Genome-wide screening and analysis of imprinted genes in rapeseed (Brassica napus L.) endosperm

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Abstract

Species-specific genomic imprinting is an epigenetic phenomenon leading to parent-of-origin-specific differential expression of maternally and paternally inherited alleles. To date, no studies of imprinting have been reported in rapeseed, a tetraploid species. Here, we analysed global patterns of allelic gene expression in developing rapeseed endosperms from reciprocal crosses between inbred lines YN171 and 93275. A total of 183 imprinted genes, consisting of 167 maternal expressed genes (MEGs) and 16 paternal expressed genes (PEGs), were identified from 14,394 genes found to harbour diagnostic SNPs between the parental lines. Some imprinted genes were validated in different endosperm stages and other parental combinations by RT-PCR analysis. A clear clustering of imprinted genes throughout the rapeseed genome was identified, which was different from most other plants. Methylation analysis of 104 out of the 183 imprinted genes showed that 11 genes (7 MEGs and 4 PEGs) harboured differentially methylated regions (DMRs). Unexpectedly, only 1 MEG out of these 11 genes had a DMR that exhibited high CG methylation rate in paternal allele and had big difference between parent alleles. These results extend our understanding of gene imprinting in plants and provide potential avenues for further research in imprinted genes.

Key words: rapeseed, endosperm, RNA sequencing, imprinting, allele-specific, DNA methylation

1. Introduction

Genomic imprinting is an epigenetic regulatory phenomenon in which there is allele-biased expression of certain genes depending on the parent of origin.1 Imprinted gene expression plays a role in fetal growth regulation and postnatal behaviour in mammals, formation of viable seeds and the inhibition of interspecies hybridization in plants.2–5

In mammals, imprinting occurs in the placenta, embryo and adult tissues, while most imprinting in plants occurs in the endosperm.6–10 The seed of angiosperms comprises three major components, the seed coat, the embryo and the endosperm, which supports embryo-genesis.11 There is evidence for cross-regulation between seed components. Seed growth and development require communication and coordination of distinct genetic programs that govern the
Identification of endosperm imprinted gene in rapeseed

In this study, using the genome of Zhonghuang 11 as reference sequence, we identified rapeseed imprinted genes and analysed their cytosine methylation levels. First, we performed deep mRNA sequencing and identified SNPs between two rapeseed lines, YN171 and 93275. Using YN171 and 93275 as parents, mRNA-seq libraries of reciprocal hybrid endosperm were constructed. Finally, based on the differential expression of parental alleles in the reciprocal endosperms, 183 imprinted genes with parent-of-origin-specific expression were identified. Further genomic DNA methylation sequencing showed that the imprinted genes used for cytosine methylation analysis were not associated with DNA methylation. Our study not only provided an opportunity to understand the mechanisms driving gene imprinting and seed development in rapeseed but also shed some light on the evolution and biological significance of gene imprinting in plants in general.

2. Experimental procedures

2.1. Plant tissue collection

Rapeseed (Brassica napus L.) lines YN171, 93275, ZY036, 6F313 and 61616 introduced in our previous studies were grown in fields at Wuhan, China.48,49 Reciprocal inter-strain F1 hybrids, YN171 (♀) pollinated with 93275 (♂) (designated Y9) and 93275 (♀) pollinated with YN171 (♂) (designated Y9) were produced by manual pollination in 2016. Other reciprocal inter-strain F1 hybrids including 93275 and ZY036, 93275 and 9T364, 93275 and 61616 were used for identification of imprinted genes.

Endosperms for RNA-seq and DNA-seq were sampled from immature seeds 30 days after pollination (DAP), from the reciprocal F1 hybrids, both 9Y and Y9, and their parental lines, 93275 and YN171, in two biological replicates. Endosperms for RT-PCR were collected at 15, 20, 25, 30 and 35 DAP separately. Each sample was collected from at least 10 individual plants. Other tissues, including leaf, stem, root, bud, pericarp, embryo and flower, were also sampled from YN171 line in two biological replicates. The endosperm was collected according to the method introduced by Li et al.50 To be specific, the seeds were kept on ice and punched rapidly at the top with a sharp steel needle to make a small hole. The prepared glass micropipette was immediately inserted into the hole and gently aspirated liquid endosperm from the embryo sac. Then the lipid endosperm was quickly transferred into EP tube on ice. All database related to the identification of rapeseed imprinted genes in this MS were uploaded to https://www.ncbi.nlm.nih.gov (BioProject ID: PRJNA470566). A table including detailed information of database was also supplied (Supplementary Table S1).

2.2. RNA-seq data generation and processing

Total RNA was extracted using a commercial kit (TaKaRa Mini BEST Plant RNA Extraction Kit). RNA quantity was measured using a NanoDrop (ND-1000, Nano-Drop). RNA-seq was performed in BGI. Specifically, libraries were constructed following the TruSeq RNA Sample Preparation Guide. The Agilent 2100 Bioanalyzer and ABI Step One Plus Real-Time PCR System (11 amplification cycles) were used for identification of imprinted genes.

Endosperm DNA methylation levels were determined using bisulfite sequencing. In parental lines and reciprocal hybrids, the genome DNA of parental and reciprocal endosperm was digested separately with four endonucleases, BbsI, HpaII, MboII and SspI. The methylated and non-methylated products were sequenced using the Illumina HiSeq X Ten platform with paired-end read lengths at 100 bp. A total of 100 bp paired-end reads were obtained, providing sufficient sequencing depth for the DNA methylation analysis. Low-quality reads were removed from the raw sequences and low-quality nucleotides were clipped from the beginning and the end of sequences using the trimmomatic-0.36 with Sliding window.
The reads were mapped to the *B. napus* genome (http://www.genoscope.cns.fr/brassicanapus/) using TopHat version 2.1.1 (+10-read-edit-dist10). To obtain gene expression level in tissues, reads that were not uniquely mapped were excluded. To determine whether the SNPs identified strictly referred to the high similarity of paralogous genes in the tetraploid rape, we compared the diversity between species and paralogous genes by calculating the ratio between the variant sites (mismatch and small Indel) and covered length at both cDNA and DNA level. We note that 94% genes with larger variation between paralogous than between species, and more than 82% genes with significant variation (Fisher exact test, \( P < 0.05 \)) which has enough sensitivity and specificity in SNP detection. For each gene, the expression level was measured by Reads Per Kilobase exon Model per Million mapped reads (RPKM).

To obtain a more comprehensive list of SNPs between the parental orthologs, the genomes of the two parental samples were also resequenced. Reads from the genome samples were mapped to the genome of Zhongshuang 11 using BWA. Samtools (1.3.1) was used to exclude reads that were not uniquely mapped and pairs that were split across different chromosomes in all of the RNA and DNA analysis. In addition, duplicate reads were removed from the DNA analysis. Alignments from same parent were merged together for SNP identification.

**2.3. Identification of SNPs between parental orthologs**
A list of SNPs between YN171 (Y) and 93275 (9) was generated using the Samtools command Mpileup and the Bcftools program. First, a list of all positions containing potential variants and consistent sites with reference in exon regions for both YN171 and 93275 were generated. SNPs between YN171 and 93275 were filtered using the following criteria: (i) depth \( \geq 10 \) (at least 10 reads covering the SNP site), (ii) exclusion of heterozygous sites in each sample by retaining the sites with the alternative bases accounted for \( >95\% \) of the reads.

**2.4. Identification of imprinted genes**
The ratio of maternal to paternal alleles in hybrid endosperm is theoretically 2:1, so the observed ratio of maternal versus paternal allele counts for each gene in Y9 and 9Y was used to perform a \( \chi^2 \) test. Genes with a parental allele bias differing from 2:1 \( (q < 0.05) \) in reciprocal hybrids were classified as potentially imprinted genes. Two replicates were considered separately and imprinted genes were identified as overlap among replicates. To identify the imprinted genes with greater confidence, more stringent standards were applied. In maternally biased endosperm genes (paternal bias greater than 2:1 at \( q < 0.05 \)), criteria was set for at least 85% of the informative reads were maternal in both directions of the reciprocal cross. For paternally biased genes (paternal bias less than 2:1 at \( q < 0.05 \)), criteria was set at least 50% of informative reads in both directions of the reciprocal cross were paternal. Genes that met these criteria in the reciprocal hybrids were defined as allele-specific imprinted genes. All statistical analysis was performed using R (version 3.2.1). Corrections for multiple testing were performed with Storey’s \( q \)-values of 0.05.

**2.5. DNA preparation and locus-specific sequencing**
Genomic DNA was extracted from parental leaf tissues (YN171, 93275) using a Qiagen DNeasy Plant Mini Kit according to the manufacturer’s instructions. Primers used for SNP identification were designed manually (Supplementary Table S2). PCR was performed using TaqPlus polymerase (Qiagen). The PCR products were sequenced by Sanger sequencing.

**2.6. Gene expression analysis by RT-PCR**
Total RNA from endosperms of reciprocal F1 hybrids and their parental lines was used for identification of putative imprinted genes. Approximately 800 ng of total RNA was used for cDNA synthesis using a reverse transcription (RT) kit (TransGen Biotech, Beijing, China) following the manufacturer’s instructions. RT-PCR primers were designed manually (Supplementary Table S2). The PCR products were sequenced by Sanger sequencing.

**2.7. Clustering analysis**
For clustering analysis, 183 predicted imprinted genes were mapped to the rapeseed chromosomes (except for 4 genes, which could not be mapped to any chromosome). Arabidopsis imprinted genes were collected from the studies published in recent years. Sliding windows of 1 MB with step size 0.1 MB were used to compare the number of mapped reference genes and imprinted genes using a hypergeometric distribution. Consecutive significant windows \( (P < 0.05) \) were combined to form a cluster. The statistical analysis was performed using R (version 3.2.1).

**2.8. Gene MapMan analysis**
MapMan analysis of the imprinted genes was performed with the Classification SuperViewer on the website (http://bar.utoronto.ca/welcome.htm) based on sequence homology with Arabidopsis genes. Categories with \( P < 0.001 \) were classified as enriched categories. PEGs and MEGs at each significant level were analysed separately.

**2.9. Genomic DNA extraction and bisulphite sequencing**
Genomic DNA was extracted from endosperm at 30 DAP by CTAB method and bisulphite sequencing by BGL specific for, at least 1 \( \mu \)g of RNase-treated DNA was used for library preparation. The DNA samples were prepared according to the following treatments: The DNA, fragmented to 100–300 by sonication (Covaris), was purified and added a single ‘A’ nucleotide to the 3’ ends of the blunt fragments. The purified genomic fragment was added with Methylated Adapter to the 5’ and 3’ ends of each strand and was purified with QIAquick Gel Extraction kit (QIAGEN) and Then bisulphite treated with an EZ DNA Methylation GoldTM kit (Zymo Research). Finally, a narrow 350–400 bp size range after PCR with uracil-tolerant DNA polymerase for 10 cycles was purified with QIAquick Gel Extraction kit (QIAGEN). Libraries were subjected to QC on a bioanalyzer before sequencing on an Illumina HiSeq2000 using 2 \( \times 150 \) paired-end reads. Low-quality reads were removed and low quality nucleotides were clipped from the beginning and the end of sequences following the same procedure used for the RNA data. Processed reads were then mapped to both the *Brassica napus* reference genome and chloroplast genome using Bismark v0.16.3 with the following parameters: -non_directional -N 1 -l 32. The generated SAM files were then fed into the deduplicate_bismark subprogram for deduplication and the bismark_methylation_extractor subprogram to extract the methylation.
The average methylation level of MEG, PEG and non-imb gene were calculated as following method: Gene body regions were separated into 60 bins, and extended 5-kb up- and downstream regions were separated into 50 bins. The average methylation levels for each bin was calculated by the ratio of the methylated Cs/Gs and total number of Cs/Gs and Ts/As from all sites with more than four reads in each bin.60

2.10. Analysis and identification of pDMRs
Reads overlapping SNP were classified as maternal, paternal using custom Perl programs, in which C > T SNPs for forward reads and G > A SNPs for reverse reads were ignored. The reads from forward and reverse strand were merged together. The metilene was used to identify CG, CHG and CHH pDMR.61 Regions containing more than five CGs/CHGs/CHHs supported with at least four reads were further analysed and significant regions were kept (P < 0.05).

Approximately 0.8–1.0 g of genomic DNA was used for bisulphite conversion using an EZ DNA Methylation GoldTM kit (Zymo Research) according to the manufacturer’s instructions. Primers were designed based on methylated sequences located on the regulatory regions and within rapeseed imprinted genes (Supplementary Table S2). Bisulphite-treated DNA was then amplified using HotStarTaq Plus polymerase (Qiagen). PCR products were cloned into a Peasy-Blunt Zero Cloning Vector (TransGen Biotech) for sequencing. At least 20 randomly chosen clones were sequenced for each gene, and the methylation levels were calculated as a percentage (%) per site for each CG type of cytosine residues.

3. Results
3.1. Genome-wide screening of putative imprinted genes in rapeseed endosperm
During rapeseed silique development, endosperm volume reached a maximum after about 30 DAP and then gradually decreased during seed maturation (Supplementary Fig. S1a). Therefore the developing endosperm tissues (30 DAP) from hybrid endosperms (Fig. 1A). Based on saturation curve of rapeseed expression genes in every hybrid endosperm sample, sequence abundance is enough for imprinted gene analysis (Supplementary Fig. S5). The two replicates were considered separately and the imprinted genes were identified as overlap among two replicates. In total, about 14,394 genes covered by at least 10 reads could be assigned to a specific allele in both of the two reciprocal hybrid endosperms (Fig. 1A).

Analysis of gene expression showed that most of genes with SNP sites in the maternal and paternal alleles had a 2:1 allelic-specific expression ratio in four endosperm samples, indicating unbiased expression (Fig. 1B). A small proportion of genes (~3.8%) exhibited parent-of-origin differences in the expression of parental alleles in all samples. Among these, 267 genes exhibited maternally biased expression (MEG) and 26 genes displayed paternally biased expression (PEG) (Fig. 1A) in the reciprocal hybrids ($\chi^2$ test, $q < 0.05$). These differentially expressed genes were considered as putative imprinted loci. To obtain a high-confidence list of imprinted genes, stringent standards were applied (for MEGs, >85% of the reads supporting expression originated from the maternal allele; for PEGs, >50% of the reads supporting expression originated from the paternal allele). Using these criteria, 183 candidate imprinted genes were identified. Interestingly, most of the imprinted genes identified in our mRNA-seq analysis exhibited partial imprinting (preferential expression of one allele) rather than complete imprinting (strict monoallelic expression). Of these imprinted genes, 167 genes were maternally expressed and 16 genes were paternally expressed (Fig. 1B, Supplementary Tables S3 and S4). We also found that when the gene was imprinted in one subgenome, the counterpart in another genome was not always imprinted. In all imprinted genes, only seven genes in A subgenome could be found to also be imprinted in C subgenome. Another 30 homologous with imprinted genes which have SNP and expression are not imprinted. Additionally, as an allotetraploid, B. napus also exhibits expression bias between the homoeologous from A$_n$ and C$_n$ genomes. For example, in the endosperms of 93,275, we found 8,401 gene pairs showing expression bias including 60 imprinted genes (Supplementary Fig. S6; Table S3).

3.2. Validation of imprinted rapeseed genes
To determine whether the imprinted genes in rapeseed are conserved in other species, the rapeseed imprinted genes were compared with the known imprinted genes identified from other plants. The results showed there were 16, 2, 1 and 1 rapeseed imprinted genes which were also imprinted in Arabidopsis, sorghum, castor bean, maize and rice, respectively (Table 1).

In addition to the imprinted genes which have been reported in other species, it is necessary to confirm the accuracy of some other imprinted genes identified in rapeseed. However, because of high homology between the A$_n$ subgenome and C$_n$ subgenome in rapeseed, it is very difficult to separate the imprinted genes from its homologs.
by designing specific primers for most of rapeseed genes. By analyzing differences in the sequences and expression of the imprinted genes and their homologs, 11 MEGs and 1 PEGs with nearly complete imprinting were screened for confirmation of the putative imprinting by measuring their allelic-specific expression patterns. RT-PCR and sequencing analysis with the total RNA from 30 DAP hybrid endosperm showed that the expression patterns of these 12 tested imprinted genes were consistent with the mRNA-seq results (Fig. 2A).

These 12 imprinted genes were also used to assess whether imprinting could be detected in other reciprocal hybrid combinations (9T364 and 93275; 61,616 and 93,275; zy036 and 93,275). The genes which had the same SNPs that were found between YN171 and 93275 were identified to be imprinted in the other reciprocal hybrid combinations (Supplementary Fig. S7), leading to the conclusion that the predicted imprinted genes were subjected to genomic imprinting and, furthermore, that mRNA-seq analysis was a reliable method to identify parent-of-origin-specific gene expression.

3.3. Characterization of imprinted genes in rapeseed

To further characterize the imprinted genes in rapeseed, the allelic-specific expression pattern of 5 of those 16 imprinted genes were analysed at earlier and later stages of endosperm development based on RNA-seq analysis. The five imprinted genes maintained their imprinting pattern at the different developmental stages, at five different time points, 15, 20, 25, 30 and 35 DAP. This result suggested that imprinted genes maintained stable regulation of expression during the development of rapeseed (Supplementary Fig. S8).
Furthermore, previous studies have found that most imprinted genes in plants show specific or preferential expression in the endosperm. To test whether this is true in rapeseed, transcriptome sequencing data was used to analyse the expression pattern of the validated rapeseed imprinted genes in other tissues of YN171, including the leaf, stem, root, bud, pericarp, embryo and flower. Figure 2B and C shows the results of clustering analysis of MEGs and PEGs based on their expression in eight tissues. Unlike some earlier studies, only about 13% (22 of 167) of the maternally imprinted genes were expressed specifically or preferentially in the endosperm in rapeseed, and in most cases the expression was elevated in the endosperm but not endosperm specifically. Further functional analyses of rapeseed imprinted genes suggested that many targets of rapeseed imprinting might have important regulatory roles. Further functional analyses of rapeseed imprinted genes (Fig. 4A). The molecular functions were enriched in receptor binding or activity, transcription factor activity, other enzyme activity, transporter activity and transferase activity (q < 0.001). MapMan analysis showed that genes related to the hormone metabolism, cell wall, transport, lipid metabolism, secondary metabolism, and cellular processes were enriched (P < 0.001) (Fig. 4B). Additionally, over 50 imprinted genes were predicted to localize in nucleus and showed binding activities and regulation of transcription functions suggesting that many targets of rapeseed imprinting might have important regulatory roles. Further functional analyses of rapeseed imprinted genes will be necessary to shed light on the functional consequences on seed development.

### 3.4. Functional annotation analysis of the imprinted genes

To annotate the function of the rapeseed imprinted genes, the protein sequences encoded by these genes were used to search the homologous Arabidopsis proteins by BLASTP. When the GO biological process annotations of the imprinted genes were assessed, the 183 rapeseed imprinted genes (q < 0.001) were allocated into several biological processes, including other biological processes, response to abiotic or biotic stimulus, response to stress, transport, other metabolic processes, other cellular processes and protein metabolism (Fig. 4A). The molecular functions were enriched in receptor binding or activity, transcription factor activity, other enzyme activity, transporter activity and transferase activity (q < 0.001). MapMan analysis showed that genes related to the hormone metabolism, cell wall, transport, lipid metabolism, secondary metabolism, and cellular processes were enriched (P < 0.001) (Fig. 4B). Additionally, over 50 imprinted genes were predicted to localize in nucleus and showed binding activities and regulation of transcription functions suggesting that many targets of rapeseed imprinting might have important regulatory roles. Further functional analyses of rapeseed imprinted genes will be necessary to shed light on the functional consequences on seed development.

### 3.5. Association analysis between allele-differential cytosine methylation and imprinted gene expression in rapeseed

To analyse the association of allele-differential cytosine methylation and imprinted gene expression in rapeseed, bisulphite sequencing
to the genomic DNA from endosperm at 30 DAP was performed to assess the patterns of DNA methylation of both maternal and paternal alleles of imprinted loci. The chloroplast genome was used to estimate the bisulphite conversion rate. We found the average methylation levels of CG, CHG and CHH were 1.39%, 0.625% and 0.51%, which showed the bisulphite conversion rate of genome sequence exceeded more than 98%. The methylation level from 5 kb upstream to 5 kb downstream of genes was analysed to investigate the DNA methylation pattern around CG, CHG and CHH residues. No significant difference was found in DNA methylation patterns of MEGs, PEGs and non-imprinted genes in CG methylation levels in the four endosperm samples originating from YN171, 93275 and the reciprocal hybrids. The overall CG methylation level was higher in PEGs than that in MEGs. However, the CG methylation level of PEGs was not significantly different from that of non-imprinted genes, although the overall levels of DNA methylation in the upstream regions of PEGs were slightly higher (Fig. 5A). At CHG and CHH residues, the methylation levels showed obvious difference in parents. CHG methylation levels showed similar in MEGs and PEGs and both of them exhibited slightly but statistically significant lower levels than non-imprinted genes, while CHH methylation levels had no significant difference between imprinted genes and non-imprinted genes (Fig. 5B and C). To eliminate the effects of non-additive gene expression in polyploidy rapeseed, we screened the homoeologous gene pairs showing significant differences in CG methylation and found 1294 gene pairs showing an inverse correlation between degree of methylation and gene expression. Among them, although 10 imprinted genes are found, most of them showed low methylation level in both of A_0 and C_n homoeologs (Supplementary Fig. S10, Table S8).

In order to detect allele-specific methylation of a gene, it must have sufficient read coverage of both the maternal and paternal genomic alleles in regions with at least one SNP between YN171 and 93275. A total of 104 genes (93 MEGs, 11 PEGs) met this criterion. Of these, seven MEGs and four PEGs showed coherent differential CG methylation between the two parental alleles in reciprocal hybrid endosperm DNAs (Supplementary Table S9). Of the 11 genes with differential CG methylation, 6 DMRs were located within the gene, 4 located at upstream region and 1 located at downstream. The identified DMRs ranged from 72 to 493 bp in length. Surprisingly,
contrary to the studies in other plants, seven identified DMRs in rapeseed imprinted genes exhibited low CG methylation rates no matter in maternal alleles or in paternal alleles and had no big difference between parent alleles. Among the other four genes with higher CG methylation rates (BnC04g0675620, BnA09g0355640, BnA04g0165340 and BnA10g0421660), only BnC04g0675620 showed obvious differential methylation levels. To further confirm the reliability of methylation analysis, two genes including 1 MEG (BnA03g0127610) and 1 PEG (BnC08g0868840) were chose to investigate allele-specific methylation patterns. The results indicated that 2 genes showed difference in methylation states between parental alleles, which was coincident with the results from bisulphite sequencing data (Supplementary Fig. S11). Taken together, we found 1 gene (BnC04g0675620) showing high methylation levels in at least 104 imprinted genes. Additionally, lower methylation levels of CHG and CHH were found in some imprinted genes, which showed no great difference between parent alleles (Supplementary Table S9).

4. Discussion

As a result of the increasing focus on the importance of epigenetic mechanisms in fundamental biological processes together with the rapid advancement of RNA-seq technologies, studies of imprinting in diverse plant species, especially in several important crops, have intensified in recent years.25–27,63 At present, with the exception of Arabidopsis and Capsella rubella, there have been no studies on gene imprinting in the endosperms of Cruciferae species.64 Rapeseed is an important oil crop, and a genome-wide survey of imprinted genes in rapeseed endosperm may improve our understanding of storage accumulation and offer another model system for seed development. The first step toward understanding the biological functions of imprinting is to identify a large set of imprinted genes in a given species, although imprinted genes may differ between species and even between different strains within a species.57

In this study, we applied high-throughput mRNA-seq analysis with highly stringent selection criteria and identified 167 MEGs and 16 PEGs from hybrid rapeseed endosperms originating from YN171 and 93275. Although imprinted genes are well conserved in mammals, it is obvious that the overlap of imprinted genes in different plant species is rather limited.23,27 Even between Capsella rubella and Arabidopsis, which share a common ancestor ~10–14 million years ago, only about 14% of the MEGs and 29% of the PEGs were found to overlap.64 In our study, although limited overlap of identified rapeseed imprinting genes with other plants supports less conserved imprinting in plants, among the 16 genes overlapped with Arabidopsis imprinted genes, BnC03g0699760, the homolog of Arabidopsis FIS2, was the most well-known imprinted gene in Arabidopsis.63 FIS2 in the endosperm is initiated and established by DME-mediated active DNA demethylation in the central cell, while the paternal alleles remain methylated and silenced.34 Unfortunately, we could not confirm if the homolog of FIS2 on A05 chromosome is also imprinted because of SNP absence. BnA04g0178370, BnA04g0178970 and BnC04g0675620 are homologs of
Arabidopsis JMJ15. JMJ15 is closely related to JMJ14, which is thought to demethylate trimethylated lysine 4 of histone H3 (H3K4me3) and is involved in DRM2-mediated maintenance of DNA methylation.\textsuperscript{21}

Previous studies have not conclusively resolved whether imprinting of genes is conserved across the different stages of seed development. Xin et al.\textsuperscript{66} showed that MEGs and PEGs in maize exhibited distinct patterns of gene imprinting during endosperm development, with PEGs predominantly detected at 7 DAP and MEGs predominantly detected at 10 DAP. In castor bean, most of the 19 imprinted genes (17 MEGs and 2 PEGs) were found to maintain their imprinting pattern throughout endosperm development.\textsuperscript{26} In this study, the imprinting status of 5 rapeseed imprinted genes across the 5 developmental stages of endosperm from 15 to 35 DAP were highly similar, suggesting that imprinting occurs early and persists in the rapeseed endosperm.

In plants, the proportions of MEGs and PEGs showing preferential expression in endosperm were dramatically different in maize (61.5\% MEGs and 76.5\% PEGs) and sorghum (96.7\% MEGs and 93.7\% PEGs).\textsuperscript{25,27} However, our analysis showed that most of the rapeseed imprinted genes (66\% MEGs and 67\% PEGs) were also expressed in other tissues, rather than endosperm specific. The difference between our findings and previous studies might be due to the number of tissues analysed. Only four other tissues were analysed in maize and five in sorghum, while we used seven other tissues in rapeseed. This conjecture was further supported by our finding that about 30\% of the imprinted genes were expressed in only two or three tissues. In castor bean, where measurements were made in a range of tissues, including the leaf, stem, root, male and female flower, embryo and endosperm, the expression of 67\% of the MEGs and 17\% of the PEGs was observed in other tissues.\textsuperscript{26} These results suggest that most imprinted genes also play important roles in the development of other tissues.

Gene expression within a cluster is often orchestrated by a common imprinting control region.\textsuperscript{67} Although physical clustering is a hallmark of imprinted genes in mammals, this does not appear to be a consistent characteristic in plants. The proportion of clustered genes in plants is highly variable. In Arabidopsis, 21 of 208 (10\%)
and 18 of 66 (27%) imprinted genes showed evidence of clustering.\textsuperscript{44,68} In maize, only 2 clusters including 4 genes were found in a set of 100 imprinted genes (4%),\textsuperscript{23} however, clear clustering of 74 of 217 (34%) imprinted genes and non-coding RNAs was observed from the same reciprocal hybrids between the two parental lines (B73 and Mo17).\textsuperscript{68} An analysis of 100 sorghum imprinted genes revealed that only 28 genes (28%) were grouped in 12 clusters. Unlike the results reported in other plant species, we found that 48.6% of the identified imprinted genes in rapeseed formed 34 distinct clusters, although the clustering criteria we used were similar to those in other studies. This suggested that clustering might play a role in imprinted gene regulation in rapeseed as it is in mammals.

In some other plant species, most of DMRs identified are uniformly hypomethylated in maternal alleles and hypermethylated in paternal alleles.\textsuperscript{26–68} In maize, the identified DMRs have an average of 14% CG methylation in maternal alleles and 94% in paternal alleles.\textsuperscript{68} In castor bean, the average is 9.6% in maternal alleles and 86% in paternal alleles.\textsuperscript{24} In this study, among the 104 genes (93 MEGs, 11 PEGs) that cytosine methylation levels could be analysed, 4 PEGs showed differential CG sites with the average CG methylation rates of over 30% in maternal alleles and about 3% in paternal alleles.\textsuperscript{24} In this study, among the 104 genes (93 MEGs, 11 PEGs) that cytosine methylation levels could be analysed, 4 PEGs showed differential CG sites with the average CG methylation rates of over 30% in maternal alleles and about 3% in paternal alleles.\textsuperscript{24} Only one MEG showing differential methylation levels exhibited high CG methylation rates. Additionally, we also found over 5000 non-coding RNAs in 93275 and YN171. By analysing the locations and expression levels of the non-coding RNA, we found 1 RNA showing exonic overlap on the opposite strand with \textit{BnC03g0568040} gene and 14 RNAs locating near 9 imprinted genes (the distance is about 1–40 kb) (Supplementary Table S10). However, all the RNAs nearly do not affect the expression of imprinted genes. Because DNA methylation is a classic epigenetic mechanism and has been considered a major regulator of gene imprinting, the methylation mechanisms of imprinting in rapeseed imprinted genes is worth being investigated further.

Authors’ contributions
J.L. and W.H. conceived the research plans; H.L. analysed the sequence data; S.F. performed the experiments; S.S., X.Z. and Z.H. provided technical assistance; J.L. and J.L. wrote the article.

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Conflict of interest
None declared.

Supplementary data
Supplementary data are available at DNARES online.

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