Comparative analysis of circular RNAs between soybean cytoplasmic male-sterile line NJCMS1A and its maintainer NJCMS1B by high-throughput sequencing

Linfeng Chen¹, Xianlong Ding¹, Hao Zhang¹, Tingting He¹, Yanwei Li¹, Tanliu Wang¹, Xiaojing Li¹, Ling Jin¹, Qijian Song², Shouping Yang¹* and Junyi Gai¹*

Abstract

Background: Cytoplasmic male sterility (CMS) is a natural phenomenon of pollen abortion caused by the interaction between cytoplasmic genes and nuclear genes. CMS is a simple and effective pollination control system, and plays an important role in crop heterosis utilization. Circular RNAs (circRNAs) are a vital type of non-coding RNAs, which play crucial roles in microRNAs (miRNAs) function and post-transcription control. To explore the expression profile and possible functions of circRNAs in the soybean CMS line NJCMS1A and its maintainer NJCMS1B, high-throughput deep sequencing coupled with RNase R enrichment strategy was conducted.

Results: CircRNA libraries were constructed from flower buds of NJCMS1A and its maintainer NJCMS1B with three biological replicates. A total of 2867 circRNAs were identified, with 1009 circRNAs differentially expressed between NJCMS1A and NJCMS1B based on analysis of high-throughput sequencing. Of the 12 randomly selected circRNAs with different expression levels, 10 showed consistent expression patterns based on high-throughput sequencing and quantitative real-time PCR analyses. Tissue specific expression patterns were also verified with two circRNAs by quantitative real-time PCR. Most parental genes of differentially expressed circRNAs were mainly involved in biological processes such as metabolic process, biological regulation, and reproductive process. Moreover, 83 miRNAs were predicted from the differentially expressed circRNAs, some of which were strongly related to pollen development and male fertility; The functions of miRNA targets were analyzed using Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG), and the target mRNAs were significantly enriched in signal transduction and programmed cell death. Furthermore, a total of 165 soybean circRNAs were predicted to contain at least one internal ribosome entry site (IRES) element and an open reading frame, indicating their potential to encode polypeptides or proteins.

Conclusions: Our study indicated that the circRNAs might participate in the regulation of flower and pollen development, which could provide a new insight into the molecular mechanisms of CMS in soybean.

Keywords: Soybean (Glycine max (L.) Merr.), Cytoplasmic male sterility, CircRNAs, High-throughput sequencing, Parental genes, Binding miRNAs
Background

Soybean (Glycine max (L.) Merr.) is an important oil and protein crop, but soybean yield is low. Utilization of heterosis has been proved to be one of the effective methods to increase crop yield, and cytoplasmic male sterility (CMS) plays an important role in heterosis utilization [1]. CMS is a maternally inherited phenotype caused by the interaction of cytoplasmic genes and nuclear genes, which lead to pollen abortion but normal pistil development [2, 3]. To date, CMS has been observed in over 200 plant species [4] and is often obtained in wild germplasm or by inter- or intra-subspecies backcrossing [5]. In soybean, CMS was first reported by Davis [6], after which many studies on CMS were conducted [7, 8]. In recent years, transcriptomics [9], proteomics [10], microRNA [11], and DNA methylation of CMS have been explored [12], but the function of circular RNAs (circRNAs) in soybean CMS has not been reported.

CircRNAs are a class of endogenous noncoding RNAs, which do not have 5’ caps and 3’ tails and form a ring structure with covalent bonds [13]. CircRNAs were first reported based on deep sequencing of RNA by Salzman et al. [14], which were neglected for decades because they were considered as transcriptional noise or reverse transcription PCR artifacts by-product [15]. With the development of high-throughput sequencing technology and bioinformatics, circRNAs have been identified in all domains of life, including eukaryotes [15, 16], archaea [17], bacteria [18], and viruses [19]. Previous studies showed that circRNAs were more resistant to degradation by RNase R than their linear counterparts [20], and had tissue, cell-type, and developmental-stage specific expression patterns [15, 21, 22]. Li et al. [23] demonstrated that the circRNAs located in the nucleus could promote transcription of their parental genes via specific RNA-RNA interactions, but another recent study revealed that circRNAs derived from organelle genome could also regulate gene expression [24]. Furthermore, circRNAs act as miRNA sponges and prevent them from inhibiting their target miRNAs [25, 26]. Most recent studies have demonstrated that some circRNAs can be translated into polypeptides or proteins by translation initiation element internal ribosome entry site (IRES) or N6-methyladenosine (m6A) [27, 28].

Although not as comprehensive as in animals, the exploration of circRNAs in plants is increasing. Differential expression of circRNAs has been reported in plants, for example, 27 rice exonic circRNAs were associated with phosphate starvation responsive expression [16], 163 tomato circRNAs with chilling responsive expression [29], and 62 wheat circRNAs with dehydration stress specific expression [30]. Moreover, some circRNAs showed time, tissue, species, or developmental-stage specific expression patterns in plants [16, 24, 31, 32]. Unlike the positive regulation of circRNAs in animals [23], over-expression of circRNAs in rice and tomato reduced expression level of their parental genes [33, 34]. In soybean, a total of 5372 circRNAs were identified, of which approximately 80% were generated from paralogous genes [35]. Meanwhile, Zhao et al. [35] found that up-regulation of circRNAs might decrease the activity of target miRNAs and increase expression of the related mRNAs.

To explore the expression profile and possible functions of circRNAs in the soybean CMS line NJCMS1A vs. its maintainer line NJCMS1B, high-throughput deep sequencing coupled with RNase R enrichment strategy was conducted. Target miRNAs of differentially expressed circRNAs and the correlated mRNAs were predicted using bioinformatics methods, and their potential functions were further analyzed. Our study investigated the possible role of circRNAs in CMS for the first time, and the results showed that the circRNAs might participate in the regulation of flower and pollen development in soybean.

Methods

Plant materials and sample collection

The soybean cytoplasmic male-sterile line NJCMS1A was developed through consecutive backcross. The cultivar N8855 was the donor and cultivar N2899 (designated as NJCMS1B afterwards) was a recurrent parent [8, 36, 37]. NJCMS1A and NJCMS1B are near-isogenic lines of isonuclear alloplasmic type with similar nucleus but different cytoplasm. NJCMS1A and NJCMS1B were grown in the summer of 2016 at Dangtu Experimental Station, National Center for Soybean Improvement, Nanjing Agricultural University, Maanshan, Anhui, China. Male sterile plants were identified by observing dehiscence of anthers, germination rate of pollen, and morphological traits of plants at maturity. Because it is difficult to determine development stage of pollen based on flower bud size in soybean, during the flowering period, the flower buds of different sizes were collected from NJCMS1A or NJCMS1B plants and mixed, with three biological replicates per line. Meanwhile, roots, stems and leaves of soybean were also collected at the flowering period with three biological replicates. The tissues were immediately frozen in liquid nitrogen and stored at − 80 °C for further use.

Total RNA extraction and RNase R treatment

Total RNA from flower buds of NJCMS1A and NJCMS1B lines was extracted using the RNeasy Plant Kit (Qiagen, DEU) according to the manufacturer’s protocol, DNA contamination was removed using the DNase I contained in the Kit. For RNase R-treated total
RNA samples, the purified DNaseI-treated total RNA was incubated for 15 min at 37 °C with 3 units RNase R per μg of total RNA (Epicentre, Shanghai, CN). RNA was subsequently purified using an RNase MinElute Cleaning Kit (Qiagen, DEU). To obtain accurate and sufficient transcriptome data, RNA from three biological replicates of NJCMS1A and NJCMS1B lines were sequenced.

**CircRNA library construction and sequencing**

rRNA-depleted RNAs in 5 μg RNA per sample was obtained using the Epicentre Ribozero™ rRNA Removal Kit (Epicentre, USA) and were further treated with RNase R (Epicentre, USA) for Trizol extraction. Sequencing libraries were prepared using NEBNext Ultra Directional RNA Library Prep Kit (NEB, USA) following manufacturer's instructions. The libraries were preliminarily quantified by Qubit and diluted to 1 ng/μl. The insert size of the libraries was detected by Agilent 2100/Caliper, which was expected to be distributed around 250–300 bp. The effective concentration of the libraries was accurately quantified by qRT-PCR, and the effective concentration was greater than 2 nM to ensure the libraries quality. Briefly, fragmentation was carried out using divalent cations under elevated temperature in NEBNext First-strand Synthesis Reaction Buffer. First-strand cDNA was synthesized using random hexamer primer and M-MuLV reverse transcriptase (RNaseH). Second-strand cDNA synthesis was subsequently performed using DNA Polymerase I and RNase H. In the reaction buffer, dTTP in dNTPs was replaced by dUTP. Remaining overhangs were blunted via exonuclease/polymerase. After adenylation of 3′ ends of DNA fragments, NEBNext Adaptor with hairpin loop structure was ligated for hybridization. In order to obtain cDNA fragments of 150–200 bp in length, the library fragments were purified with the AMPure XP system (Beckman Coulter, Beverly, USA). Subsequently, 3 μl USER Enzyme (NEB, USA) was added to adaptor-ligated cDNA of 150–200 bp and kept at 37 °C for 15 min followed by 5 min at 95 °C before PCR. PCR was performed with Phusion high-fidelity DNA polymerase, universal PCR primer, and index (X) primer. Finally, the library was purified (AMPure XP system) and then qualified by the Agilent (AMPure XP system) and then qualified by the Agilent Bioanalyzer 2100 system. Clustering of the index-coded samples was performed on a cBot Cluster Generation System using HiSeq PE Cluster Kit v4 cBot (Illumina) according to the manufacturer's instructions. After cluster generation, the libraries were sequenced on an Illumina HiSeq 2500 platform and 125 bp paired-end reads were generated. Library construction and sequencing were carried out by Novogene (Novogene, Beijing, China).

**Identification of circRNAs and differential expression analysis**

Raw data (raw reads in fastq format) was first processed by a custom perl script. Sequence reads with adapter, poly-N, and low-quality were eliminated. The remaining reads were used for read quality and GC content estimation, and downstream calculation. Soybean reference genome and gene annotation were downloaded from the website at https://phytozome.jgi.doe.gov/pz/portal.html#. The index of the reference genome was built using Bowtie v2.0.6. Differential expression analysis was carried out with DESeq2 (version 1.6.3) [39]. Differentially expressed circRNAs were identified with the cutoff threshold of |log2 (fold-changes)| ≥ 2 based on the method used by Wang et al. [30] and Liu et al. [31].

**Prediction of miRNA targets of circRNAs, mRNA targets of miRNA, and annotation of functions**

miRNA binding sites of differentially expressed circRNAs were predicted by psRobotTar in psRobot [40]. Meanwhile, the circRNA-miRNA interaction network was delineated by Cytoscape [41]. The obtained miRNAs were used to predict the target mRNAs by psRobot with default parameters [40]. The parental genes of differentially expressed circRNAs and the predicted target mRNAs were classified into different functional processes based on Gene Ontology (GO) term enrichment using the Web Gene Ontology Annotation Plot (WEGO) [42] and agriGO [43, 44], respectively. The KOBAS [45] was used for KEGG pathways enrichment analysis. The gene annotation of *A. thaliana* at http://www.arabidopsis.org/ was used to define functions of homologous genes in soybean.

**PCR amplification and sanger sequencing**

To validate the soybean circRNAs identified in this study, a set of 12 differentially expressed circRNAs were amplified with divergent primer. As a control, a pair of convergent primers were designed for gma-circRNA00002 (Additional file 1). All primers were synthesized by Invitrogen (Shanghai, China). The total and RNase R-treated
RNA of each sample were used as templates. PCR products were separated using agarose gel, and each band was excised and purified using the AxyPrep DNA Gel Extraction Kit (Axygen, Suzhou, China) for Sanger-sequencing.

**Quantitative real-time PCR validation**

Quantitative real-time PCR (qRT-PCR) was carried out to validate differential expression levels of circRNAs. Divergent primers were designed in order to obtain amplicon from circle template (Additional file 1). According to the instruction of the iScript Select cDNA Synthesis Kit (BIO-RAD, USA), 1 μg of the total RNA untreated with RNase R was reverse-transcribed with random primers. Expression of circRNAs was quantified using the iTag Universal SYBR Green Supermix (BIO-RAD, USA) on the Bio-Rad CFX96 machine (CFX96 Touch, BIO-RAD, USA). All real-time PCR assays were performed with three biological replicates, and the expression of the housekeeping gene GADPH was used as a reference for data normalization. The qRT-PCR aliquot contained 2 μL cDNA, 0.6 μL of each upstream and downstream primers (10 μM), 10 μL Takara SYBR Premix Ex Taq, and 6.8 μL RNase-free ddH2O and performed with an initial denaturation at 95 °C for 30 s, followed by 40 cycles at 95 °C for 5 s, 60 °C for 30 s. The amplification curve and melting curve were examined to evaluate specific amplification. The circRNAs relative expression levels (log2 values) were calculated using the $2^{-\Delta\Delta C_t}$ method. Student’s t-test was performed to compare differences of circRNAs expression between NJCMS1A and NJCMS1B. The probability threshold of significance was $P < 0.05$.

**Protein-coding potential prediction of circRNAs**

The IRES element is required to initiate translation of a mRNA sequence without a 5’-cap structure [46]. If a circRNA has at least one IRES element, it may be able to encode a protein. To predict the IRES elements in soybean circRNAs, we blasted sequences of the circRNAs to all the IRES sequences in the website (http://iresite.org/) at an E-value < 0.05 [47]. To predict the ORFs of predictive circRNAs, we used the prediction algorithm at the website (https://github.com/kadenerlab/cORF_pipeline). Briefly, the longest ORF spanning the circRNA junction was selected for further analysis. The possible coding products of the circRNAs with protein-coding potential were used to predict the conserved domains using the Conserved Domain Database (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi).

**Results and discussion**

**Identification of circRNAs in soybean**

Total amount of sequence for each RNA libraries was ≥9 Gb with Q20 ≥ 93%, Q30 ≥ 85%, and an error rate ≤ 0.05 (Table 1), the depth and accuracy of high quality sequence was sufficient for subsequent analysis.

A total of 2867 circRNAs were identified, which included 1722 from NJCMS1B and 1643 from NJCMS1A (Fig. 1a, Additional file 2). Among them, 452 (15.8%) were derived from exons of a single protein-coding gene (exonic circRNAs), 821 (28.6%) from introns (intronic circRNAs), 293 (10.2%) from exons and introns of one or more genes (exon-intron circRNAs), and 1301 (45.4%) from intergenic regions (intergenic circRNAs) (Fig. 1b). CircRNAs primarily derived from exons in several plants have been reported, e.g. 5152 (85.7%) circRNAs in A. thaliana [16], 615 (72.0%) in tomato [29], and 1453 (93.7%) in maize [48]. However, only 2494 (46.4%) exonic circRNAs were identified in soybean by Zhao et al. [35], which was similar to our result. The length of circRNAs in this study ranged from 150 to 44,756 bp, but most (92.8%) were < 2000 bp (Fig. 1c). The length distribution of soybean circRNAs in this study was similar to that reported by Zhao et al. [35]. Additionally, the sequence alignment between the circRNAs inform this study and previous study (Zhao et al. [35]) using blastn indicated that only 78 circRNAs were homologous (Additional file 3) at e-value < 1e−5 and identity > 85%. The small proportion of homologous circRNAs in these two studies might be associated with tissue specific expression pattern of circRNA (Additional file 4) and use of different circRNAs prediction software. Besides, the number of circRNAs in different chromosomes and the densities of circRNAs in different chromosomal regions were also different (Fig. 1d).

**Expression profiling of circRNAs and qRT-PCR validation**

A total of 1009 circRNAs with at least two-fold (log2) change of expression level between NJCMS1A and NJCMS1B were identified by high-throughput sequencing. Of these, a total of 360 circRNAs were up-regulated, 649 circRNAs were down-regulated in NJCMS1A vs. NJCMS1B (Additional file 5).

The back splicing sites and expression profiles of the 12 randomly selected from 1009 circRNAs were further experimentally validated. We treated total RNA samples with RNase R to verify the special stability (resistance to exonuclease-mediated degradation) of circRNAs and eliminated interference from linear RNA. Divergent primers were designed to guarantee that amplifications were from circular templates and a pair of convergent primers for gma-circRNA0002 were designed as a control. Total RNA samples and RNase R-treated samples were used as reaction templates to amplify circRNAs. As
a result, all 12 pairs of divergent primers yielded amplification products with the expected length from cDNAs that with and without RNase R treatment. To demonstrate that the amplification products from the divergent primers were derived from the corresponding circRNAs and spanning the junction sites, we collected the amplification products for further detection by Sanger sequencing (Fig. 2 and Additional file 6). As shown in Fig. 3, among the 12 random-selected circRNAs, 10 showed consistent expression patterns with RNA-seq results. The coincidence rate between qRT-PCR and RNA-seq was 83.3%. Two inconsistent expression of circRNAs candidates (gma-circ2468 and gma-circ2848) may be caused by the low expression levels of the two circRNAs.

### Table 1 Summary of circRNA sequencing data

| Sample   | Replicates | Number of raw reads | Number of clean reads | Clean bases (Gb) | Error rate (100%) | Q20 (100%) | Q30 (100%) | GC content (100%) |
|----------|------------|---------------------|-----------------------|------------------|-------------------|-------------|-------------|-------------------|
| NJCMS1A-1 | 84,474,228 | 80,699,898          | 12.1                  | 0.05             | 93.38             | 85.41       |             | 54.16             |
| NJCMS1A   | 67,528,014 | 63,913,764          | 9.58                  | 0.02             | 97.85             | 94.07       |             | 53.75             |
| NJCMS1A-2 | 98,550,080 | 92,713,788          | 13.9                  | 0.03             | 97.29             | 93.12       |             | 54.55             |
| NJCMS1B-1 | 74,471,198 | 68,964,198          | 10.34                 | 0.03             | 97.39             | 93.08       |             | 53.11             |
| NJCMS1B   | 77,162,554 | 70,527,006          | 10.58                 | 0.03             | 97.04             | 93.03       |             | 53.94             |
| NJCMS1B-3 | 88,067,914 | 84,489,922          | 12.68                 | 0.05             | 93.53             | 85.55       |             | 54.67             |

**Fig. 1** CircRNA sequencing data. (a) Venn diagram shows the number of the identified circRNAs in NJCMS1A and NJCMS1B. (b) Source statistics of the circRNAs. (c) Sequence length distribution of the circRNAs in different samples. (d) Circle plot shows the distribution of the identified circRNAs in soybean chromosomes and their expression levels. The outermost layer represents all soybean chromosomes (Chr01-Chr20). The middle blue lines show the distribution of the circRNAs in soybean chromosomes, while the denser lines indicate more circRNA distribution. The innermost green lines show expression levels of the circRNAs, and the height of the lines indicates the level of expression.
Tissue specific expression patterns validation by qRT-PCR

Previous studies showed that circRNAs were tissue-preferentially expressed [15, 24]. To verify this, two highly expressed circRNAs (gma-circRNA0002 and gma-circRNA2483) based on the high throughput sequencing were selected for qRT-PCR analysis. These two circRNAs were expressed only in leaves and flower buds of NJCMS1A and NJCMS1B, but not in roots and stems (Fig. 4). The expression level of gma-circRNA0002 was only significantly different in the flower buds, while the expression levels of gma-circRNA2483 were significantly different in the leaves and flower buds between NJCMS1A and NJCMS1B.

Functional categorization of parental genes of differentially expressed circRNAs

It has been reported that circRNAs could regulate expression of their parental genes [33, 34]. To understand the possible functions of circRNAs in CMS of soybean, the parental genes of the differentially expressed circRNAs were predicted. A total of 545 parental genes were obtained from the 1009 differentially expressed circRNAs. GO classification showed that these genes are involved in a wide range of biological processes, such as metabolic process (GO:0008152), biological regulation...
(GO:0065007), and cellular process (GO:000987) (Fig. 5). Interestingly, a fraction of parental genes was classified into the categories of reproduction (GO:0000003) and reproductive process (GO:0022414). Among the cellular components, cell (GO:0005623), cell part (GO:0044464), and organelle (GO:0043226) accounted for a large proportion. In the molecular functions, the two main categories were binding (GO:0005488) and catalytic (GO:0003824). GO analysis of the parental genes showed that the differentially expressed circRNAs from NJCNS1A and NJCMS1B were associated with various functions in different biological processes, cellular components, and molecular function, indicating that circRNAs may play an important role in the fertility of soybean. KEgg pathway analysis identified 7 pathways including valine, leucine and isoleucine degradation, selenocompound metabolism, RNA transport, synthesis and degradation of ketone bodies, ascorbate and aldarate metabolism, ubiquitin mediated proteolysis, and porphyrin and chlorophyll metabolism (Additional file 7). These pathways are mainly related to amino acid degradation and material metabolism.

Putative functions of parental genes of differentially expressed circRNAs in flower development and male fertility

Based on the gene annotation of A. thaliana, the function of many parental genes were related to flower development and male fertility. We randomly enumerated 20 parental genes (Table 2) and selected four of them to describe their functions in detail. Glyma.06G173500, the parental gene of gma-circRNA0717, was a homolog of the Glucan Synthase-Like 8 (GSL8) gene in A. thaliana. GSL8 and Glucan Synthase-Like 10 (GSL10) were two members of the A. thaliana GSL gene family, which are independently required for male gametophyte development and plant growth. Experiments showed that GSL8 and GSL10 T-DNA insertions led to pollen sterility [49]. GSL8 was believed to be involved in the synthesis of cell wall component callose [50]. It was hypothesized that gma-circRNA0717 may play a vital role in male gametophyte development. Glyma.13G230500, the parental gene of gma-circRNA1856, was an ortholog of the maternal effect embryo arrest 18 (MEE18) gene in A. thaliana. Mutation of MEE18 may affect pollen gametogenesis, pollen germination, pollen tube growth, polarity or guidance, and pollen tube-embryo sac interactions or fertilization [51]. X-ray induced 1 gene (XRI1) was a novel DNA repair factor and was essential for male and female meiosis, and homozygous XRI1 mutants caused complete sterility in A. thaliana [52]. In this study, Glyma.18G164900, the parental gene of gma-circRNA2481, was a homolog of the XRI1 gene. Glyma.18G118100, the parental gene of gma-circRNA2454, was a homolog of squalene epoxidase 1 (SQE1) gene in A. thaliana. Previous studies showed that SQE1 mutants displayed severe growth defects in A. thaliana, including short stature, short roots, and complete infertility [53]. We speculated that differences in circRNAs expression levels may influence the functions of pollen and male gametophytes, and result in CMS in NJCMS1A by interacting with their parental genes.

Characterization of binding miRNAs of differentially expressed circRNAs

CircRNAs have been reported to act as miRNA sponges regulating gene expression [25]. To verify if circRNAs

![Fig. 5 Gene Ontology (GO) annotation of parental genes of the differentially expressed circRNAs between NJCMS1A and NJCMS1B](image-url)
have a similar function in soybean, we predicted potential binding sites of miRNAs of the differentially expressed circRNAs. We observed that 72 differentially expressed circRNAs, of which 28 were up-regulated and 44 were down-regulated, contained 83 predicted circRNA-binding miRNAs. Of these circRNAs, only 24 had two to four miRNA binding sites (Additional file 8). Based on the predicted results, a circRNA-miRNA interaction network was delineated by Cytoscape (Fig. 6). The results showed that a single circRNA could target different miRNAs; for example, gma-circRNA0534 targeted to 16 miRNAs. Meanwhile, a single miRNA could be targeted by diverse circRNAs. For instance, gma-miR156 was targeted by 27 circRNAs.

Among the predicted miRNAs, three miRNAs (gma-miR156e, gma-miR4349, and gma-miR4993) were differentially expressed in our previous study [11]. In addition, we found that some miRNAs, such as miR156, miR162, miR169, and miR172, were important for pollen development and male fertility in soybean. miR156 is one of the highly conserved miRNA families, which was first reported in *A. thaliana* [54]. Squamosa promoter binding like (*SPL*) family proteins are involved in almost all physiological and biochemical processes such as morphogenesis, development stage transition, sporulation, flower and fruit development, external stress response, and hormone signal transduction [55]. Previous studies showed that miR156 regulated the development stage transition [56], flowering process [57], fertility maintenance [58], and fruit ripening [59] in plants by regulating members of the *SPL* family. Gma-circRNA0888, gma-circRNA1395, and gma-circRNA1746 could all target gma-miR156aa and gma-miR156z, which are members of miR156 family and probably involved in flower development. Furthermore, gma-circRNA1001 was predicted to target gma-miR162a, gma-miR162b, and gma-miR162c. NADP-dependent isocitrate dehydrogenase (*NADP-ICDH*) was targeted by gma-miR162a, gma-miR162b, and gma-miR162c according to the degradome analysis performed prior by NJCMS1A and NJCMS1B [11].

| Table 2 Twenty fertility-related parental genes of differentially expressed circRNAs |
|-----------------|-----------------|-----------------|-----------------|
| circRNAs ID | Parental genes | Homologue Genes (Arabidopsis thaliana) | Function description |
| gma-circRNA0457 | Glyma.04G155600 | AT2G24120 | male gametophyte development |
| gma-circRNA0717 | Glyma.06G173500 | AT2G36850 | male gametophyte development; callose component |
| gma-circRNA1324 | Glyma.09G266600 | AT3G61050 | male gametophyte development; pollen tube growth |
| gma-circRNA2270 | Glyma.16G218700 | AT5G22110 | male gametophyte development; embryo development |
| gma-circRNA2801 | Glyma.20G205100 | AT1G06750 | male gametophyte development; pollen tube growth |
| gma-circRNA0057 | Glyma.01G140600 | AT1G71820 | pollen germination; pollen tube growth |
| gma-circRNA0685 | Glyma.06G135600 | AT3G03810 | pollen tube development; chloroplast component |
| gma-circRNA0914 | Glyma.07G002300 | AT1G78900 | pollen development |
| gma-circRNA1244 | Glyma.09G103400 | AT2G37270 | pollen tube development |
| gma-circRNA1301 | Glyma.09G194900 | AT3G65930 | pollen germination; plasma membrane component |
| gma-circRNA1432 | Glyma.10G217200 | AT1G76850 | pollen germination; pollen tube growth |
| gma-circRNA1856 | Glyma.13G230500 | AT2G34090 | pollen gametogenesis; pollen germination; pollen tube-embryo sac interactions |
| gma-circRNA2481 | Glyma.18G164900 | AT5G48720 | pollen development; DNA repair; male meiosis |
| gma-circRNA2791 | Glyma.20G182800 | AT3G10380 | pollen germination; pollen tube growth |
| gma-circRNA1934 | Glyma.14G119800 | AT1G43850 | flower development; embryo development |
| gma-circRNA2102 | Glyma.15G000300 | AT1G16710 | flower development; protein amino acid acetylation |
| gma-circRNA2454 | Glyma.18G118100 | AT1G58440 | flower development; endomembrane system component |
| gma-circRNA2637 | Glyma.19G212600 | AT4G02300 | flower development; cell wall component |
| gma-circRNA0464 | Glyma.04G163400 | AT5G61150 | negative regulation of flower development |
| gma-circRNA2163 | Glyma.15G265300 | AT1G79280 | negative regulation of flower development; stamen development |

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may result in significant ROS accumulation and male sterility in soybean. Moreover, gma-miR169e was targeted by gma-circRNA1393, and gma-miR169e could target dihydrolipoyl dehydrogenase ($E_3$), an important part of the pyruvate dehydrogenase complex. Pyruvate dehydrogenase complex is a critical pathway that supports energy generation for pollen and pollen tube growth [62]. In our previous study, qRT-PCR analysis showed that the target gene $E_3$ was down-regulated in NJCMS1A [11]. In addition, gma-circRNA0195 was predicted to target gam-miR172j, which could target Glyma.01 g188400, the homolog of the APETALA2 (AP2) of A. thaliana. In A. thaliana, miR172 could control flowering time and floral organ formation by regulating expression of the AP2-like transcription factor. A previous study indicated that over-expression of miR172 caused flower development abnormalities, and its phenotype was similar to ap2 mutants, leading to abnormal gametophyte development [63].
Functional annotation of predicted target mRNAs

To further explore the role of circRNAs in CMS in soybean, the 83 binding miRNAs of differentially expressed circRNAs were used to predict their possible target mRNAs by psRobot with the default parameters. In total, 1166 target mRNAs were predicted and further used for functional analysis (Additional file 8). A direct acyclic graph (DAG) of biological processes was obtained using the agriGO online server (Fig. 7). The DAG can indicate submitted terms and the inter-relationships between terms. From this figure, we could identify several pathways with significant enrichment, especially those involved in signal transduction and programmed cell death (PCD). A previous study showed that pollen development depended on the interaction of multiple signaling pathways, in which calmodulin was a key element [64]. Disruption of signaling pathways might cause abnormal pollen development, resulting in male sterility in soybean. In addition, PCD in plants is a cellular process similar to apoptosis, which contains fragmentation of nuclear DNA and is controlled by mitochondrion-driven signals [3]. Plant PCD plays a role in development processes such as senescence, seed germination, organ development, root tip elongation, xylem and aerenchyma formation, and disease resistance [65]. The development of plant male gametophytes in anthers requires cooperative interactions between sporophytic (anther wall) and gametophytic (microspore) cells as well as proper PCD-controlled cellular degeneration of the tapetum and the anther wall tissue [66]. Therefore,
premature or delayed PCD leads to abnormality of pollen development and tapetal function, and even male sterility [67–69]. For example, the PET1-CMS cytoplasm in sunflower causes premature PCD of the tapetal cells, which then leads to abnormal anther development [70]. The KEGG pathway analysis was also used to further explore the function of predicted target mRNAs of differentially expressed circRNAs. The target mRNAs were significantly enriched in six pathways: beta-Alanine metabolism, fatty acid degradation, lysine degradation, ascorbate and aldarate metabolism, limonene and pinine degradation, and mRNA surveillance pathway (Table 3). Several studies have shown that in most plant species, the contents of amino acids, proteins, and soluble sugars in male sterile lines were lower than those in their maintainer lines [71, 72]. In this study, the pathways of amino acid degradation and material metabolism may be important causes of CMS in soybean. GO categories and KEGG pathway analyses showed that a large number of target mRNAs were involved in pollen development and male fertility, which implied that the differentially expressed circRNAs might play an important role in CMS of soybean.

**Prediction of protein-coding potential of soybean circRNAs**
Recent studies have demonstrated that some circRNAs can be translated into polypeptides or proteins by translation initiation element internal ribosome entry site (IRES) [27]. To verify whether soybean circRNAs have a similar function, we predicted the protein-coding potential by blasting sequences of the circRNAs to all the IRES sequences at the website. A total of 165 soybean circRNAs contained at least one IRES element and an open reading frame (Additional file 9), which might have protein-coding potential. Furthermore, conserved domains of the possible protein-coding products were predicted by Conserved Domain Database (Additional file 10) [73], which might have important functions. Recent studies have shown that the protein-coding products of circRNAs can influence the function of their parental genes by interacting directly or indirectly with them [74, 75].

In this study, some parental genes of circRNAs with protein-coding potential were associated with flower and pollen development. For example, gma-circRNA0736 was predicted to encode an 81-amino acid protein, and its parental gene was a homolog of VERNALIZATION INDEPENDENCE 4 (VIP4) in A. thaliana. The VIP4 mutants could cause slightly early flowering and variable fertility under standard growth conditions [76]. Gma-circRNA1793 was predicted to encode a 154-amino acid protein, and its parental gene was a homolog of no pollen germination related 2 (NPGR2) in A. thaliana. The NPGR2 encodes a calmodulin-binding protein that is essential for pollen germination [77]. The protein-coding products of these circRNAs may affect flower and pollen development in soybean by affecting the function of the parental genes.

**Conclusion**
In this study, a total 2867 circRNAs, of which 1009 were differentially expressed between the soybean CMS line NJCMS1A and its maintainer NJCMS1B, were identified by high-throughput deep sequencing. Tissue specific expression patterns were verified by quantitative real-time PCR. The parental genes of differentially expressed circRNAs were mainly enriched in biological processes such as metabolic process, biological regulation, and reproductive process. A large number of parental genes were related to flower development and male fertility. A total of 83 binding miRNAs were predicted among the differentially expressed circRNAs, which included well-known flower and pollen development-related miRNAs. The target mRNAs predicted for the 83 binding miRNAs were significantly enriched in signal transduction and programmed cell death. A total of 165 soybean circRNAs contained at least one IRES element and an open reading frame (ORF), indicating their potential to encode polypeptides or proteins. Our study indicated that circRNAs might participate in the regulation of flower and pollen development, which could provide a new insight into the molecular mechanisms of CMS in soybean.

### Table 3 KEGG pathway enrichment of target mRNAs of differentially expressed circRNAs in soybean

| Pathway ID | Description                           | Gene count | P-value |
|------------|---------------------------------------|------------|---------|
| gmx00410   | beta-Alanine metabolism               | 9          | 0.004   |
| gmx00071   | Fatty acid degradation                | 7          | 0.027   |
| gmx00310   | Lysine degradation                    | 5          | 0.030   |
| gmx00053   | Ascorbate and aldarate metabolism     | 6          | 0.032   |
| gmx00903   | Limonene and pinine degradation       | 3          | 0.034   |
| gmx03015   | mRNA surveillance pathway             | 12         | 0.045   |

Pathway with the threshold of P-value < 0.05 was listed.
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Availability of data and materials

The sequencing data have been deposited in the Sequence Read Archive (SRA) at the National Center for Biotechnology Information (NCBI) under accession number SRP160000.

Authors’ contributions

JG, SY, and LC conceived and designed the experiments. LC performed the experiments and analyzed the data. LC, XD, HZ, TH, YL, TW, XL, and LJ contributed reagents, materials, and analysis tools. LC and SY wrote the paper. JG, SY, QS, and LC revised the paper. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The plant materials were collected from germplasms bank of National Center for Soybean Improvement. The collection and usage of samples followed the ethics of the People’s Republic of China.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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