast, a loss of synchronous locule development was observed in the C5-type CMS anthers during all anther development stages (marked by the black arrows) (Figures 10G–L). There were no morphological differences in the developed locules between the maintainer and C5-type CMS cabbage lines at the sporogenesis cell stage and microspore mother cell stage (Figures 10A, B, 10G, 11). However, the tapetum prematurely separated from the locule wall at the tetrad stage in the C5-type CMS line (marked by the blue arrows) (Figure 10I), rather than separating from the

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**FIGURE 8**

Y2H assay of the interaction of ORF222a with ATP17-2. Serial dilutions of cotransformed yeast were spotted on synthetic dropout SD/-Trp-Leu medium to select for cotransformants and SD/-Trp-Leu-His-Ade medium to select for positive interactions. p53 was used as a positive control.
connective tissue during the uninucleate microspore stage, as in the maintainer cabbage line (Figure 10D). In the uninucleate microspore stage of the C5-type CMS line, haploid microspores formed but crowded together (Figure 10J). The tapetal cells were still quite intact at the uninucleate microspore stage and mature pollen stage in the C5-type CMS line (marked by the blue arrows) (Figures 10J, K). Premature separation of the tapetum from the connective tissue and delayed cell collapse of tapetal cells hindered the release of haploid microspores, which resulted in the structural collapse of microspores. The residues of microspores that failed to develop were visible, and the epidermis, fibrous layer, and intermediate layer of pollen sacs were tightly connected, generally without dehiscence (Figures 10K, L). The tapetal cellular debris was still observed at the anthesis stage (marked by the blue arrow) (Figure 10L). Taken together, these findings indicate that premature separation of the tapetum from the locule wall and abnormal degradation of tapetal cells lead to overlap of microspores, which might be the cause of CMS in C5-type CMS cabbage lines.

Discussion

Mitochondrial origin of nap-type CMS in C5-type CMS B. oleracea

Most CMS systems of B. oleracea (cabbage) were transferred from other cruciferous crops. Because of incomplete pollen abortion and abnormal flower organ growth, the pol and Nig CMS systems cannot be used for cabbage breeding. The Ogu-CMS system is the main type of male sterility system in cabbage breeding (Ji et al., 2020). However, a high degree of cytoplasmic genetic uniformity may lead to genetically vulnerable plants (Yamagishi and Terachi, 2017), which prompted us to generate and apply a new type of CMS in cabbage. In this study, we generated the C5-type CMS cabbage line via intergeneric hybridization with nap-type CMS B. napus and then consecutive backcrossing. Sequencing revealed that the mitochondrial genome of the C5-type CMS cabbage line is quite homologous to that of the nap-type CMS B. napus strains 51218 and Westar but significantly different from that
of common cabbage (Figures 3B, 4C, 5). Both the chloroplast and mitochondrial genomes of progeny by sexual propagation are usually identical to those of maternal parents without recombination (Park et al., 2021). Here, the original cytoplasm of C5-type CMS cabbage was inherited from a certain nap-like CMS B. napus haplotype. However, the mitochondrial genome sequence of this B. napus haplotype has not yet been released.

C5-type CMS is associated with energy deficiency

Many ORFs in CMS lines have been found to disrupt energy metabolism through cotranscription with the F$_1$F$_0$-ATP synthase subunit. Here, we found that orf188a, orf322a, and orf322a were cotranscribed with rps7, cox1, and atp9, respectively (Figure 7). Most CMS-related genes have been found to be chimeric ORFs consisting of fragments of coding sequences of known mitochondrial genes from mtETC pathways, including cox1, atp6, atp8, atp9, and some other mitochondrial genes (Chen and Liu, 2014). For instance, orf463a for radish DCGMS-type CMS possesses a 128 bp fragment corresponding to part of the cox1 sequence (Wang et al., 2020). orf346 for Nsa-type CMS also shows partial sequence identity with the cox1 gene (Sang et al., 2021). In our previous study, we found that the Ogura-type CMS B. oleracea gene orf138, which originated from radish, encodes an atp8-like gene. The CMS-related gene orf154a is partially homologous to the ATP synthase subunit 1 (atpA) gene (Zhong et al., 2021). In nap-type CMS Brassica, chimeric orf222 encodes a protein with a segment of ATP8 at the N-terminus plus a sequence of unknown origin (Geddy et al., 2005). orf288 for Hau-type CMS B. juncea contains a 94 bp sequence with partial sequence homology to nad5 and partial sequence identity to atp9 (Heng et al., 2014). In our results, we found that the C5-type CMS-related gene orf188a encodes the atp6-like gene and that orf222a encodes 58 amino acids identical to ATP8 at the N-terminus (Supplementary Figure S1). The CMS-related genes encode partial fragments consistent with known functional mitochondrial genes. It has been speculated that these chimeric genes and the cotranscription events may disrupt mitochondrial functions by interfering with the expression of native mitochondrial genes, affecting the assembly or activity of different mitochondrial complexes (Shaya et al., 2012). In addition, it has been reported that chimeric genes confer the CMS phenotype by directly interacting with genes involved in the mtETC or ATP synthase complex. In maize, the chimeric gene atp6c confers CMS by interacting with ATP8 and ATP9, which impairs the assembly of the mitochondrial ATP synthase complex. Ultimately, this results in reductions in the quantity and activity of assembled ATP synthase required for anther development (Yang et al., 2022). In our study, we found that ORF222a impaired the step of F$_1$F$_0$-ATP synthase assembly by interacting strongly with an ATP17 homolog (Bo7g114140) in B.
oleracea (Figure 8 and Supplementary Figure S2). We also found that the ATP and ATPase activity levels in the anther samples of the C5-type CMS cabbage line were remarkably lower than those in the maintainer line (Figure 9). In addition, cytological sections showed that four locules develop synchronously through all stages of maintainer anther development (Figures 10A–F). In contrast, C5-type CMS anthers can display a loss of synchronous locule development (Figures 10G–I), which could be triggered by energy deficiency. Taken together, all of these findings indicate that C5-type CMS is associated with energy deficiency.

The abnormal development of tapetal cells probably causes male sterility in the C5-type CMS cabbage line

Programmed cell death (PCD) of the tapetum is one of the most critical steps for fertility. It is generally assumed that abnormal tapetal cells fail to produce enough nutrients and enzymes for microspore development and release from the tetrad, which often triggers the abortion of microspores (Liu et al., 2015). Additionally, defective tapetum PCD has previously been reported to cause anther indehiscence in an autophagy-deficient mutant of rice (Kurusu et al., 2014). In our results, the premature separation of the tapetum from the connective tissue and delayed cell collapse of tapetal cells hindered the release of haploid microspores, which directly resulted in both the structural collapse of microspores and anther indehiscence in the C5-type CMS line (Figure 10). This result is consistent with the development of nap-type CMS anthers in the Brassica napus (Geddy et al., 2005). However, it is quite different from the case in the Ogura-type CMS cabbage line, in which abnormal proliferation of tapetal cells hinders and destroys the development of haploid microspores through spatial constriction (Zhong et al., 2021).

Conclusion

We generated a new type of CMS cabbage line, named the C5-type CMS line, which showed abortion of microspores and anther indehiscence. orf188a, orf222a, orf261a, orf286a, and orf322a were specifically identified as candidate CMS genes via sequencing of the mitochondrial genome of the C5-type CMS cabbage line. We found that orf114a, orf188a, orf261a and orf322a were cotranscribed with atp8, rps7, cox1, and atp9, respectively (Figure 7). The most interesting finding was that ORF222a interacts with an ATP17 homolog (Bo7g114140) in R. oleracea, which might impair the step of F1Fo-ATP synthase assembly. We also found that the ATP and ATPase activity levels in the anther samples of the C5-type CMS cabbage line were remarkably lower than those in the maintainer line. These candidate CMS genes may perturb ATP synthesis during the anther development, which triggers a loss of synchronous locule development, premature separation of the tapetum from the connective tissue, and delayed cell collapse of tapetal cell in anthers, finally leading to CMS.

Data availability statement

The data presented in the study are deposited in the NCBI repository, accession number ON960289. The data will be released on Sep 17, 2022. We also provide confirmation of deposition.

Author contributions

XZ carried out the sequence data analysis, and drafted the manuscript. XZ, XY, JC, RH and YG designed and coordinated all experiments. JK supervised the work and edited the manuscript. All authors read and approved the final manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

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