Transcriptomic Analysis for Different Sex Types of *Ricinus communis* L. during Development from Apical Buds to Inflorescences by Digital Gene Expression Profiling

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The castor plant (*Ricinus communis* L.) is a versatile industrial oilseed crop with a diversity of sex patterns, its hybrid breeding for improving yield and high purity is still hampered by genetic instability of female and poor knowledge of sex expression mechanisms. To obtain some hints involved in sex expression and provide the basis for further insight into the molecular mechanisms of castor plant sex determination, we performed DGE analysis to investigate differences between the transcriptomes of apices and racemes derived from female (JXBM0705P) and monoecious (JXBM0705M) lines. A total of 18 DGE libraries were constructed from the apices and racemes of a wild monoecious line and its isogenic female derivative at three stages of apex development, in triplicate. Approximately 5.7 million clean tags per library were generated and mapped to the reference castor genome. Transcriptomic analysis showed that identical dynamic changes of gene expression were indicated in monoecious and female apical bud during its development from vegetation to reproduction, with more genes expressed at the raceme formation and infant raceme stages compare to the early leaf bud stage. More than 3000 of differentially expressed genes (DEGs) were detected in *Ricinus* apices at three developmental stages between two different sex types. A number of DEGs involved in hormone response and biosynthesis, such as auxin response and transport, transcription factors, signal transduction, histone demethylation/methylation, programmed cell death, and pollination, putatively associated with sex expression and reproduction were discovered, and the selected DEGs showed consistent expression between qRT-PCR validation and the DGE patterns. Most of those DEGs were suppressed at the early leaf stage in buds of the mutant, but then activated at the following transition stage (5-7-leaf stage) of buds in the mutant, and ultimately, the number of up-regulated DEGs was equal to that of down-regulation in the small raceme of the mutant. In this study, a large number of DEGs and some suggestions involved in sex expression and reproduction were discovered using DGE analysis, which provides large information and valuable hints for next insights into the molecular mechanism of sex determination. It is useful for other further studies in *Ricinus*. 

**Keywords:** castor bean, digital expression profile, pistillate, sex determination, transcriptome analysis
INTRODUCTION

Castor (*Ricinus communis* L.), an important oilseed crop belonging to the Euphorbiaceae family, grows in an indeterminate annual or perennial depending on climate and soil types in tropical, sub-tropical, and warm temperate regions (Anjani, 2012). Because of the high fatty acid content in its seeds (more than 45%) and the rich ricinoleic acid content in its oil (80–90%), castor is a versatile raw material in industrial chemistry: for example, castor oil can be used for the production of lubricants, nylon, hydraulic and brake fluids, paints, dyes, coatings, inks, cold resistant plastics, waxes and polishes, pharmaceuticals, perfumes, and biodiesel (Jeong and Park, 2009; Halilu et al., 2013). To date, several high-yield varieties and hybrids have been developed (Baldanzi and Pugliesi, 1998; Amaral, 2003; Savy Filho, 2005; Pranavi et al., 2011), but to meet the tremendous global demand for castor oil, cultivars with even higher yield and oil content are needed (Anjani, 2012). However, hybrid breeding for improved yield and high purity in *Ricinus* is still constrained by the genetic instability of females and the unknown mechanism of sex expression.

In castor plant, the standard type of inflorescence is gradient monoeccious raceme (female flowers at the apex and staminate flowers on the lower portion) (Shifriss, 1960). However, a wide variation of inflorescence patterns occurs in natural cultivation, including other kinds of racemes such as strictly pistillate (bearing only female flowers), male (only staminate flowers), apically interspersed (monoeccious with interspersed male flowers in the apical pistillate region), and entirely interspersed (female and male flowers uniformly interspersed) (George and Shifriss, 1967). In addition, inflorescence setting with one or a few hermaphrodite flowers occurs occasionally (Jacob, 1963). Accordingly, for castor individuals, there are several sex models, including normal monoeccious, sex reversal, interspersed sexuality, and strictly female (Shifriss, 1956; Jacob and Atsmon, 1965; George and Shifriss, 1967).

In the past, scientists focused on the sexuality of *Ricinus* to interpret the inheritance and instability of sex variation. Shifriss (1956) described the sex tendency, sex patterns, inheritance and reversion of *Ricinus*, and proposed a hypothesis: Gene *F* controls a genetically stable series of sex variants ranging from female (*f*) to strongly male inbreds, and an unknown factor could affect male tendency, sex reversion, and sex instability by suppressing gene *F* or mutating itself. Later, he described sex variations of *Ricinus* in two genetic systems, which he tentatively named as “conventional” and “unconventional” (Shifriss, 1960). Monoeccious variants and rare recessive female mutants were ascribed to the conventional form, sex reversals and non-reverted females belonged to the unconventional form. Monoeccism is governed by two major groups of genes: qualitative genes that determine flower type, and polygenes regulating gradient differentiation and racial differences in sex tendency. In addition, gene modifiers such as *id* and *th* were also considered to affect the pattern of sex differentiation. He postulated that genetic instability of spontaneous mutants, so-called position-effect variegation, may result from a rearrangement undergoing two basic kinds of genetic changes: mutation into new hereditary potentialities, and transformation between an “active” and “inactive” state. Moreover, further evidence regarding the former findings (Shifriss, 1956, 1960) revealed that femaleness transmits more effectively to progeny through female inflorescences of sex-reversal plants than through reverted monoeccious inflorescences of the same plant (Jacob and Atsmon, 1965). With regard to interspersed sexuality, Shifriss (1956, 1960) believed that the interspersed pattern of sex differentiation is determined by hereditary factors for femaleness and genes for interspersed staminate flowers (*id*). George and Shifriss (1967) investigated more deeply into the inheritance of interspersed inflorescence patterns, and concluded that the level of expressivity of interspersed staminate flowers depends on the dosage of two independent genes (*id1* and *id2*), their loci, and the environment.

In addition to genetic regulation, sex expression or variation of *Ricinus* is simultaneously affected to some degree by fluctuations in the environment, including temperature, vegetative activity, nutrition level, pruning, and seasonal variations (Shifriss, 1956), as well as by plant hormones (Shifriss, 1961; Philapos and Narayanawamy, 1976; Kumar and Rao, 1980; Mohan Ram and Sett, 1980; Varkey and Nigam, 1982; Tan et al., 2011). Despite a great deal of progress, the hypotheses described above are vulnerable in the absence of cytological information, and the molecular mechanisms of sex variation and genes determining sex expression in *Ricinus* remain poorly understood.

Genome-wide analyses have dramatically improved the efficiency of gene discovery. Next-generation sequencing (NGS) technologies provide new approaches for global measurements of gene expression. Due to its high efficiency and low cost, NGS has become an attractive alternative method for more efficient study of the genome, epigenome, and transcriptome (Oshlack et al., 2010; McIntyre et al., 2011). Many plant species have benefited from this technology, and large-scale genome sequences and transcriptome data are available in both model and non-model species (Goff et al., 2002; Yu et al., 2002; Kaplan et al., 2006; Ramsey et al., 2008; Huang et al., 2009; Wang et al., 2011; Xu et al., 2013). Cucumber, as a model plant for study of floral sex expression, its sex-controlling genes, sex-modifying plant hormones, and interactions with environmental conditions are clearly understood (Galun, 1962; Trebitsh et al., 1987, 1997; Malepszy and Niemirowicz-Szczytt, 1991; Yin and Quinn, 1995; Yamasaki et al., 2000; Kater et al., 2001; Mibus and Tatlioglu, 2004; Li et al., 2009b), and a number of candidate genes required for sex determination have been identified by transcriptome profile analysis (Guo et al., 2010; Wu et al., 2010). In *Ricinus*, however, a gap still exists between interpreting the molecular mechanism of sex expression and isolating the sex-determining genes. The complete sequencing of the castor bean genome (Chen et al., 2010) provides tremendous opportunities for genomic analysis and facilitates identification of genes of biological interest. Digital gene expression (DGE) analysis, a powerful and recently developed tool, is based on ultra-high-throughput sequencing of millions of signatures in the genome, allowing identification of specific genes and direct quantitation of transcript abundance (Bentley, 2006; Hong et al., 2011). DGE analysis detects gene expression more quantitatively than microarray assays (‘t Hoen et al., 2008) while providing similar
assessments of relative transcript abundance. In comparison to high-throughput mRNA sequencing (RNA-seq), DGE more accurately detects expression differences in poorly expressed genes and exhibits much less transcript length bias (Hong et al., 2011). Thus, DGE has been widely utilized to discriminate differences in transcriptional responses among different tissues and organs in plants in the contexts of biotic and abiotic stress, metabolite biosynthesis, and developmental biology (Tian et al., 2013; Wei et al., 2013; Zhang et al., 2013; Guo et al., 2014; Yu et al., 2014; Zhao et al., 2014).

In this study, we performed DGE using the Illumina HiSeq 2000 to investigate the differences between the transcriptomes of apices and racemes from female (JXBM0705P) and monoecism (JXBM0705M) of Ricinus. In an attempt to get some hints associated with sex expression in Ricinus, we conducted (after high-throughput tag sequencing) an integrated bioinformatic analysis to identify expression patterns of genes and critical pathways in the female and monoecious lines at three stages of apex development. Assessment of the changes in gene expression between female and monoecious apices yielded sets of up-regulated and down-regulated genes associated with sex expression. Based on these differentially regulated genes (DEGs), some DEGs putatively related to sex expression were selected to confirm by quantitative RT-PCR analysis. Thus, comparison of gene expression patterns at three developmental stages of monoecious and female apices provides important hints for further insight into the molecular mechanisms underlying Ricinus sex variation.

MATERIALS AND METHODS

Plant Material

Seeds of nearly isogenic monoecious (JXBM0705M) and female (JXBM0705P) castor lines were kindly provided by Professor Huang (Castor Oil Research Institute of Jiaxiang, Zibo, Shandong Province). JXBM0705M is a wild monoecious line that bears male and female flowers (Figure 1A), and JXBM0705P, a pistillate line that bears only female flowers (Figure 1B), was developed by consecutively selecting spontaneous mutants of JXBM0705M. Seeds were sown and cultivated in the Yangluo experimental field of the Oil Crops Research Institute of the Chinese Academy of Agricultural Sciences, under standard field conditions, from spring to autumn of 2012. A previous study revealed that apices are leaf buds prior to the 5-leaf stage (i.e., five fully expanded leaves) and change to floral buds during the 6–9-leaf stage (Wang et al., 2012). According to this criterion, and combined with anatomical observation of a series of apical buds of the cultivar, in order to investigate global changes in the transcriptome during the development of buds from leaf apices to racemes, apical buds at the 3–4-leaf stage (ABML1 for monoecism and ABPL1 for female) and the 5–7-leaf stage (ABML2 for monoecism and ABPL2 for female) (Figure 1C), and small racemes 2–3 cm in size (RML for monoecism and RPL for female, Figure 1D), were collected in triplicate to construct 18 DGE libraries. Excised apices and infant racemes were immediately frozen in liquid nitrogen and stored at −80°C until use.

RNA Preparation, Library Construction, and Sequencing

Total RNA was extracted from buds and racemes using the GeneJET Plant RNA Purification Mini Kit (Thermo Fisher Scientific, Waltham, USA). RNA was dissolved in diethylpyrocarbonate (DEPC)-treated water and stored at −70°C. The quality of total RNA was checked on 1% agarose gels, and purity was evaluated by OD260/280 ratio using a NanoDrop ND 1000 spectrophotometer (NanoDrop, Wilmington, DE, USA). Concentration and integrity of RNA were verified using an Agilent 2100 Bioanalyzer. The extracted RNA was prepared for next sequencing and quantitative RT-PCR validation.

A total of 8 μg of total RNA was subjected to oligo (dT) magnetic bead adsorption to purify mRNA. Oligo (dT) was then used as a primer for the synthesis of first- and second-strand cDNA. The 5’-ends of tags could be generated using two types of endonucleases, namely, NlaIII or DpnII. Bead-bound cDNA was subsequently digested with the restriction enzyme NlaIII, which recognizes and cleaves CATG sites. Fragments other than 3’-cDNA fragments connected to oligo (dT) beads were washed away, and Illumina adaptor 1 was ligated to the sticky 5’-end of the digested bead-bound cDNA fragments. The junction of the Illumina adaptor 1 and the CATG site constitutes the recognition site for MmeI, an endonuclease with separate recognition and cleavage sites. MmeI cleaves 17 bp downstream of the CATG site, yielding tags with adaptor 1. After removing 3’ fragments by magnetic bead precipitation, adaptor 2 was ligated to the 3’-ends of tags, yielding tags with different adaptors at both ends.
to form a tag library. After linear PCR amplification, fragments were purified by PAGE gel electrophoresis. During the quality control steps, the Agilent 2100 Bioanalyzer and ABI Step One Plus Real-Time PCR System were used for quantitation and quality verification of the sample library. Finally, the library was sequenced on an Illumina HiSeq 2000.

Data Analysis and Mapping of DGE Tags
Millions of raw 49 bp sequences were generated. Image analysis, base calling, generation of raw tags, and tag counting were performed using the Illumina pipeline (t’Hoen et al., 2008). Prior to mapping of tags to the reference database, empty tags (i.e., those with no tag sequence between the adaptors), adaptors, low-quality tags (tags containing one or more unknown nucleotides “N”), and tags with a copy number of 1 were removed from raw sequences to obtain clean tags.

To evaluate the normality of the entire DGE dataset, the distribution of clean tag expression was analyzed. The saturation analysis of the 18 libraries was also performed to estimate whether sequencing depth was sufficient for transcriptome coverage. To identify the gene expression patterns in apical buds and small racemes of two Ricinus lines, all clean tags were annotated by mapping to the castor genome (Chan et al., 2010) using the SOAP2 software (Li et al., 2009a), with a maximum of one nucleotide mismatch allowed. All tags mapped to reference sequences were filtered, and the remaining tags were designated as ambiguous tags. Mapping events on both sense and antisense sequences were included in data processing. For gene expression analysis, in order to identify genes that were differentially expressed between monoeocious and female apices at the same developmental stages, as well as genes that exhibited distinctive expression between different developmental stages in the same line, we converted the number of raw clean tags in each library to TPM (number of transcripts per million tags) (t’Hoen et al., 2008; Morrissy et al., 2009) to obtain normalized gene expression levels. Then, nine pairs of DGE profiles of different sample libraries (ABML1 vs. ABPL1, ABML2 vs. ABPL2, RML vs. RPL, ABML1 vs. ABML2, ABML2 vs. RML, ABML1 vs. RML, ABPL1 vs. ABPL2, ABPL2 vs. RPL, and ABPL1 vs. RPL, where a is the control and b is the experimental group in “a vs. b”) were compared to assess the diversity of gene expression. A rigorous algorithm for identifying DEGs (differentially expressed genes) between two samples, false discovery rate (FDR), was applied to determine the threshold P-value in multiple tests and analyses. Criteria for DEGs were as follows: FDR, ≤0.001, and [log2Ratio], ≥1 (Audic and Claverie, 1997; Benjamini and Yekutieli, 2001). Following, we performed functional annotation to assign the unambiguously mapped genes identified in all libraries to Gene ontology (GO) terms. Gene annotation was conducted using Blast2GO (Conesa et al., 2005). Gene ontology (GO) was used to determine the possible functions of all DEGs by searching the GO database (http://www.geneontology.org/), and Web Gene Ontology Annotation Plot (WEGO) was also used for GO classification of genes identified in each DGE library (Ye et al., 2006). Moreover, the GO distribution of DEGs in comparisons of each pair of libraries was determined and compared. To further characterize gene function, pathway enrichment analysis of the DGE results was conducted by BLAST search of the KEGG database (http://www.kegg.jp/kegg/). Clustering analysis of differential gene expression patterns was also performed using MultiExperiment Viewer (MeV) (Chu et al., 2008; Howe et al., 2011). A P-value of 0.05 was selected as the threshold for a gene set to be considered significantly enriched.

Quantitative RT-PCR Analysis
Quantitative RT-PCR analysis was used to verify the DGE results. The RNA samples used for the qRT-PCR assays were identical to those used for the DGE experiments. Three biological replicates and two technical replicates were performed for each sample. The first-strand cDNA was synthesized from the total RNA (1 ug) from each sample using the PrimeScript RT reagent kit with gDNA Eraser (perfect Real Time) (TaKaRa, RR047A). The primers designed for qPCR analysis was listed in Additional File S1. Actin gene (forward primer: 5'-TGCTGACAGAATGAGCAAG-3'; reverse primer: 5'-AATCCACATCGCTGGAAGG-3') was used as an internal control gene for normalization. Quantitative PCR reactions used ABI7500 quantitative PCR, which was performed using SYBR® Premix Ex Taq™II(Tli RNaseH Plus) (TaKaRa, RR820A) kit according to the manufacturer’s instructions. Reaction conditions were 95°C for 5 s in order to activate enzyme reaction. Two step cycles were then used: 95°C for 5 s, then 60°C for 30 s, 40 cycles; solubility curve conditions were 95°C for 15 s, 60°C for 1 min, 95°C for 30 s, 60°C for 15 s. The specificity of the SYBR green PCR signal was further confirmed by melting curve analysis and agarose gel electrophoresis, and the relative expression levels of genes were calculated with the 2^ΔΔCt method.

Hormone Measurements
Contents of several phytohormones including auxin or indole-3-acetic acid (IAA, the major form of auxin in plants), abscisic acid (ABA), jasmonic acid (JA), and gibberellins (GAs) were analyzed to compare and verify the physiological differences between the monoeocious and the pistillate. The samples used for the hormone measurement were the same as those used for the DGE experiments (Three development stages of apices from leaf bud to raceme for the monoeism and female, respectively; ABML1, ABML2, and RML for monoeicism, ABPL1, ABPL2, and RPL for female.) Three biological replicates were designed for each sample. Quantification of endogenous IAA, ABA, JA, and GAs (GA1, GA3, GA4, and GA7) were performed as described previously (Chen et al., 2012). The data of detection was analyzed using Microsoft Excel 2010 and SAS V8 softwares.

RESULTS
Analysis of DGE Libraries and Tag Mapping
The 18 DGE libraries were sequenced and generated approximately 6 million raw tags for each library, and more than 96% of the raw tags were clean (Table 1). There produced approximately 5.7 million clean tags per library, of which 148,526–174,340 were distinct. The number of unique distinct clean tag sequences ranged from 134,541 to 159,229 (Table 1).
# Table 1: Categorization and abundance of tags.

| Summary                              | ABML1          | ABPL1          | ABML2          | ABPL2          | RML           | RPL           |
|--------------------------------------|----------------|----------------|----------------|----------------|---------------|---------------|
| Raw data                             | 5,919 ± 211,429 | 5,888 ± 122,404 | 6,004 ± 92,543 | 6,000 ± 135,373 | 5,930 ± 123,287 | 6,045 ± 233,096 |
| Distinct tags                        | 371 ± 4765     | 374 ± 40,008    | 391 ± 12,149   | 440 ± 11,500   | 369 ± 807     | 447 ± 47,302  |
| Clean tags                           | 5,697 ± 209,466 | 5,661 ± 132,174 | 5,767 ± 88,697 | 5,729 ± 121,449 | 5,722 ± 123,544 | 5,771 ± 203,535 |
| Total number                         | 5,199 ± 567    | 5,170 ± 650    | 5,269 ± 111    | 5,232 ± 678    | 5,233 ± 678   | 5,287 ± 678   |
| Distinct tag numbers                 | 148 ± 12,470   | 155 ± 6806     | 170 ± 3920     | 172 ± 5900     | 173 ± 4430    | 174 ± 340,17   |
| All tags mapping to genes            | 5,696 ± 967,781 | 5,476 ± 137,781 | 5,477 ± 134,778 | 5,490 ± 156,338 | 5,516 ± 151,205 |                |
| Total % of clean tags                | 94.66 ± 0.83   | 95.35 ± 1.86   | 94.98 ± 1.72   | 95.6 ± 0.58    | 95.92 ± 0.82  | 95.62 ± 1.79  |
| Distinct tag numbers                 | 135 ± 3278     | 134 ± 8871     | 139 ± 5476     | 154 ± 2420     | 141 ± 2952    | 159 ± 9689    |
| Distinct tags (% of clean tags)      | 90.02 ± 0.98   | 90.69 ± 1.97   | 89.91 ± 1.71   | 90.65 ± 1.4    | 91.60 ± 1.2   | 91.4 ± 1.89   |
| Unambiguous tag mapping to gene      | 168 ± 7762     | 162 ± 5683     | 163 ± 16,927   | 169 ± 5762     | 172 ± 6399    | 173 ± 11,236  |
| Total number                         | 2.96 ± 0.07    | 2.86 ± 0.21    | 2.83 ± 0.26    | 2.96 ± 0.16    | 3.01 ± 0.09   | 3.01 ± 0.1    |
| Distinct tag numbers                 | 7665 ± 306     | 7323 ± 585     | 8164 ± 599     | 8722 ± 426     | 8223 ± 295    | 9013 ± 514    |
| Distinct tags (% of clean tags)      | 5.08 ± 0.15    | 4.94 ± 0.31    | 5.26 ± 0.2     | 5.11 ± 0.15    | 5.34 ± 0.04   | 5.18 ± 0.13   |
| All tag-mapped genes                 | 50 ± 577       | 38 ± 1085      | 45 ± 4807      | 54 ± 1599      | 44 ± 1911     | 41 ± 6629     |
| % of ref genes                       | 41.99 ± 0.48   | 32.17 ± 0.9    | 37.57 ± 3.98   | 44.78 ± 0.96   | 37.01 ± 1.59  | 34.48 ± 0.52  |
| Unambiguous tag-mapped genes         | 4867 ± 191     | 4582 ± 227     | 5050 ± 333     | 5215 ± 197     | 5144 ± 137    | 5292 ± 144    |
| % of ref genes                       | 4.03 ± 0.16    | 3.79 ± 0.19    | 4.18 ± 0.28    | 4.32 ± 0.16    | 4.26 ± 0.12   | 4.38 ± 0.12   |
| Mapping to genome                    | 31 ± 3790      | 30 ± 8907      | 33 ± 1786      | 27 ± 2213      | 26 ± 2537     | 30 ± 6426     |
| Total % of clean tags                | 0.55 ± 0.07    | 0.54 ± 0.15    | 0.57 ± 0.12    | 0.48 ± 0.04    | 0.47 ± 0.05   | 0.52 ± 0.1    |
| Distinct tag numbers                 | 1494 ± 115     | 1607 ± 173     | 1879 ± 390     | 1715 ± 412     | 1230 ± 113    | 1548 ± 187    |
| Distinct tags (% of clean tags)      | 0.99 ± 0.07    | 1.08 ± 0.12    | 1.21 ± 0.2     | 1.22 ± 0.05    | 0.85 ± 0.05   | 0.89 ± 0.05   |
| Unknown tags                         | 272 ± 43,697   | 233 ± 100,607  | 257 ± 93,700   | 224 ± 28,803   | 206 ± 41,217  | 223 ± 104,814 |
| Total % of clean tags                | 4.79 ± 0.76    | 4.11 ± 1.7     | 4.45 ± 1.59    | 3.92 ± 0.55    | 3.61 ± 0.77   | 3.85 ± 1.7    |
| Distinct tag numbers                 | 13,565 ± 1338  | 12,378 ± 2813  | 13,810 ± 2853  | 14,271 ± 2285  | 11,541 ± 1925 | 13,563 ± 4350 |
| Distinct tags (% of clean tags)      | 8.99 ± 0.91    | 8.23 ± 1.98    | 8.89 ± 1.67    | 8.35 ± 1.18    | 7.48 ± 1.08   | 7.69 ± 1.85   |
The RPL libraries and ABPL2 libraries contained the two highest numbers of distinct clean tags; the other libraries had similar numbers.

Our results showed that mRNAs transcribed from major genes were often present at fewer than 10 copies, and only a small proportion of genes were expressed at higher levels (Figure 2; Additional Files S2, S3). The distributions of total and distinct clean tag copy numbers had highly similar properties in libraries from all six sample types, with most tags coming from highly expressed genes (Figure 2). Among the distinct clean tags, more than 56% were present at 2–5 copies, 33.81–36.84% were present at 5–100 copies, and fewer than 6.27% had copy numbers higher than 100. However, more than 70% of total clean tags had counts above 100 in each library, indicating that the overall DGE data among the 18 libraries was normally distributed.

A reference gene database that included 120,799 R. communis Unigene sequences was preprocessed for tag mapping. Among the sequences, genes with a CATG site accounted for 90.03% (Additional File S3). Distinct clean tags (4.94–5.34%) were mapped unambiguously in the unigene database, whereas 7.48–8.99% of distinct clean tags were not mapped to the unigene virtual tag database, in libraries from the six sample types (Table 1). Around 95% of total clean tags were mapped onto the R. communis genome with a perfect match or 1 bp mismatch to sense or antisense genes, and approximately 90% of distinct clean tags were successfully mapped. Ultimately, tag mapping onto the R. communis genome generated 50,722 tag-mapped genes (Table 1; Additional File S4).

In this research, the number of detected genes increased with the amount of sequencing amount until the number of tags reached 3 million or higher (Additional File S5). Distinct clean tags (4.94–5.34%) were mapped unambiguously in the unigene database, whereas 7.48–8.99% of distinct clean tags were not mapped to the unigene virtual tag database, in libraries from the six sample types (Table 1). Around 95% of total clean tags were mapped onto the R. communis genome with a perfect match or 1 bp mismatch to sense or antisense genes, and approximately 90% of distinct clean tags were successfully mapped. Ultimately, tag mapping onto the R. communis genome generated 50,722 tag-mapped genes for ABML1, 38,867 for ABPL1, 45,392 for ABML2, 54,090 for ABPL2, 44,709 for RML, and 41,658 for RPL (Table 1; Additional File S4).

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In total, more than 88% of genes were transcribed from both strands, indicating the importance of RNA-mediated gene regulation in bud transformation from leaf to inflorescence.

### Global Gene Expression of Buds at Different Stages and Analysis of Differential Gene Expression

In this study, we identified 5526 and 5203 genes in the transcriptomes of monoecious and pistillate apical buds, respectively, at the 3–4-leaf stage; 5779 and 6118 genes at the 5–7-leaf stage; and 5870 and 5988 genes in racemes (Figure 3A). Our results showed that whether monoecism or female, the transcriptome of their terminal buds during the three developmental stages was dynamic. In total, 7495 and 7610 genes were expressed in monoecious and pistillate apical buds over the three stages, respectively. Of which, 4066 and 4013 were constitutive, 1881 and 1924 were specific to a single stage, and 1548 and 1637 were expressed at two stages, correspondingly for monoecism and female (Figures 3B,C; Additional File S6).

The complex changes of gene expression indicated that the transition from leaf apical bud to raceme is an involute process in castor bean. More genes were expressed in the buds closer to raceme formation and incipient racemes than in early apical buds (Figure 3), and more stage-specific transcripts were detected in the 5–7-leaf stage bud and the small raceme than in the early leaf bud (Figure 3D; Additional File S6). These observations suggest that identical molecular pathways are involved in apical bud development from vegetation to reproduction in the monoecism and female, the developmental process may require expression of a larger number of unique genes involved in regulatory processes and related pathways.

Comparison of the monoecious line with the female line revealed that 1386 DEGs were up-regulated and 1946 DEGs were down-regulated at the 3–4-leaf stage, but the majority of genes were up-regulated during the subsequent two stages: 1952 up-regulated and 1408 down-regulated genes at the 5–7 leaf stage, and 1667 up-regulated and 1451 down-regulated genes at the initial raceme stage (Table 2; Figures 4A–C; Additional File S7). The number of up-regulated DEGs in the female was greater than the number of down-regulated DEGs during the developmental transition from leaf bud to infant raceme, which might indicate that females require a number of genes involved in silencing of male flower expression. Similarly, it was showed that up-regulated genes outnumbered down-regulated genes in these comparisons between different development stages, except for the RPL vs. ABPL2 (Table 2; Figures 4D–I; Additional File S7), which also revealed that the expression of many genes tended to increase; a few genes were activated, whereas others were suppressed, during bud development from vegetation to reproduction.

### Functional Annotation and DEG Clustering Analysis

GO analysis revealed that these well-annotated sequences belonged to three main categories (cellular component, molecular function, and biological process) and mainly...
FIGURE 3 | Transcriptome analysis of apical buds and racemes of the monoecious line (ML) and pistillate line (PL). (A) Transcriptome sizes of the monoecious and pistillate lines at three developmental stages. (B,C) Venn diagram showing the overlaps between three stages (two apical bud stages and raceme stage) of ML and PL. The number in parentheses after each stage designation is the total transcripts detected in that stage(s). (D) Analysis of transcriptome changes from apical buds to raceme development of the ML and PL. Transcripts shared by three stages of ML and PL are not shown; numbers above the x-axis represent transcripts present in the indicated stage that are stage-specific (dark red), not shared with the previous stage (orange), or shared with the prior stage but missing in at least one other stage (yellow-green). Numbers below the x-axis represent transcripts present in the prior stage that were not detected in the current stage (green and blue).

TABLE 2 | Gene expression levels across different sample libraries.

|                  | ABML1/ RML | ABML2/ RML | ABML1/ RML | ABPL1/ RPL | ABPL2/ RPL | ABPL1/ RPL | ABML1/ ABPL1 | ABML2/ RPL | RML/ RPL |
|------------------|------------|------------|------------|-----------|------------|------------|-------------|------------|---------|
| Significantly up-regulated | 17         | 48         | 68         | 32        | 75         | 103        | 2           | 15         | 6       |
| Up-regulated      | 1873       | 1991       | 2083       | 2613      | 2055       | 2535       | 1384        | 1937       | 1661    |
| Not DEGs          | 3372       | 3184       | 3207       | 3043      | 2940       | 2863       | 3089        | 3664       | 3855    |
| Down-regulated    | 1471       | 1757       | 1494       | 1118      | 2190       | 1293       | 1929        | 1401       | 1443    |
| Significantly down-regulated | 13         | 14         | 12         | 4         | 33         | 9          | 17          | 7          | 8       |
| Total expression genes | 6746       | 6994       | 6864       | 6810      | 7293       | 6803       | 6421        | 7024       | 6973    |

Significantly up-regulated: log_{2} Ratio, ≥1, and probability, ≥0.8; Up-regulated: log_{2} Ratio, ≥1, and probability, <0.8; Not DEGs: genes not differentially expressed (0 ≤ |log_{2} Ratio| < 1); Down-regulated: log_{2} Ratio, ≤ −1, and probability, <0.8; Significantly down-regulated: log_{2} Ratio, ≤ −1, and probability, ≥0.8.

distributed into 43 categories, including the most predominant pathways such as “Intracellular membrane-bounded organelle,” “Adenyl ribonucleotide binding,” “Transition metal ion binding,” “Intrinsic to membrane,” “Cellular protein modification process,” “Plastid,” “Membrane and cell part,” “Binding and protein binding,” “Metabolic process,” and “Transcription, DNA-dependent.” The distributions among the pairwise comparisons were similar, except for small differences in the numbers of genes in each category and the total numbers of main category (Additional Files S8, S9). In addition, all of the annotated genes were mapped to terms in the KEGG database to search for significantly enriched genes involved in spliceosome, cell cycle, homologous recombination, or signaling pathways (Additional File S10).

In general, the 7687 DEGs in all comparisons were clustered as the union of DEGs. A total of 617 transcripts occurring
simultaneously in the three comparisons of three development stages between the monoecism and female were identified, and the comparison of the ratio values of these genes were used for clustering as the intersections of DEGs (Figure 5). Among the nine major clusters, 62 genes grouped into Cluster A were down-regulated in the three comparisons of developmental stages between the monoecism and the female, whereas the 97 genes in Cluster D were up-regulated in the three comparisons. GO analysis of these clustered genes indicated that the known genes were mainly involved in membrane or vesicle structures of intracellular or cytoplasmic components, activity and binding functions, response to light or hormone stimulus, transport, signal transduction, metabolic processes, and developmental processes involved in differentiation and reproduction (Additional File S11).

A heat map was generated of DEGs between different growth stages for both the monoecious and female buds (Figure 6). There were 621 DEGs in the four comparisons that clustered as the intersections of DEGs (Additional File S12). As shown in Figure 6, the DEGs in “ABML1 vs. ABML2” and “ABPL1 vs. ABPL2”...
FIGURE 5 | Hierarchical clustering (HCE) analysis of differentially expressed transcripts between monoecious and pistillate lines at three development periods (ABML1 vs. ABPL1, ABML2 vs. ABPL2, and RML vs. RPL. “a” was the control and “b” was experimental group in “a vs. b”). The clusters from A to I indicate the nine major clusters resulting from HCE analysis. Each line refers to data from one gene. The color bar represents the log_{10} RPKM and ranges from green (low expression) to red (high expression).

vs. ABPL2” were closely correlated to the DEGs in “ABML2 vs. RML” and “ABPL2 vs. RPL,” respectively, indicating that gene expression differences between monoecious and female lines were not obvious during the same development processes. However, this analysis revealed a greater number of DEGs between different development processes, with a relatively distant relationship, which suggests that transformation of apical bud from leaf to flower is a complex process.

Genes Associated with Sex Expression and Reproduction in Castor Bean

Based on prior knowledge of the putative involvement in sex determination and the expression levels and functions of DEGs between the monoecious and female buds in three development stages, several subgroups were assumed to be putatively related to sex expression and reproduction (Table 3). We identified seven genes involved in response to hormone stimulus, whose expression changed significantly in mutant females over the three developmental stages (Table 3). These genes included those that encode dynamin-2A, auxin response factor, PCI domain-containing protein, ATP-binding protein, spermidine synthase, Xaa-Pro amino peptidase, and a conserved hypothetical protein.

These genes are associated with tissue or organ developmental processes, and several of them take part in signal transduction, auxin transport, phytohormone biosynthesis, and metabolism (such as polyamine and abscisic acid).

In this study, we detected eight transcription factors that exhibited significantly different expression patterns between the pistillate and monoecious lines during the three developmental stages. These transcriptional regulators were identified as DNA-dependent transcription factors, including MADS box proteins, DNA-binding proteins, the Axial regulator YABBY5, transcription initiation factors, auxin response factor, and
## Table 3 | Some selected differentially expressed genes detected by digital expression profiling in apical bud of castor bean.

| Functional group | Gene ID | Gene annotation | log2ratio |
|------------------|---------|-----------------|-----------|
| **RESPONSE TO HORMONE STIMULUS** | | | |
| g|255573875|ref|XM_002527811.1| Dynamin-2A | −8.05 | −3.54 | −8.54 |
| | g|2555388885|ref|XM_002510462.1| Auxin response factor (transcription DNA-dependent) | −3.58 | 1.00 | −3.66 |
| | g|255546086|ref|XM_002514057.1| PCI domain-containing protein | −3.66 | 3.58 | 4.84 |
| | g|255543197|ref|XM_002512816.1| ATP-binding protein | 1.15 | −6.23 | 4.84 |
| | g|255550142|ref|XM_002516076.1| Spermidine synthase (auxin transport, polyamine biosynthetic process) | −5.71 | 1.36 | −1.54 |
| | g|255568254|ref|XM_002525056.1| Xaa-pro amino peptidase (auxin transport) | −1.77 | 1.50 | −1.78 |
| | g|255560413|ref|XM_002521176.1| conserved hypothetical protein (ABA and carotenoid metabolic process) | −4.56 | 4.56 | 3.58 |
| **TRANSCRIPTION, DNA-DEPENDENT** | | | |
| g|255558565|ref|XM_002520262.1| MADS box protein | −2.52 | −4.90 | −2.80 |
| | g|255583281|ref|XM_002510462.1| Auxin response factor | −3.58 | 1.00 | −3.66 |
| | g|255525732|ref|XM_002527155.1| Axial regulator YABBY5 | −1.18 | 8.07 | 5.69 |
| | g|255538455|ref|XM_002510247.1| DNA-binding protein | −2.08 | 1.31 | 4.14 |
| | g|255558989|ref|XM_002520472.1| protein with unknown function | −1.56 | 3.62 | −1.57 |
| | g|255558973|ref|XM_002520464.1| transcription initiation factor | 3.58 | 3.66 | 4.14 |
| **SIGNAL TRANSDUCTION** | | | |
| g|255555566|ref|XM_002518774.1| gcn4-complementing protein | −5.16 | −1.07 | −4.56 |
| | g|255550049|ref|XM_002516030.1| protein with unknown function | −4.92 | 3.62 | −1.57 |
| | g|255568372|ref|XM_002535052.1| Histidine-containing phosphotransfer protein | −5.39 | 4.87 | −3.62 |
| | g|255584316|ref|XM_002532848.1| Cyclic nucleotide-gated ion channel | −3.58 | 3.66 | −4.14 |
| **SEX DIFFERENTIATION** | | | |
| g|255572726|ref|XM_002519847.1| arginine/serine-rich splicing factor | −4.56 | 3.66 | 3.58 |
| **POLLINATION** | | | |
| g|25560126|ref|XM_002521035.1| acid phosphatase | −2.42 | 1.27 | 1.90 |
| **PROGRAMMED CELL DEATH** | | | |
| g|255539009|ref|XM_002510674.1| Ricinus communis cysteine protease | −4.22 | 3.66 | −4.54 |
| **DEVELOPMENTAL PROCESS INVOLVED IN REPRODUCTION** | | | |
| g|255577146|ref|XM_002529411.1| 1,4-alpha-glucan branching enzyme | −1.02 | 4.20 | −3.62 |
| | g|255588742|ref|XM_002534658.1| DNA replication helicase dna2 | −1.45 | 5.71 | −1.96 |
| | g|255568769|ref|XM_002525310.1| sentrin/sumo-specific protease | −1.66 | 2.32 | 1.29 |
| | g|255578470|ref|XM_002530054.1| U4/U6 small nuclear ribonucleoprotein Prp4 | −6.24 | 4.20 | 4.22 |
| | g|255544821|ref|XM_002513426.1| sorting and assembly machinery (sam50) protein | −3.66 | 4.20 | 4.14 |
| **HISTONE DEMETHYLATION/METHYLATION** | | | |
| g|255565532|ref|XM_002523711.1| eukaryotic translation initiation factor 2c | −3.58 | 1.29 | 1.67 |
| | g|255575536|ref|XM_002529823.1| set domain protein | 2.78 | −4.20 | −4.14 |
| | g|255553976|ref|XM_002517983.1| DNA (cytosine-5)-methyltransferase | −4.89 | 3.58 | −3.62 |
| | g|255558643|ref|XM_002520301.1| s-adenosyl-methyltransferase mara | 1.23 | 3.66 | 5.33 |

The RNA polymerase sigma factor rpoD1. Auxin response factor is respond to hormone stimulus and participates in hormone-mediated signaling pathways, also plays a role in tissue and organ developmental processes. Furthermore, one subgroup is composed of four genes related to signal transduction, including Gcn4-complementing protein, histidine-containing phosphotransfer protein, an unknown protein, and cyclic nucleotide-gated ion channel, which exhibited consistent expression changes: down-regulation at the early leaf-bud stage, up-regulation at the 5–7-leaf stage, and down-regulation at the young raceme stage. Another two genes listed above (PCI domain-containing protein and ATP-binding protein) play roles in signal transduction in addition to response to hormone stimulus.
One gene (arginine/serine-rich splicing factor) associated with sex differentiation and another gene (acid phosphatase) related to pollination were identified as differentially expressed between monoecious and pistillate apical bud development. In female apical buds, both genes were distinctly down-regulated at the early leaf-bud stage, but up-regulated at the 5–7-leaf and initial raceme stages. In addition, expression of a PCD (programmed cell death)-related gene that encodes cysteine proteinase was altered in pistillate apices, with down-regulation at the early leaf-bud and initial raceme stages and up-regulation during the period of transition from leaf bud to floral bud. Several genes involved in reproduction exhibited the same expression patterns in pistillate terminal bud: sentrin/sumo-specific protease, the U4/U6 small nuclear ribonucleoprotein Prp4, sorting and the assembly machinery (sam50) protein were down-regulated at the early leaf stage, but up-regulated at the 5–7-leaf and small raceme stages, (Table 3); notable exceptions were 1,4-alpha-glucan branching enzyme and DNA replication helicase (dna2). Histone modification and DNA methylation or demethylation, play an important role in the regulation of gene expression or silencing during developmental processes. We identified four genes involved in histone methylation and demethylation (Table 3), including eukaryotic translation initiation factor 2c, set domain protein, DNA (cytosine-5)-methyltransferase, and s-adenosyl-methyltransferase (mraw). These genes were also significantly differentially expressed between the monoecious and female apical bud development.

**DEGs Were Confirmed by Quantitative RT-PCR**
To confirm the differentially expressed genes in the monoecious and female apical buds, 14 genes were selected for quantitative RT-PCR analysis at the three apical bud developmental stages. Representative genes selected for qPCR validation were those involved in response to hormone stimulus, transcription factor, signal transduction, sex differentiation, pollination, reproduction and histone methylation/demethylation because those phenomena were putative to associate with plant sex determination (Guo et al., 2010; Wu et al., 2010). The expression of the 13 genes (Dynamin-2A, Auxin response factors, ATP-binding protein, Spermidine synthase, auixin transport, conserved hypothetical protein, MADS box protein, two unknown proteins, arginine/serine-rich splicing factor, acid phosphatase, DNA replication helicase dna2, and eukaryotic translation initiation factor 2c) indicated by qRT-PCR was basically congruent with the DGE analysis patterns (Figure 7). Only one gene (s-adenosyl-methyltransferase mraw) did not show consistent expression between the qRT-PCR and DGE data sets.

**Differential Hormone Levels in the Apices and Racemes Between the Monoecism and Female**
DGE analysis and qRT-PCR verification showed that some genes which response to hormone stimulus were expressed differentially in apices and racemes between the monoecism and female, and possibly associated with castor sex expression. To validate whether the levels of phytohormone in the apices and inflorescences of the monoecious line and the pistillate line vary or not, the IAA, ABA, JA, and GAs contents of the two lines at three developments were quantitatively measured. The result showed that the auxin in the tow castor lines indicated a similar variation pattern in that IAA was equivalent or increased from the 3–4-leaf stage to the 5–7-leaf stage, and later significantly reduced (P < 0.01) (Figure 8A; Additional File S13). It was worthwhile to note that the auxin level in pistillate line was remarkably higher than that in the monoecism at each of the three development stages. Four kinds of GAs were performed in the measurement, but only GA4 were detected with a relatively low level, and other three GA were not detected. The GA4 level in the apices except little racemes also showed significant differences between the monoecism and female (Figure 8A; Additional File S13). The level of GA4 in the female was significantly lower (P < 0.01) than that in the monoecism at the stage of early leaf bud, but markedly higher (P < 0.01) at the 5–7-leaf stage and similar at the stage of young inflorescence.

The ABA levels in the monoecism and female revealed an identical change pattern from early leaf apices to infant racemes, with a notable consecutive reduction (Figure 8B; Additional File S13). The ABA level in female was significantly higher (P < 0.01) than that in monoecism at early leaf apices, but notable lower (P < 0.01) at following 5–7-leaf stage, and comparative level at little raceme stage. In addition, the variation in endogenous JA levels showed a similar pattern as IAA in the two castor lines during the three development stages (Figure 8B; Additional File S13). However, opposite to IAA, the JA level in the pistillate line was extremely significant lower (P < 0.01) than that in the monoecism at the two apical development stages, and significantly higher (P < 0.01) than that in monoecism at the stage of little raceme.

**DISCUSSION**
Sex expression in plants is controlled by genetic factors and non-genetic fluctuations (Shiffriss, 1956; Ainsworth, 1999), and phytohormones play important roles in sex determination in some species (Yin and Quinn, 1995; Ainsworth, 1999; Tanurdzic and Banks, 2004). In cucumber, ethylene, a true sex hormone, was proven to be a “female” hormone that exerts a strong feminizing effect (Dellaporta and Calderon-Urrea, 1993); Auxin also promotes cucumber femaleness, potentially playing an indirect role in sex determination by promoting the action of ethylene (Yamasaki et al., 2000). The level of gibberellic acid (GA), which acts downstream of ethylene or through a parallel pathway (Ainsworth, 1999), is correlated with male tendency. Sex expression in castor bean is also regulated by plant hormones (Shiffriss, 1961; Tan et al., 2011), but in contrast to cucumber, ethylene and ethylene-like substances (NIA 10637) result in masculinization and can transform female flowers into male ones in monoecious plants (Philipos and Narayanaswamy, 1976); moreover, GA causes feminization in castor, and spraying monoecious inbreds with GA can markedly increase female tendency (Shiffriss, 1961). In addition, previous work showed...
that the activity of auxin-like substances causes feminization of castor bean, a process influenced by kinetin and morphactin (Kumar and Rao, 1981). Morphactin causes maleness in R. communis (Rkey, 1978; Varkey and Nigam, 1982), whereas daminozide promotes femaleness with reduction in the position of bearing the first female flower and the ratio of male and female flowers (Chauhan et al., 1987). To date, however, the key sex hormone in castor bean remains unknown. In this
study, we identified several genes involved in the response to hormone stimulus (Table 3; Figure 7), including auxin transport and auxin response factor genes and other genes involved in plant hormone-mediated biosynthetic and metabolic process, such as polyamine biosynthesis and abscisic acid and carotenoid metabolism. In the pistillate line, three genes (auxin response factor, spermidine synthase, and Xaa-Pro amino peptidase) related to auxin response and transport, were down-regulated in early leaf apices and young racemes, but up-regulated during the transformation from leaf apices to floral buds. In addition, the result of hormone measurement detected that auxin level in the female was higher than that in the monoeocious at each of the three development stages and extremely significant higher at the 5–7-leaf stage. The levels of JA, ABA, and GA in the apices also displayed significant differences between the pistillate line and the monoeocious line, except the GA and ABA level in the infant racemes (Figure 8). Our results suggested that DEGs involved in the pathways of plant hormone response and hormone-mediated biosynthetic and metabolic processes, may participate in or affect the regulation of sex expression and reproduction by changing the hormone level and transport, which will be elucidated in further research. Our result was consistent with previous reports (Heslop-Harrison, 1957; Rkey, 1978). Those authors showed that variation in sex expression in plants is associated with variations in the hormone level (Heslop-Harrison, 1957), and that auxins promote maleness (Wittwer and Hillyer, 1954). Interference in auxin transport, or quick degradation of the hormone, can decrease the auxin level, resulting in altered sex expression (Thomson and Leopold, 1974; Gaganias and Berg, 1977). The fact that lateral branches of castor bean plants produce only male flowers after decapitation supports the view that inhibition of auxin transport can result in maleness (Rao, 1969). Furthermore, our previous research (Wang et al., 2012) demonstrated that IAA and ABA levels in the inflorescence and flower vary remarkably between the pistillate and the monoeocious, also suggesting that some phytohormones such as IAA etc. and related genes are important for sex determination in R. communis. In plants, phytohormones modulate various growth and developmental events through signal transduction pathways (Bleecker and Kende, 2000; Davies, 2004). Notably, in this study, three genes involved in signal transduction exhibited expression patterns similar to those of genes involved in auxin response and transport (i.e., down-regulated in early leaf apices and small racemes, but up-regulated during the transformation stage from leaf to floral bud) in the pistillate line. However, other genes involved in signal transduction were down-regulated throughout all three stages. Subsequent investigations of these genes and crosstalk between other hormones and auxin may reveal their specific roles in sex expression or sex determination.

Castor is a monoeocious species, with pistillate flowers borne on short pedicels along the upper portion of the raceme and staminate flowers borne similarly on the lower portion (Brigham, 1967). Castor flowers are unisexual and lack petals. Male and female flowers differ radically in general morphology and size: male flowers are devoid of rudiments of organs of the opposite sex, whereas pistillate flowers occasionally exhibit vestiges of stamen (Wu et al., 1998). The occasional appearance of rudimentary androecium in Ricinus female flowers suggests that like unisexual flowers of other species (Atsmon and Galun, 1960; Cheng et al., 1983; Bracale et al., 1991; Malepszy and Niemirowicz-Szczyt, 1991; Veit et al., 1993), castor flowers (especially pistillate flowers) pass through a “bisexual stage” during which development of all floral organs is initiated. Castor sex determination probably involves PCD of opposite sex organs in the flower and inflorescence, similar to what is observed in several other plant species such as maize, cucumber, and campion (Ye et al., 1991; Dellaporta and Calderon-Urrea, 1993; Kater et al., 2001); these plants follow a sex determination pathway involving arrest of preformed sexual organs in bisexual primordia in which some PCD-related genes participate (DeLong et al., 1993). In this study, we detected a PCD-related gene (cysteine protease) that was significantly up-regulated in female apices at the 5–7-leaf stage, but obviously down-regulated at the two other stages (early leaf apices and initial inflorescence). This...
result was consistent with recent findings that PCD-related genes, including cysteine protease, are more highly expressed at the peak of anther abortion in CMS and GMS lines of cotton (Lorrain et al., 2003; Wei et al., 2013). It is possible that the cysteine protease might play a role in Ricinus apical development, and thus be associated directly or indirectly with sex expression. Our study identified hundreds of genes potentially involved in sex determination, including one (arginine-serine-rich splicing factor) associated with sex differentiation; this gene was up-regulated during apical transformation and initial raceme formation in female apical buds.

Transcription factors such as the maize DELLA protein D8 and the melon zinc finger protein CmWIP1 have been functionally associated with the sex determination process (Peng et al., 1999; Martin et al., 2009). Moreover, several transcription factors were identified as differentially expressed during sex determination in cucumber (Guo et al., 2010; Wu et al., 2010), including zinc finger transcription factors, Aux/IAA transcription factor, auxin-induced protein, BR11, BR11-associated receptor kinase, MYC transcription factor, BEL1-like homeodomain protein, bHLH proteins, WRKY DNA-binding protein, and NAC domain protein, IF-2, and so on. Consistent with the results in cucumber, we identified eight DNA-dependent transcription genes in castor bean that were significantly differentially expressed between the three apical developmental stages in the pistillate line. Of these, auxin response factor, MADS box protein, IF-2, and two DNA-binding proteins were also confirmed to be differentially expressed during sex determination. These transcription factors detected in this study were mainly binding proteins, response factors, initiation and regulation factors, and even MADS box proteins; the relationship between these transcription factors and plant sex determination remains to be determined, and should be investigated in future work. In addition, we also identified a few genes involved in histone demethylation/methylation and reproductive processes. DNA methylation and histone modification play essential roles in genome management and control gene expression or silence (Cheng and Blumenthal, 2008). DNA methyltransferase, the main enzyme involved in DNA methylation, was detected in female apices; it was obviously up-regulated at the stage of transformation from leaf to floral apices, and down-regulated at the other two developmental stages, relative to monoecious apices (Table 3). DNA methyltransferase is not only associated with DNA methylation, but is also associated with many important biological activities, including cell proliferation and senescence (Berger, 2007). The increase in methyltransferase gene expression during the transitional stage may change the level of DNA methylation, leading to the subsequent repression of male flower gene expression and the appearance of only female flowers in female racemes. Genes associated with the reproduction process, identified as DEGs here and in future research, will contribute to elucidation of the molecular mechanisms underlying Ricinus sex determination and sex patterns.

DATA DEPOSITION

The Illumina pair-end reads of Ricinus communis L. obtained in the study are available at NCBI SRP064760.

AUTHOR CONTRIBUTIONS

YX conceived and designed the study, amended the manuscript. TM participated in its design and performed the experiments, analyzed the data and drafted the manuscript. XJ and WL prepared the samples, RNA extraction, cDNA library construction and helped with the sequencing. FC collected and prepared the samples. HJ provided the castor seeds and took part in field cultivation and bud collections. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2015.01208

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

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