Sequence Analysis and Expression of orf224 Gene Associated with Two Types of Cytoplasmic Male Sterility in Brassica napus L.

Jianmin Liu^[a,b], Maoteng Li^[a,b], Hao Wang^[c], Longjiang Yu^[a,b], and Dianrong Li^[c,*]

^[a] Institute of Resource Biology and Biotechnology, College of Life Science and Technology, Huazhong University of Science and Technology, Wuhan, 430074, China.
E-mail: limaoteng426@163.com
^[b] Key Laboratory of Molecular Biophysics, Ministry of Education, Wuhan, 430074, China
^[c] Hybrid Rapeseed Research Center of Shaanxi Province, Dali, 715105, China.
E-mail: lidr@peoplemail.com.cn

* Authors for correspondence and reprint requests

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Polima and Shaan 2A are the two most widely used forms of cytoplasmic male sterility (CMS) in the utilization of heterosis of rapeseed (Brassica napus) in China. A previous study indicated that the mitochondrial gene, orf224, was the only gene with a differential expression pattern among the normal, sterile and fertility-restored lines in rapeseed. DNA sequences of orf224, including coding sequences from Shaan 2A and Polima CMS, were then amplified and analyzed. DNA sequence alignment indicated both the coding sequences were 675 bp in length and had 99.9 and 99% homology in nucleotides and amino acids, respectively, and shared certain similarity to homologues from other Brassica spp. and Arabidopsis thaliana. The probable promoter regions of orf224 were conserved between B. napus and A. thaliana, but the upstream regions of probable promoter regions were completely divergent from each other. Additionally, analysis of the primary and secondary structure of the proteins encoded by orf224 from the two lines predicted that the proteins contain α-helix, extended strand, and random coil. After cloning a \textit{in vitro} experiment showed that these two proteins could be expressed in \textit{Escherichia coli} BL21.

\textbf{Key words:} Shaan 2A CMS, Polima CMS, orf224 Gene

Introduction

Cytoplasmic male sterility (CMS) is a maternally inherited character, which results from abnormal microsporogenesis or microgametogenesis leading to no pollen production or dysfunctional pollen. The hybrid seed yield of rapeseed (Brassica napus) was reported to be as much as 60% higher than in parental lines therefore it was frequently used in commercial F1 hybrid production (Horlow et al., 1992; Lin et al., 2006; Pascal and Francoise, 2004; Singh and Brown, 1991). Four types of CMS of rapeseed exist in nature, \textit{i.e.} nap CMS, Polima CMS, Shaan 2A CMS, and Ogu CMS. Nap CMS is also known as the Shiga-Thompson system, and was first reported by Shiga and Baba (1973) and Thompson (1972). However, the sensitivity of this CMS to temperature is a critical restriction for its application. Ogu CMS, first observed by Ogura in a Japanese radish groups, also has serious defects in hybrid production due to its deficiency in chlorophyll and the absence of a restoring gene (Gourret et al., 1992). Polima CMS, which might have originated from a Polish cultivar through genetic mutation, was discovered by Fu in 1972 (Fu, 1990). Shaan 2A CMS was discovered by Li in 1976 (Li, 1980) in the progeny of hybridization of S74 × (Fengshou 4 + 7207), and this CMS was first applied utilizing of rapeseed heterosis, and resulted in the world’s first hybrid rapeseed ‘Qinyou 2’.

At present, Polima and Shaan 2A CMS are the only two CMS widely applied in heterosis utilization in China. Although Shaan 2A CMS and Polima CMS have different origin, they are sometimes assumed to be the same type of CMS, because they share the same restorer lines and maintainer lines (Yang and Fu, 1991), and have the same cytoplasm as revealed by restriction fragment length polymorphism (RFLP) and mtDNA fragment polymorphism analyses (Yang et al., 1998). Previous studies of the molecular basis of Polima CMS have demonstrated that the orf224/atp6 coding region is the only mitochondrial gene region expressed differently in normal, sterile and
fertility-restored lines (Handa and Nakajima, 1992; Singh et al., 1996; Yuan et al., 2003).

To analyze the cytoplasmic difference in Shaan 2A and Polima CMS, the orf224 gene was selected for DNA sequence comparison. Cloning, expression in *Escherichia coli* BL21, and sequence analysis of the orf224 gene in Shaan 2A and Polima CMS were performed. The results indicated only one base difference between them. The difference upstream of these two genes and the phylogenetic relationship between *B. napus* and *A. thaliana* were analyzed. Secondary structure analysis revealed that the orf224 protein contained an α-helix, extended strand, and random coil. The results provide theoretical support for producing other sterile crops.

**Materials and Methods**

**Materials**

Shaan 2A CMS and Polima CMS lines were provided by Dr. Dianrong Li, Chinese National Rapeseed Genetic Improvement Center (Shaanxi, Dali, China).

**Bacterial strain and vectors**

*Escherichia coli* DH5α was used as the host for plasmid amplification, and *E. coli* BL21 (DE3) (Novagen, Darmstadt, Germany) was used for gene expression. pMD18-T simple vector and pET-32a (+) plasmid were purchased from Takara Biotechnology Co. Ltd. (Dalian, China) and Novagen, respectively.

**DNA isolation and primer design**

Total DNA was isolated from leaf tissue according to Li et al. (2007). The primers for amplifying orf224 were designed as follows, according to Lin et al. (2006):

orf224L: 5′-CGC|CATGGATGCCTCAACTGGATAAATTCAC-3′ (NcoI),
orf224R: 5′-CGGAGCT|CTCAGCGAAAGAGATCAAGGA-3′ (SacI).

**PCR amplification**

PCR amplification was carried out in 20 µL of reaction system including 10 ng of total DNA, 20 mM Tris [2-amino-2-(hydroxymethyl)-1,3-propanediol]-HCl (pH 8.4), 20 mM KCl, 1.5 mM MgCl₂, 0.4 mM of each dNTP, 10 pm of each primer, and 1.5 U of Pfu DNA polymerase (Tiangen Biotech Co. Ltd., Beijing, China). PCR was performed in an i-Cycler thermal cycler (BIORAD, Hercules, CA, USA). The PCR procedure for orf224 amplification was: 95 °C for 5 min, followed by 30 cycles at 94 °C for 60 s, 55 °C for 60 s, and 72 °C for 60 s, and then extension at 72 °C for 10 min.

**DNA fragments purification and construction of pMD18-T-orf224 vector**

The PCR product of the expected size was separated on 1.2% agarose gel and purified with 3S spin agarose gel DNA purification kit (Shanghai Biocolor Bioscience & Technology Co. Ltd., Shanghai, China) following the manufacturer’s instructions. Subsequently, the purified PCR product was cloned into pMD18-T simple vector. The recombined vector (pMD18-T-orf224) was transformed into *E. coli* DH5α. Positive recombinants were verified by PCR, and sequenced at Beijing Sunbiotech Co. Ltd.

**Bioinformatic analysis**

The nucleotide sequence and inferred amino acid sequence of orf224 of Shaan 2A and Polima CMS were aligned with CLUSTAL X software and edited with the Jalview Multiple Alignment Editor V1.8. Sites containing gaps were excluded. Cluster trees were constructed with MEGA version 3.1 (Kumar et al., 2004), and the neighbour-joining method was applied to p-distance. The secondary structure prediction was performed online (http://bioinf.cs.ucl.ac.uk/psipred/psiform.html/).

**Construction of pET32a (+)-orf224 plasmid**

pMD18-T-orf224 and pET32a (+) were digested by NcoI and SacI restriction enzymes according to the manufacturer’s protocol (Fermentas, Vilnius, Lithuania). The prepared NcoI-orf224-SacI insert was ligated into the treated pET32a (+) vector using T₄ DNA ligase at 16 °C overnight. The *E. coli* DH5α and *E. coli* BL21 (DE3) competent cells were prepared by the calcium chloride method and were used for transformation. The transformed bacteria were selected on media containing ampicillin. Meanwhile, the positive colonies were further analyzed by PCR, restriction enzyme digestion, and sequencing to ensure plasmid integrity.
Induction of orf224 expression with IPTG

Three 5-mL cultures of the recombinant E. coli BL21 were prepared in LB (Luria-Bertani) medium containing 100 µg/mL ampicillin. The cultures were incubated at 37 °C with shaking to an OD_{600} of 0.6–0.8. Then isopropyl-β-D-thiogalactopyranoside (IPTG) was added to the first two 5-mL cultures for a final concentration of 1 mM. The third culture was saved as control. All cultures were incubated while shaking at 37 °C. Samplings were done by removing 1 mL of cultures after 3 and 4 h, respectively. The cell pellets obtained after centrifugation at 9168 × g for 2 min were stored at −20 °C.

Protein analysis by polyacrylamide gel electrophoresis (PAGE)

The cell pellets were resuspended in 100 µL of sterile distilled water. Then 100 µL of 2 × SDS loading buffer were added and mixed thoroughly. Samples were heated for 8 min at 100 °C and cooled immediately at 0 °C to denature the proteins and then analyzed by SDS-PAGE. The resolving and stacking gels used were 15% (w/v) and 5% (w/v), respectively (Laemmli, 1970). The gel was allowed to run for 1 h at 60 V and 2 h at 100 V, followed by staining with Coomassie Brilliant Blue R250. The low molecular weight standard was 14.4–94.0 kDa.

Results

Amplification of the orf224 gene from Shaan 2A and Polima CMS

The A_{260}/A_{280} and A_{260}/A_{230} of total DNA were assessed with a Beckman DU7500 instrument (Brea, CA, USA), and the ratios of both A_{260}/A_{280} and A_{260}/A_{230} were in the range 1.8–1.9, which indicated that the total DNA had high quality and could be used for PCR amplification.

The orf224 gene of Shaan 2A and Polima CMS was amplified by the designed primers. The agarose gel analysis showed that both Shaan 2A and Polima CMS had a specific band of 500–750 bp length (Fig. 1a). The four purified PCR products were cloned into pMD18-T simple vector and then verified by NcoI and SacI double digestion (Fig. 1b). The recombined vector was transformed into E. coli DH5α, and the positive recombinant clones were sequenced.

Sequence analysis and secondary structure of orf224 in Shaan 2A and Polima CMS

According to gene sequencing results, the length of orf224 in both CMS lines was 675 bp (GenBank accession numbers for orf224 of Shaan 2A and Polima CMS were EU254234 and EU254235, respectively); it encoded a polypeptide including 224 amino acids. The homology of the two genes in nucleotides and amino acids was 99.9 and 99%,
Fig. 2. Sequence alignment of the orf224 gene in Shaan 2A and Polima CMS lines and a homologous sequence (including a upstream 300-bp fragment) in the mitochondrial genome of *B. napus* (GenBank number: AP006444) and *A. thaliana* (GenBank number: Y08501). The solid line frame indicates the orf224 gene regions.
respectively (Figs. 2 and 3), and they had 83% homology with a related gene in the mitochondrial DNA of *B. napus* (AP006444) (Handa, 2003) and 57% homology with an associated gene in the mitochondrial genome of *A. thaliana* (Y08501) (Unseld *et al*., 1997). Meanwhile, sequence alignment showed that orf224 gene regions were conserved in *B. napus*, and the probable promoter regions were conserved between *B. napus* and *A. thaliana* (Fig. 4). However, the upstream regions of probable promoter regions were completely divergent from each other. The results were coincident with the experimental analysis of Handa (2003).

The orf224 protein included 224 amino acids, with a molecular mass of 26 kDa. A theoretical pI of 9.08 for Shaan 2A and a pI of 8.9 for Polima CMS were deduced in the ExPaSy web server (http://expasy.org/tools/protparam.html). There were 24 negatively charged residues (Asp + Glu) and 29 positively charged residues (Arg + Lys) of the orf224 protein of Shaan 2A CMS, but the respective numbers were 25 and 29 residues for Polima CMS. The instability index (II) was computed to be 53.17, which showed that the protein was unstable. Grand average of hydropathicity was –0.09. The secondary structure had seven
\[ \alpha \]-helices, four extended strands, and eight random coils. However, no \( \beta \)-turn was observed (Fig. 5).

**Construction of recombinant plasmid and protein expression analysis**

The expression vector pET32a (+) was used to construct pET32a (+)-orf224. *NcoI* and *SacI* cloning sites were chosen for insertion. The constructed plasmid was amplified in *E. coli* DH5\( \alpha \) and expressed in *E. coli* BL21 (DE3). Ligation of the orf224 gene insert into pET32a (+) and subsequent transformation of *E. coli* DH5\( \alpha \) yielded ampicillin-resistant colonies. The selected colonies were purified to obtain a plasmid, and then double digests on the plasmid with *NcoI* and

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Fig. 5. The secondary structure of the orf224 protein.
SacI determined the presence of the orf224 gene (Fig. 1c).

Plasmid pET32a (+)-orf224 was transformed into E. coli BL21 (DE3) competent cells, and one of the positive colonies was cultured in LB medium, containing 100 µg/mL ampicillin, and grown at 37 °C. When the OD₆₅₀ value reached 0.6–0.8, IPTG was added to a final concentration of 1 mM, and incubation was continued for 3 and 4 h. The cells were centrifuged at 9168 × g for 2 min. The expression protein was analyzed by SDS-PAGE. Compared with the uninduced cells, the IPTG-induced cells had an additional protein band of 26 kDa on SDS-PAGE gel (Fig. 6), which corresponded to the calculated molecular weight of the protein. The amount of additional protein induced after 3 and 4 h was equal.

Discussion

CMS occurs widely in higher plants and is widely used in plant heterosis. Much research has revealed that mitochondria have the closest relationship with CMS (L'Homme and Brown, 1993; L'Homme et al., 1997; Schnable and Wise, 1998; Song and Hedgcoth, 1994). Erickson et al. (1986) found a remarkable difference between mtDNA of Polima CMS and of cam CMS. Further research revealed that some regions of the mitochondrial genome were absolutely necessary to CMS (Schnable and Wise, 1998). It was shown that the orf224/atp6 region was correlated with the Polima CMS (Wang et al., 1995) and the orf222 was related to nap CMS (L'Homme et al., 1997); the homology of nucleotides and proteins of these two regions was 85 and 79%, respectively (L'Homme et al., 1997). orf138 was only found in Ogu cytoplasm (Bellaoui et al., 1997). Wan et al. (2007) compared hau CMS with pol CMS, nap CMS, ogu CMS, and tour CMS in the fields of genetics, cytology, and molecular biology. They found that the restorer and maintainer relationship of hau CMS differed from other CMS lines, and they could distinguish between other five CMS lines using RFLP. Shaan 2A CMS and Polima CMS have the same restorer lines and maintainer lines. Many researchers thought that these two CMS belong to the same line, according to research on the general shape, cytology and restriction maps (Liu et al., 1998). Wang et al. (2002) analyzed the sequence of the orf224 gene of Shaan 2A CMS and Polima CMS, and found only one nucleotide difference. However, Lin et al. (2006) found differences in five bases between Shaan 2A CMS and Polima CMS.

The present research on the sequence of the orf224 gene found only one base difference, which led to one amino acid change. The results confirmed that Shaan 2A and Polima CMS were not of the same CMS. The secondary structure of the orf224 protein was predicted, and it was observed that the orf224 protein was expressed immediately after induction with IPTG in E. coli BL21 (ED3). Although this is a preliminary conclusion, it is very important for studying the relationship between the orf224 gene and CMS.

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