Development of SSR Markers Associated with Polima CMS Fertility Restorer in Brassica napus L. and Its Application

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Abstract

The Polima system of cytoplasmic male sterility (Pol CMS) was widely used for hybrid seed production, but with high probability of occurrence of maternal false hybrid due to the fertility of Polima CMS (Pol CMS) line greatly affected by the ambient temperature, therefore, the detection of maternal false hybrids is the key to purity identification. In present study, the SSR markers were designed based on the flanking sequences of Pol CMS restorer gene Rfp, and application of SSR markers in hybrid seed purity test of rapeseed. The main results were as follows: the sequences of Rfp were subjected to basic local alignment search tool queries against the Brassica rapa genome to determine chromosome positions, and 32 SSR markers were developed based on the upstream and downstream sequences of Rfp. Among the 32 SSR markers, one codominant marker was polymorphic between Pol CMS sterile and restorer line, the PCR product length was 162 bp in Pol CMS sterile lines, and 184 bp in restorer lines. Based on the PCR sequence, two insert fragments (15 bp and 7 bp) were identified in Pol CMS sterile lines, and 32 SSR markers were developed based on the upstream and downstream sequences of Rfp. The SSR marker co-segregated with plant fertility of individuals in the “fengyou737” F2 population. The purity of “fengyou737”, “fengyou320” and “fengyou958” were identified by means of PCR-SSR markers and field-planting, and the results of two ways were highly consistent, indicated that the codominant SSR marker tightly linked to Rfp, and the marker can detect paternal and maternal false hybrids. The present findings will provide technical assistance for hybrid seed purity test and genetic diversity analysis in rapeseed.

Keywords Brassica napus L.; Pol CMS; Restorer gene; SSR

Polima cytoplasmic male sterility (Pol CMS) is the first cytoplasmic male sterility type of Brassica napus with production value. It is also the sterile source of the most widely used hybrid pollination control system in rapeseed in China (Fu, 1995). The sterility of the male sterile lines in this system is controlled by the interaction between the mitochondrial male sterile genes and the nuclear genome, and the fertility restorer gene (Rfp) is located in the nuclear genome (Yang et al., 1996). The fertility of this type of sterile line is easily affected by ambient temperature, causing the occurrence of self-pollination by its own pollen in the process of large area hybrid seed production, which leads to the production of the maternal pseudo hybrid. Because the female pseudo hybrid is sterile, it not only affects the consistency of varieties, but also seriously affects the production advantage of varieties. Therefore, strict identification of seed purity and quality must be carried out before the application and production of rapeseed hybrids (Jean et al., 1997; Natasa et al., 2006; Natasa et al., 2010; Zhao et al., 2010). At present, the laboratory identification of rapeseed purity quality mainly includes three categories: isozyme electrophoresis, gliadin electrophoresis and molecular marker techniques. Among them, isozyme electrophoresis and gliadin electrophoresis have the advantages of simple technique, rapid identification and low cost, however, the polymorphism of isozyme and storage protein was relatively low, for some hybrid combinations, it is difficult to distinguish their parents from their hybrids, so the practical application of this technique is greatly limited. Molecular markers based on genomic DNA sequences, especially SSR markers developed in the early 1990s (Wang et al., 2010; Zhang et al., 2013). Compared with other molecular markers, this type of marker is relatively simple in operation and low in cost. Therefore, it has become the preferred type of molecular marker in the identification of purity and authenticity of crop varieties (He et al., 2009; Song et al., 2013; Wang et al., 2014; Zhu et al., 2015; Wang et al., 2016). However, with the narrowing of genetic diversity
of breeding resources, the workload of SSR marker screening has been increasing. At the same time, breeders often start parental reproduction and cross production when the phenotypic traits are basically the same in the selection of hybrid parents. However, there is still separation of genetic information at the molecular level, which makes it difficult to ensure the accuracy of the purity identification of hybrid lines (Wang et al., 2009; Zhan et al., 2015; Li et al., 2016).

The results showed that the mono-dominant gene Rfp was responsible for the fertility recovery of Pol CMS of B. napus (Yang et al., 1996). In recent years, researchers have done a lot of genetic mapping of Rfp, and developed a series of molecular markers linked to the fertility restorer genes (Liu et al., 2007; Yang et al., 2008). Rfp gene has been located in the N9 or N18 linkage group respectively. Until 2014, Rfp was mapped to the N9 linkage group of B. napus, and was restricted to the 29.2k region of A9 in B. rapa by molecular marker analysis. This region contains 7 ORFs, of which ORF2 is most likely involved in fertility recovery (Liu et al., 2012; Liu, 2012). The researcher then transferred the ORF2 into sterile lines of B. napus, the results showed that ORF2 had the function of fertility recovery (Chen et al., 2014). Further sequence analysis showed that the coding region of ORF2 was 1 953 bp long (Liu et al., 2016). Meanwhile, the genome sequencing information of B. rapa. and B. napus was published in 2011 and 2014, respectively, which also provided a useful reference platform for the development of molecular markers.

In light of the actual demand for purity identification of Pol CMS hybrids, the sequence information of Rfp candidate gene region was used in this study to perform BLAST analysis on target sequences in B. napus genome and B. rapa genome to determine the position and sequence of the PPR gene related to fertility restoration. And then, SSR markers were developed in the nearby region and used in the purity identification study of Pol CMS hybrids. It was expected to provide a more accurate and efficient detection method for the purity identification of rapeseed hybrids.

1 Results and Analysis

1.1 Development of fertility restorer gene-linked SSR Markers

Based on the 1 953 bp sequence of ORF2 published by Liu et al. (2016), a highly homologous region (99% homologous) was obtained on the A9 chromosome of B. rapa. A total of 32 SSR loci were obtained in the upper and lower 15K region of this region, and 32 pairs of SSR primers were designed and synthesized online. The 32 pairs of primers were used for PCR amplification of genomic DNA of 10 tested parental materials, and one pair of SSR primer RfpSSR19 (F: AGTTAGCTCCTCTGTGGTTTGC; R: CTACAGCTCGATTAGGGATCGT), and its amplification site was located at NC_024 803.1:34 437 588 to 34 437 748 of B. napus genome. It showed significant polymorphism in tested 4 male sterile lines and 6 restorer lines (Figure 1), which was a co-dominant marker of sterile lines and restorer lines. Sequencing results of the specific fragments showed that the size of the amplified fragments was 162 bp in the sterile line and 184 bp in the restorer line. Compared with the sterile lines, there were one 15 bp and one 7 bp inserts in the amplified fragments of the restorer lines (Figure 2). The amplified product of RfpSSR19 was compared in B. napus, and the marker was located in the region from -3 962 bp to -3 801 bp upstream of the initiation site of Rfp translation, indicating that the obtained marker RfpSSR19 was closely linked to the Rfp gene.

1.2 Field location test of single plant of F2 segregating population

At the blooming stage, the F2 segregating population of the selfed progeny of the pol CMS hybrid Fengyou 737 was tested for fertility, and the SSR marker RfpSSR19 was used to analyze the genotypes of 192 randomly selected individual plants. The results showed that the single plant fertility of 184 bp fragment was sterile, and the PCR amplification band pattern was co-segregated with the single plant fertility (Figure 3).
1.3 Comparison of purity identification results of tested hybrid samples

The selected SSR primers were used for PCR amplification of the genomic DNA of the seedlings of the tested hybrid varieties Fengyou 737, Fengyou 320 and Fengyou 958, and clear and stable amplification bands were obtained (PCR amplification results of Fengyou 737 hybrid samples) (Figure 4), and the reliable SSR purity identification results were obtained. As can be seen from the purity identification results of field planting (Table 1), the obtained results of SSR marker identification and field planting identification are highly consistent, and the
The difference of identification results of 9 tested samples is less than 4%, indicating that marker RpfSSR19 is reliable for purity identification results of test materials used in this study.

Figure 4 Amplification patterns of SSR-PCR in Fengyou 737 hybrids purity identification

Table 1 Purity of field planting identification and SSR identification of hybrid samples

| Cultivars  | No.  | SSR purity (%) | Field purity (%) |
|-----------|------|----------------|------------------|
| Fengyou 737 | 737-1 | 89.5           | 91.6             |
|           | 737-2 | 92.3           | 91.5             |
|           | 737-3 | 87.9           | 90.2             |
|           | 320-1 | 89.3           | 92.2             |
| Fengyou 320 | 320-2 | 88.8           | 89.1             |
|           | 320-3 | 87.9           | 90.8             |
|           | 958-1 | 92.1           | 90.5             |
| Fengyou 958 | 958-2 | 90.3           | 93.6             |
|           | 958-3 | 88.5           | 91.7             |

2 Discussion

The temperature sensitivity of male sterile lines is a problem in Pol CMS hybrid seed production system. For a long time, the monitoring of seed purity and quality, especially the accurate identification of female pseudo hybrids in hybrid lines, has been an indispensable link in rapeseed hybrid production (Zeng et al., 2009). In this study, a specific SSR marker closely linked to the Rfp gene was developed based on the actual requirements for purity identification of Pol CMS hybrids in rapeseed. The production and application tests showed that the marker could be used for purity and quality identification of different Pol CMS hybrids, and the identification results were highly consistent with the field fertility identification. The present study solves the problem that SSR markers for the purity of rapeseed hybrids need to be screened from genome-wide databases for the paternal and maternal lines of hybrid crossings (Mei et al., 2006; Tang et al., 2007). Therefore, a large number of preliminary works on the synthesis and screening of primer sequences is reduced, lowering the cost and early investment of identification, which is conducive to the popularization and application of the technology. At the same time, it overcomes the technical difficulty that common SSR markers can only be used to detect the genotypes of paternal and maternal lines of specific rapeseed hybrid crossings, which are greatly affected by biological and mechanical hybrid pseudohybrids, resulting in a certain deviation between identification results and field fertility identification.

At present, molecular markers based on PCR amplification, especially SSR markers, are abundant in genetic information due to their large number, high polymorphism, stable results and simple operation (Liu and Li, 2007; Chen et al., 2009), has become the preferred method for indoor purity identification of rapeseed hybrids. In this study, based on the advantages of common SSR markers, a specific SSR marker located in the -3 962 bp to -3 801 bp region upstream of the translation initiation site of Rfp gene was developed. Because it is difficult to recombine with the restorer gene, this marker can be used to identify the fertility of individual plants in the progeny of Pol CMS. Therefore, the results of purity identification by using it are more consistent with the individual fertility of field planting identification, and it has more practical significance in the purity identification of hybrids. At the same time, this marker is a co-dominant marker, which can effectively determine the pseudohybrid types of male and female parents in the hybrid samples, and can be used to determine the hybrid seeds of Pol CMS parents. The
results obtained in this study can provide a new technique and method for the department of seed management and law enforcement.

The molecular marker identification purity of the 9 samples tested in this study was generally lower than that of field planting, but the differences were less than 4%. It is speculated that it may be due to the stronger growth competitive advantage of hybrids in the process of field planting identification, which is easier to survive as a plant than its parents, leading to a higher results of fertility identification at flowering stage. In addition, the polymorphism of the markers obtained in this study was only 22 bp difference between sterile and restorer lines, which required a long time to separate by agarose gel electrophoresis. Therefore, markers with more obvious differences of fragment size could be developed by further primer design to improve the efficiency of purity identification.

3 Materials and Methods
3.1 Experimental materials
The test materials: 12 parents of Pol CMS hybrid of Brassica napus, including 5 sterile lines and 7 Pol-CMS restorer lines (Table 2). Three samples of Fengyou 737, Fengyou 320 and Fengyou 958 were produced from Pol-CMS hybrid varieties, and Fengyou 737 was self-crossed into F2 population.

Table 2 Name and source of 12 rapeseed lines tested in this study

| No. | Material name           | Source                                                                 |
|-----|------------------------|------------------------------------------------------------------------|
| 1   | 20A                    | Pol-CMS sterile line transformed with germplasm from JiangXi           |
| 2   | 317A                   | Pol-CMS sterile line bred from reciprocal cross population between 16A and double-low line 317 |
| 3   | 167A                   | Pol-CMS sterile line bred from 17A                                    |
| 4   | 17A                    | Pol-CMS sterile line bred from Xiang5A with pol CMS and double-low line Zhongshuang 10 by hybridization and subsequent orientation selection |
| 5   | 169A                   | Pol-CMS sterile line bred from Xiang5A with pol CMS and double-low line SC4 by hybridization and subsequent orientation selection |
| 6   | 13R                    | Pol-CMS restorer line transformed with Xiangyou 13                    |
| 7   | XWR                    | Pol-CMS restorer line bred from Pol-CMS hybrid in Hubei               |
| 8   | Z2R                    | Pol-CMS restorer line bred from Pol-CMS hybrid Zhongyouza 2           |
| 9   | 320R                   | Pol-CMS restorer line bred from Fengyou 730 with pol CMS and double-low line 320 by emasculation hybridization and subsequent orientation selection |
| 10  | SC4                    | Pol-CMS restorer line transformed with germplasms from Sichuan        |
| 11  | SC3                    | Pol-CMS restorer line transformed with germplasms from Sichuan        |
| 12  | 95-5                   | Pol-CMS restorer line bred from Pol-CMS hybrids in Hubei              |

3.2 Sampling of test samples
At the end of September 2018, 12 Pol CMS hybrid parental materials were sown in the experimental field of Hunan Crop Research Institute (the last crop was rice). At the seedling stage (about 7-leaf stage), 5 young leaves of each material were picked out and placed in a 2 mL centrifuge tube and stored at -20℃ for primer screening.

Each sample of Fengyou 737, Fengyou 320 and Fengyou 958 was randomly selected about 1 000 seeds and divided into two parts, one for molecular marker purity identification and the other for field planting purity identification.

About 500 F2 population seeds of the tested variety Fengyou 737 were selected and sown in the experimental field of Hunan Crop Research Institute. Each sample identification plot was in 8 rows, with a line length of 2.0 m and a row spacing of 0.5 m. The seeds were evenly sown and the seedlings were not thinned during the whole growth period.

3.3 Development of SSR markers linked to Rfp
The 1953 bp sequence of ORF2 published by Liu and Zhang (2016) was performed BLAST analysis on the B. rapa genome and B. napus genome (http://brassicadb.org/brad/) to identify the physical locations of highly homologous Rfp genes related to fertility restoration. Based on the location and its area on both sides of the 5 K,
10 K, 15 K for SSR loci, via http://wsmartins.net/websat/ online design SSR primers. Then, the designed SSR primers were used for PCR amplification on genomic DNA of different restorer lines and male sterile lines, and the primers with differences were screened as linkage markers of Rfp gene.

3.4 Verification of cosegregation of fertile segregating populations
From the 7th to 9th leaf stage, 192 single plants of Fengyou 737 self-pollinating F2 population were sequentially selected for tagging, and genomic DNA of single plant was extracted. PCR amplification was performed using the primers screened above, and the amplified products were electrophoresis separated by 3.0% Agarose gel. Finally, observation, photo taking, and band reading were performed in the gel imaging system. The amplified band types of each individual plant were recorded. In the flowering stage, the fertility performance of 192 plants with tag number was investigated.

3.5 Identification of purity of hybrid samples by molecular markers
After the samples of the three varieties were germinated, the genomic DNA of young embryos was extracted according to Chen Ying’s CTAB method (Chen, 1993). The genomic DNA of young embryos of Fengyou 737, Fengyou 320 and Fengyou 958 seed production samples were amplified by PCR using primers with different PCR products between sterile and restorer lines. The amplified products were separated by electrophoresis with 3.0% agarose gel. Finally, the purity of samples was calculated by observing, taking photos and reading bands in the gel imaging system. SSR purity (%)=(number of tested plants-number of maternal band types-number of paternal band types)×100/number of tested plants.

3.6 Purity identification of hybrid samples in field planting
The samples to be tested of the three varieties were sown in the test field of Hunan Crop Research Institute. Each sample to be tested was identified in 8 rows with a line length of 2.0 m and a row spacing of 0.5 m. The seeds were evenly sown, and no seedlings were thinned during the whole growth period. Purity of field identification (%)=(number of tested plants-number of sterile plants-number of heteromorphic plants)×100/number of tested plants.

Authors’ contributions
WTH was the executor of the experimental design and experimental research of this study. WTH, LB and GYM completed the data analysis and drafted the manuscript. LB, GYM, ZYT and LXH participated in the experimental design and analysis of the experimental results. FLY, QL and DLC revised the manuscript and checked the data. LM conceived of the project, directed the design of the study, data analysis, draft and revision. All authors read and approved the final manuscript.

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