Generation and classification of transcriptomes in two Croomia species and molecular evolution of CYC/TB1 genes in Stemonaceae

Ruisen Lu a, Wuqin Xu a, Qixiang Lu a, Pan Li a, Jocelyn Losh a, Faiza Hina a, Enxiang Li b, Yingxiong Qiu ab

a Key Laboratory of Conservation Biology for Endangered Wildlife of the Ministry of Education, and College of Life Sciences, Zhejiang University, Hangzhou, 310058, China
b College of Life Sciences, Nanchang University, Nanchang, 330031, China

1. Introduction

Stemonaceae Engl., a monocotyledonous flowering plant family, belongs to Pandanales and consists of four genera: Croomia Torr., Stemona Lour., Stichoneuron Hook. f. and Pentastemona Steenis, with about 37 known species (Chase et al., 2016; Christenhusz et al., 2016). The family is native to seasonal climate areas across Southeast Asia and tropical Australia, with one species [Croomia pauciflora (Nutt.) Torr.] in North America (Li et al., 2008). Because the roots and rhizomes of some Stemona and Croomia species are found to contain many bioactive and structurally unique alkaloids, such as protostemonine, stichoneurine, and croomine groups (Greger, 2006; Lin et al., 2006, 2008; Kongkiatpaiboon et al., 2011; Chen et al., 2017), they are widely used in Southeast Asian folk medicine to treat cough, traumatic injury, and enteric helminth worms (Lee et al., 2008; Chen et al., 2017). Thus, the family Stemonaceae has been a focus of phytochemical and pharmacological research (Kongkiatpaiboon et al., 2011; Chen et al., 2017). Among the four genera, Croomia exhibits a typical EA-ENA disjunct distribution across the Pacific Ocean, comprised of three herbaceous perennial species: Croomia heterosepala (Bak.) Oku. and Croomia japonica from East Asia; C. pauciflora from Southeastern North America (Li et al., 2008; Fang et al., 2013). Due to their small range size and small number of populations, all three species of Croomia are treated as rare and endangered in China, Japan, and the United States (Estill and Cruzan, 2001; Wang and Xie, 2004). However, except for a few traditional molecular markers (e.g. ISSR, nSSR) developed and used in former studies (Li et al., 2008; Fang et al., 2013), no genomic resources have been reported for these endangered Croomia species, which hinders the studies on

A B S T R A C T

The genus Croomia (Stemonaceae) is an excellent model for studying the evolution of the Eastern Asia (EA)—Eastern North America (ENA) floristic disjunction and the genetic mechanisms of floral zygomorphy formation. In addition to the presence of both actinomorphic and zygomorphic flowers within the genus, species are disjunctively distributed between EA and ENA. However, due to the limited availability of genomic resources, few studies of Croomia have examined these questions. In this study, we sequenced the floral and leaf transcriptomes of the zygomorphic flowered Croomia heterosepala and the actinomorphic flowered Croomia japonica, and used comparative genomic approaches to investigate the transcriptome evolution of the two closely related species. The sequencing and de novo assembly of transcriptomes from flowers of C. heterosepala (ChFlower), flowers of C. japonica (CjFlower), and leaves of C. japonica (CjLeaf) yielded 57,193, 62,131 and 64,448 unigenes, respectively. In addition, estimation of Ka/Ks ratios for 11,566 potential orthologous groups between ChFlower and CjFlower revealed that only six pairs had Ka/Ks ratios significantly greater than 1 and are likely under positive selection. A total of 429 single copy nuclear genes (SCNGs) and 21,460 expression sequence tags-simple sequence repeats (EST-SSRs) were identified in this study. Specifically, we identified seven CYC/TBI-like genes from Stemonaceae. Phylogenetic and molecular evolution analyses indicated that these CYC/TBI-like genes formed a monophyletic clade (SteTBL1) and were subject to strong purifying selection. The shifts of floral symmetry in Stemonaceae do not appear to be correlated with TBL copy number.

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population genetics, conservation management, and adaptive divergence of these species.

In addition, our previous molecular phylogenetic analyses using chloroplast DNA sequence variation of the trnL-F region have supported the monophyly of *Croomia* and a sister relationship of *C. pauciflora* to the two Asian species (Li et al., 2008). Molecular dating suggests the two Asian species diverged in the Mid-to-Late Pleistocene (0.84–0.13 mya), and *C. pauciflora* diverged at the Plio-Pleistocene boundary (<2.6 mya). Although chloroplast markers are often used to infer phylogenetic relationships, very low resolution of the dataset hampered any further statistical inference. Moreover, since the possible presence of multiple haplotypes and/or independent chloroplast lineages does not correspond with the actual species divergence, our previous molecular dating may only represent the divergence of a particular genetic locus but not the divergence of species. Thus, application of coalescent-based models or related simulation-based approaches to multi-locus sequence data should allow us to explore in more detail the prevalent time scales, geographic modes, and demographic history of population and species divergences (Li et al., 2012; Dolman and Joseph, 2016). However, the genus lacks sequence data at a genomic scale to facilitate such investigations.

The two Asian species form a parapatric species pair with abutting ranges in South Japan, but *C. japonica* also occurs disjunctively on the adjacent Asiatic mainland in East China. There is considerable difference in floral characters of the two species (Rogers and Harpending, 1992). For example, all four tepals of *C. japonica* are homomorphic with a re-curved edge, whereas those of *C. heterosepala* have a straight edge, and one tepal is much larger than the other three (Okuyama, 1944; Ohwi, 1965; Li et al., 2008). Thus, like other species of Stemonaceae, the flowers of *C. japonica* are radially symmetrical (i.e., actinomorphic). By contrast, *C. heterosepala* represents the sole species with bilaterally symmetrical (i.e., zygomorphic) flowers within this family. *Croomia* and its related genera, therefore, provide an ideal model to investigate the origin and maintenance of flower symmetry; the first step to addressing this problem is to identify the genes responsible for flower symmetry. More recently, functional studies in model *E. antirrhinum majus* (Preston and Hileman, 2009); *Oryza sativa* (Yuan et al., 2009) and non-model species [e.g. *Caprifoliaceae* and *Loniceraceae* (Howarth et al., 2011); *Commelinaceae* (Preston and Hileman, 2012); *Ranunculaceae* (Jabbour et al., 2014)] have demonstrated that TCP-like genes (sometimes referred to as TB1-like or TBL in monocots), which belong to the class II TCP (TB1, CYC and PBF) family, are involved in the establishment and maintenance of zygomorphic flowers. However, to date, few studies are available for monocots (Bartlett and Specht, 2011; Preston and Hileman, 2012; Hoshino et al., 2014), and no TCP or CYC/TB1 genes have been identified from Stemonaceae.

Transcriptome sequencing or RNA-Seq is one of the most efficient and cost-effective methods currently available for gene discovery and developing massive genome-wide markers in non-model organisms (Wen et al., 2015). Recent studies have demonstrated the utility of these data for resolving the relationships of diverse lineages of organisms (i.e. RNA-Seq phylogenetics) (Zhou et al., 2017), inferring demographic histories (Zhu et al., 2016), estimating genomic variation (Kawakami et al., 2014), and identifying genetic bases of adaptive divergence (Wen et al., 2015; Mao et al., 2016). Furthermore, transcriptome analysis provides valuable insights into genes and gene activities responsible for differences in organ morphology during the developmental processes (Liu et al., 2013; Ma et al., 2014). In this study, using the Illumina HiSeq 2000 platform, we obtained the RNA sequence data of two flower samples (C. heterosepala and *C. japonica*) and one leaf sample (C. japonica). De novo assembly of these transcripts was conducted to characterize the transcriptomes of the two flowers and one leaf, respectively. Based on transcriptome data of *Croomia*, a set of putative single-copy genes were screened in a genus framework by filtering strict orthologs. By pairwise comparison of the orthologous sequences from each species pair, candidate genes under adaptive selection in speciation or population differentiation were identified. In addition, we developed a large set of EST-SSR loci and validated a subset of them, yielding massive potential molecular markers. Finally, we isolated and sequenced CYC/TB1 (TCP) genes across four genera (7 species) of Stemonaceae to reconstruct the CYC/TB1 gene tree and estimate its molecular evolution in this family.

## 2. Materials and Methods

### 2.1. Plant samples, cDNA library preparation, and illumina sequencing

During April 2010, living plants of *C. heterosepala* from Nara, Japan (34°31′N, 135°41′E) and *C. japonica* from Mt. Tianmu, China (30°20′N, 119°26′E) were transplanted to the Botanical Garden of Zhejiang University, Hangzhou, China. Fresh flowers of *C. heterosepala* (ChFLOWER) and *C. japonica* (CjFLOWER) at the anthesis stage, as well as juvenile leaf samples of *C. japonica* (CjLeaf), were harvested from these living plants and immediately frozen in liquid nitrogen for storage at −80 °C until total RNA extraction. For each tissue, total RNA was extracted from a mixture of three individuals using TRIzol Reagent (Invitrogen Life Technologies, USA) according to the manufacturer’s recommendations. RNA quality and quantity were assessed using gel electrophoresis and a NanoDrop spectrophotometer 2000 (Thermo Scientific, Wilmington, DE, USA). Sequencing libraries were generated from 3 μg total RNA using NEBNext Ultra™ RNA Library Prep Kit for Illumina® (NEB, USA) following manufacturer’s protocol, and index codes were added to attribute sequences to each sample. After validating the library quality with an Agilent 2100 Bioanalyzer system, the constructed libraries were sequenced on an Illumina HiSeq 2000 platform and paired-end reads were generated. All of the sequencing data were deposited in the Sequence Read Archive (SRA) of the NCBI database under accession numbers: SRX3328777 (ChFLOWER), SRX3328635 (CjFLOWER), and SRX3328351 (CjLeaf).

### 2.2. Raw data processing and de novo assembly

Raw sequence reads were firstly filtered through in-house Perl scripts. In this step, clean reads were obtained by removing reads containing adapters, reads with more than 5% unknown bases (N bases), and reads with more than 20% of low-quality bases (quality value ≤ 20). At the same time, Q20, Q30, GC content and the sequence duplication level of the clean data were calculated. All the downstream analyses were based on these high-quality clean reads. The high-quality clean reads from ChFLOWER, CjFLOWER, and CjLeaf were then assembled de novo separately using TRINITY (Haas et al., 2013) with “min_kmer_cov” set to 2 and all other parameters set to default. These assembled unigene sets were further processed by sequence splicing and redundancy removal using TGICL software (Pertea et al., 2003) to retrieve non-redundant unigenes. Furthermore, the individual assembled unigene sets from CjFLOWER and CjLeaf were pooled together and assembled using TGICL software (Pertea et al., 2003) into non-redundant unigenes that represented the *C. japonica* transcriptome (CjTranscriptome).

### 2.3. Functional annotation and prediction of protein-coding sequence (CDS) regions

To predict the putative functions of the assembled unigenes in ChFLOWER, CjFLOWER, CjLeaf and CjTranscriptome, the unigenes were...
utilized for homology searches against the National Center for Biotechnology Information (NCBI) non-redundant (Nr) protein database, Swiss-Prot protein database (http://www.expasy.ch/sprot), the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways database (Kanehisa et al., 2008) and the Cluster of Orthologous Groups (COG) database (http://www.ncbi.nlm.nih.gov/COG/) using BLASTX (Altschul et al., 1990) with an E-value cut-off of 1e−5. Based on the best BLASTX hits from the NCBI Nr database, Gene Ontology (GO) terms (Ashburner et al., 2000) of the unigenes were performed using BLAST2GO version 2.6.0 (Conesa et al., 2005) with an E-value (GO) terms (Ashburner et al., 2000) of the unigenes were performed cut-off of 1e−5. To obtain information on general functional categories, the distributions of level-2 GO terms for all unigenes were plotted with the Web Gene Ontology Annotation Plot (WEGO) (Ye et al., 2006) for three main categories: biological process, molecular function and cellular component (http://www.geneontology.org). In addition, unigenes were further queried against the Nr (NCBI non-redundant nucleotide sequences) database using BLASTN with an E-value cut-off of 1e−5.

The protein-coding region sequences (CDS) of unigenes were predicted according to the Blast results against the Nr, Swiss-Prot, KEGG and COG databases (E-value < 1e−5) and translated into amino acid sequences using the standard codon table. If the results between different databases were in conflict with each other, a priority order of Nr, Swiss-Prot, KEGG, and COG was followed. For those unigenes that could not align to any of the above databases, ESTScan (Iseli et al., 1999) was used to predict CDS regions and determine the amino acid sequences.

2.4. Ka/Ks ratios of orthologous pairs between ChFlower and CjFlower transcriptomes and mining of single copy nuclear genes (SCNGs)

Based on the predicted CDS regions of ChFlower and CjFlower transcriptomes, putative orthologous groups between these two flower transcriptomes were obtained using OrthoMCL v2.0.9 (Li et al., 2003) with default parameters by identifying clusters with only one sequence from each flower transcriptome. Those orthologs were removed following three criteria: (A) the alignment lengths were <150 bp, (B) Ka or Ks values were not applicable, (C) Ks values were more than 0.1, which is a benchmark of potential paralogs (Bustamante et al., 2005; Ai et al., 2015). The YN algorithm (Yang and Nielsen, 2000) implemented in KAKS_CALCULATOR v1.2 (Zhang et al., 2006) was employed to calculate non-synonymous rates (Ka), synonymous rates (Ks) and Ka/Ks (ω) ratios of each putative pair of orthologs.

For the mining of putative single copy nuclear genes (SCNGs), 959 single copy nuclear genes shared by Arabidopsis thaliana, Populus trichocarpa, Vitis vinifera, and O. sativa (usually referred as APVO genes, Duarte et al., 2010) were used for our analysis. We retrieved the protein sequences encoded by APVO genes from the TAIR10 database (Duarte et al., 2010) and queried these sequences against the putative orthologous genes between ChFlower and CjFlower using TBLASTN with a threshold E-value of 1e−10. All the queries with BLAST hits were considered to be putative single copy nuclear genes (SCNGs) in the Croomia species.

2.5. Detection and validation of EST-SSRs

EST-SSRs were identified from unigenes of CjTranscriptome using the program MISA (http://pgrc.ipk-gatersleben.de/misa) (Dieringer and Schlötterer, 2003) with thresholds of 12, 6, 5, 5, 4, and 4 repeat units for mono-, di-, tri-, tetra-, penta-, and hexa-nucleotide motifs, respectively. Primer pairs for the SSRs were designed using Primer Premier v5.0 (Premier Biosoft International, Palo Alto, California, USA). To assess the reliability and cross-species transferability of the identified EST-SSRs, 100 primer pairs were randomly selected to test for amplification across 15 populations (two samples per population) of C. heterosepala, C. japonica and C. pauciflora (Table S1).

The PCR amplification reactions were carried out in 30 μl reaction volume containing 17 μl 2x master mix (TSINGKE, Beijing, China), 9 μl of ddH2O, 1 μl of each primer, and 2 μl of genomic DNA. PCR amplifications were performed using a thermal cycler GeneAmp PCR System 9700 (Applied Biosystems, Foster City, CA, USA) following the procedures in Zhu et al. (2016). The PCR products, along with a 100 bp marker (TaKaRa, Dalian, Liaoning, China), were electrophoresed on 12% non-denaturing polyacrylamide gels stained with silver staining to assess amplification success and polymorphism. The polymorphic primer pairs were synthesized again with fluorescent dyes (TAMRA, HEX, or 6-FAM) at the 5′ end of the forward primer and were then used to amplify DNA from all 15 populations (240 individuals in total) of C. heterosepala, C. japonica and C. pauciflora for genetic diversity analysis (Table S1). PCR amplification followed the conditions described above. Fragments of PCR product were separated on a 3730xl DNA Analyzer (Applied Biosystems) and then the alleles were manually scored and determined using GENEMARKER v2.2.0 (SoftGenetics, PA, USA) with GeneScan 500 LIZ as an internal size standard. Finally, the number of observed alleles (Ho) as well as observed (Hd) and expected (He) heterozygosities, and the polymorphism information content (PIC) values were calculated using CERVUS v3.0.3 (Kalinowski et al., 2007). The significance of departures from Hardy–Weinberg equilibrium (HWE) and linkage disequilibrium (LD) between all pairs of polymorphic loci were analyzed using GENEPOP v4.0.7 (Rousset, 2008). Frequencies of null alleles were estimated in the FRENA package (Chapuis and Estoup, 2007) following the expectation maximization (EM) method (Dempster et al., 1977).

2.6. Analysis of CYC/TB1-like genes of TCP transcription factors in Stemonaceae

In order to identify CYC/TB1-like genes in C. heterosepala and C. japonica, three strategies were adopted in this study. First, TCP domain sequences of O. sativa (23 accessions) and A. thaliana (24 accessions) downloaded from PlantTFDB v4.0 (planttfdb.chpku.edu.cn) were used as query entries to perform BLASTP searches against the protein sequences from both ChFlower and CjFlower transcriptomes (E-value < 1e−5). In addition, all the output protein sequences were scanned using InterProScan (https://www.ebi.ac.uk/interpro/search/sequence-search) to remove those sequences without a whole TCP domain. Second, high-quality reads that mapped to Commelinaceae TB1-like genes (Preston and Hileman, 2009) were directly assembled into CYC/TB1-like genes. Third, CYC/TB1-like genes were amplified from genomic DNA of C. heterosepala, C. japonica, and five other Stemonaceae species (C. pauciflora, Stemona tuberosa, S. japonica, Stichoneuron caudatum, and Pentastermona egregia) representing all four genera. The primers and the methodology for amplification used in this study were described in Howarth and Donoghue (2005). To distinguish and classify the TCP genes of ChFlower and CjFlower transcriptomes, TCP genes obtained from the first strategy were divided into class I PCF-like, class II CIN-like, and class II CYC/TB1-like groups. The amino acid sequences of TCP domains from O. sativa, A. thaliana, ChFlower and CjFlower were aligned using ClustaW (Thompson et al., 1994) and manually edited if necessary. Then, the Neighbor-Joining (NJ) phylogenetic tree was constructed using MEGA v7 (Kumar et al., 2016) with 1000 bootstrap replicates. To further determine ortholog or copy number of newly-determined Stemonaceae CYC/TB1-like sequences, a total of 72 amino acid sequences, which have been confirmed to be related to the formation of flower symmetry (Yuan et al., 2009; Bartlett and Specht, 2011;
Preston and Hileman, 2012; Hoshino et al., 2014; Citerne et al., 2017) plus 7 newly-determined Stemonaceae sequences were used to generate Bayesian inference (BI) and maximum likelihood (ML) trees. The best-fitting model (JTT + G + I) was selected by ProtTest v2.4 (Abascal et al., 2005). Bayesian inference analyses were conducted in MrBayes v3.2 (Ronquist and Huelsenbeck, 2003). The Markov chain Monte Carlo (MCMC) algorithm was run for two million generations with trees sampled every 500 generations. The first 25% of generations were discarded as burn-in. A 50% majority-rule consensus tree was constructed from the remaining trees to estimate posterior probabilities (PPs). Maximum likelihood analysis was conducted using PhyML v3.0 (Guindon et al., 2010) with 1000 bootstrap replicates.

Both the CYC/TB1-like phylogeny and codon alignments were used to estimate the synonymous (dS) and nonsynonymous (dN) substitution rates (r) in CODEML as part of the PAML package (Yang, 2007). Analyses were conducted under three models: the ‘one-ratio’ model (M0) assumes the equal r (dN/dS) value for the entire tree, the ‘free-ratios’ model (M1) assumes an independent r value for each branch and the ‘two-ratio’ model (M2) assumes that one or more of branches (the ‘foreground’ branch) have a distinct r value different from the remaining branches (the ‘background’ branches). Moreover, 12 hypotheses of the ‘two-ratio’ model (M2) were tested by selecting each clade (clades labeled in Fig. 3) as foreground: Hypothesis 1 (H1): ZinTBL2, H2: ComTB1a, H3: PoaTBL2, H4: ZinTBL1a, H5: ZinTBL1, H6: AlstTCP1, H7: SteTBL1, H8: PoaTBL1, and H9: AcoTBL were selected as foreground, respectively; H10: clade (ZinTBL1a and ZinTBL1b) was selected as foreground; H11: clade (ZinTBL2, ComTB1a and PoaTBL2) was selected as foreground; H12: specific r values are estimated for each of clades leading to ZinTBL2, ComTB1a, PoaTBL2, ZinTBL1a, ZinTBL1, AlstTCP1, SteTBL1, PoaTBL1, and AcoTBL. To evaluate the goodness of fit of the data, a likelihood ratio test (LRT) was performed by using the latter models with respect to the M0 model.

3. Results
3.1. RNA-sequencing and de novo assembly

The Illumina sequencing of cDNA libraries of ChFlower, CjFlower, and CjLeaf yielded 49,881,224, 50,481,150 and 60,582,220 raw reads, respectively. After filtering and trimming raw reads, 43,683,202, 44,355,666 and 54,416,848 clean reads of ChFlower, CjFlower and CjLeaf were generated, with the Q20 percentage over 97% (Table 1). Through de novo assembly, 76,976, 77,344, 79,538 contigs with an N50 length of 1,452, 1,461, 1,641 bp and mean length of 916, 951, 1027 bp were obtained for ChFlower, CjFlower and CjLeaf, respectively. These contig sequences were further assembled into 57,193 unigenes (average length = 1042 bp, N50 = 1603 bp) for ChFlower, 62,131 unigenes (average length = 1044 bp, N50 = 1575 bp) for CjFlower and 64,448 unigenes (average length = 1106 bp, N50 = 1737 bp) for CjLeaf (Table 1).

By further combining and reassembling the CjFlower and CjLeaf unigene sets, we obtained 86,457 unigenes with a mean length of 1121 bp and an N50 of 1732 bp, which represented the C. japonica transcriptome (CjTranscriptome). Additionally, not only the number of unigenes (86,457) but also the percentage of longer unigenes (>1000 bp, 42.12%) in CjTranscriptome were greater than those in CjFlower and CjLeaf transcriptomes (Table 1). The distribution of unigene length also indicated a common pattern that smaller size
groups possessed more unigenes, and more than half ranged from 200 to 1000 bp (Table 1).

3.2. Functional annotation and CDS predication

Based on a BLAST search, 39,919 (69.80%) unigenes from ChFlower, 44,209 (71.15%) unigenes from CjFlower, 43,375 (67.30%) unigenes from CjLeaf and 58,895 (68.12%) unigenes from CjTranscriptome had at least one Blast match against Nr, Swiss-Prot, KEGG, COG, GO or Nt databases (Table 2). Based on GO function classifications of the two flower transcriptomes, 18,436 (32.23%) unigenes from ChFlower and 20,376 (32.80%) unigenes from CjFlower were assigned into three main categories and 54 subcategories, and the distribution of GO terms was consistent with each other (Fig. 1).

Among these subcategories, ‘metabolic process’ (ChFlower/CjFlower: 11,178, 60.63%/12,352, 60.62%) and ‘cellular process’ (9,971, 54.08%/11,106, 54.51%) were the two mostly dominant GO terms for the biological process category, while ‘cell’ (9,143, 49.59%/10,131, 49.72%) and ‘cell part’ (9,143, 49.59%/10,131, 49.72%) were most abundant in the cellular component category, and ‘catalytic activity’ (9,640, 52.29%/10,575, 51.90%) and ‘binding’ (8,199, 44.47%/8,993, 44.14%) were highly representative for the molecular function category (Fig. 1). Beyond that, the distributions of CjLeaf and CjTranscriptome unigenes in three categories are displayed in Fig. S1.

Based on the BLAST searches against the four protein databases, the coding regions of 37,111, 41,059, 40,029 and 54,155 unigenes were separately extracted from ChFlower, CjFlower, CjLeaf and CjTranscriptome. Moreover, 1248 unigenes in ChFlower, 1390 unigenes in CjFlower, 1187 unigenes in CjLeaf and 1607 unigenes in CjTranscriptome with CDS were gained according to the ESTScan methods (Table 2), and the most abundant size class was 200–500 bp for each transcriptome (Table S2).

3.3. Orthologous pairs detection, Ka/Ks ratios, and mining of SCNGs

After filtering the preliminary orthologous pairs based on the three criteria (see before), we detected 11,566 potential orthologous groups between ChFlower and CjFlower, and proceeded to estimate the Ka, Ks and Ka/Ks values of these groups (Table S3). The mean values of the Ka, Ks and Ka/Ks ratios were 0.007, 0.024 and 0.670, respectively. Although there are 827 orthologous pairs with Ka/Ks ratios >1, only six pairs showed Ka/Ks ratios significantly >1 (P < 0.05), suggesting that these pairs are likely under positive selection. Taking a more appropriate threshold of 0.5 for the Ka/Ks ratio as an indicator of positive selection (Swanson et al., 2004; Zhang et al., 2013), another 13 orthologous pairs with Ka/Ks values between 0.5 and 1 and P values < 0.05 were identified (Table 3). The functions of these positively selected genes are described based on the best hits against the Nr database (Table 3).

Of the APVO SCNG dataset (Duarte et al., 2010) used to perform TBLASTN queries against orthologous pairs between ChFlower and CjFlower (see above), a total of 429 genes were found to have hits against our orthologous genes (Table S4), which are most likely the single copy nuclear genes in Croomia species.

3.4. Identification, validation and cross-species transferability of EST-SSRs

In this study, we identified a total of 16,384 sequences containing 21,460 EST-SSRs from the C. japonica (CjTranscriptome)
unigenes, with 3681 sequences containing more than one SSR. Among the 21,460 EST-SSRs, the most frequent repeat types were dinucleotide (10,169, 47.4%), followed by trinucleotide (6,789, 30.2%), and mononucleotide (2,871, 13.4%) (Fig. 2, Table S5). The dominant dinucleotide repeat types were AG/CT, accounting for 77.6% of repeats, followed by AT/AT (14.6%), and AC/GT (7%), while the CG/CG motifs (0.8%) were least abundant. Among the trinucleotide motifs, the most common repeats were CGG/CGG (32.0%), followed by AGG/CCT (21.1%), and AAG/CTT (14.0%) (Fig. 2, Table S5).

Of the 100 primer pairs (Table S6) selected for testing polymorphism, 46 primer pairs (46%) successfully produced PCR fragments with expected size, and 10 primer pairs (10%) exhibited polymorphism in all three Croomia species (Fig. S2). Therefore, these 10 polymorphic loci were further used to evaluate the diversity of all 240 Croomia individuals (Table 4). Across the 10
analyzed loci, the observed number of alleles ($N_A$) varied from 5 to 16, with a mean of 9.5 alleles per locus. The observed ($H_O$) and expected ($H_E$) heterozygosities ranged from 0.271 to 0.758 and from 0.303 to 0.897, respectively. The PIC values varied between 0.289 and 0.866, with an average of 0.678. In addition, 6 out of 10 loci showed significant deviation from HWE ($P < 0.001$) (Table 4), although no null alleles were found at these EST-SSR loci.

### Table 1

Summary statistics of the assembly results for C. heterosepala flowers (ChFlower), C. japonica flowers (CjFlower), C. japonica leaves (CjLeaf) and C. japonica flowers + leaves (CjTranscriptome).

| Characteristics | C. heterosepala | C. japonica |
|-----------------|-----------------|-------------|
| ChFlower        |                 |             |
| Raw reads       | 49 881 224      | 50 481 150  |
| Clean reads     | 43 683 202      | 44 355 666  |
| GC percentage   | 50.46           | 49.68       |
| Q20 percentage  | 97.60           | 97.60       |
| Total number of contigs | 76 976 | 77 344 |
| N50 length of contigs (bp) | 1452 | 1461 |
| Mean length of contigs (bp) | 916 | 951 |
| Total number of unigenes (bp) | 57 193 | 62 131 |
| Mean length of unigens (bp) | 1042 | 1044 |
| N50 length of unigenes (bp) | 1603 | 1737 |
| GC percentage (%) | 50.46 | 49.68 |
| 200–500 bp (% of total unigenes) | 23 157 (37.34%) | 22 099 (35.57%) |
| 500–1000 bp (% of total unigenes) | 13 622 (23.82%) | 14 847 (23.90%) |
| 1000–1500 bp (% of total unigenes) | 8 655 (15.13%) | 10 000 (16.10%) |
| 1500–2000 bp (% of total unigenes) | 6 076 (10.62%) | 7 053 (11.35%) |
| Functional annotations |                 |             |
| Nr               | 37 970 (66.39%) | 42 032 (67.65%) |
| Swiss-Prot       | 26 629 (46.56%) | 29 573 (47.60%) |
| KEGG            | 25 382 (44.38%) | 27 938 (44.97%) |
| COG             | 17 633 (30.83%) | 19 404 (32.23%) |
| GO              | 18 436 (32.23%) | 20 376 (32.80%) |
| ALL             | 39 919 (69.80%) | 44 209 (71.15%) |
| CDS annotations |                 |             |
| Homolog         | 37 111 (64.89%) | 41 059 (66.08%) |
| ESTscan         | 1248 (2.18%)    | 1390 (2.24%) |
| ALL             | 38 359 (67.07%) | 42 449 (68.32%) |

### Table 2

Annotation of unigenes from the transcriptomes of C. heterosepala flowers (ChFlower), C. japonica flowers (CjFlower), C. japonica leaves (CjLeaf), and C. japonica flowers + leaves (CjTranscriptome).

| Characteristics | C. heterosepala | C. japonica |
|-----------------|-----------------|-------------|
| ChFlower        |                 |             |
| Functional annotations |       |             |
| Nr               | 37 970 (66.39%) | 42 032 (67.65%) |
| Swiss-Prot       | 26 629 (46.56%) | 29 573 (47.60%) |
| KEGG            | 25 382 (44.38%) | 27 938 (44.97%) |
| COG             | 17 633 (30.83%) | 19 404 (32.23%) |
| GO              | 18 436 (32.23%) | 20 376 (32.80%) |
| ALL             | 39 919 (69.80%) | 44 209 (71.15%) |
| CDS annotations |                 |             |
| Homolog         | 37 111 (64.89%) | 41 059 (66.08%) |
| ESTscan         | 1248 (2.18%)    | 1390 (2.24%) |
| ALL             | 38 359 (67.07%) | 42 449 (68.32%) |

### Table 3

Candidate orthologs likely under positive selection in the transcriptomes C. heterosepala flowers (ChFlower) and C. japonica flowers (CjFlower), respectively.

| Gene ID of orthologous genes | Ka/Ks value | P-value (Fisher) | Description |
|-----------------------------|-------------|-----------------|-------------|
| ChFlower                    |             |                 |             |
| CL3372.Contig2              | 1.904       | 0.0027          | uncharacterized protein |
| CL7139.Contig1              | 26.629      | 0.0142          | RING-H2 finger protein ATLS8-like |
| Unigene30940                | 5.860       | 0.0142          | hypothetical protein Csa_7G372310 |
| Unigene1132                 | 2.262       | 0.0263          | | |
| Unigene9837                 | 4.382       | 0.0230          | extensin-2-like, partial |
| ALL                         | 39 919 (69.80%) | 44 209 (71.15%) | RING-H2 protein RAFL-like 19 |
| CDS annotations             |             |                 |             |
| Homolog                     | 37 111 (64.89%) | 41 059 (66.08%) | probable disease resistance protein Atg122280 |
| ESTscan                     | 1248 (2.18%) | 1390 (2.24%) | uncharacterized protein LOC105055788 |
| ALL                         | 38 359 (67.07%) | 42 449 (68.32%) | protein RAFL-like 19 |

### 3.5. Identification, phylogeny, and evolutionary analysis of CYC/TB1 genes

In total, 10 and 11 TCP proteins with an open reading frame (ORF) exceeding the TCP domain were retrieved from ChFlower and CjFlower, respectively, through the first strategy (Supplementary data). The Neighbor-Joining (NJ) tree constructed from multiple
amino acid alignment of the TCP domain sequences classified the TCP proteins into two distinct clades (Class I and Class II), however, all the Class II transcripts presented in subclade CIN-like genes with none of the transcripts belonging to the Class II TCP proteins into two distinct clades (Class I and Class II), however, first and second strategies will be discussed below. Through the third strategy, the single amplification product with a length of 417 bp for the remaining branches, which were considered to be under positive selection. Furthermore, SteTBL1 genes supported by all these three best models (H4, H8, and H12 of the M2 model) were under strong purifying selection with values ranging from 0.1402 to 0.1523, as expected for a gene that codes for a functionally important protein.

### 4. Discussion

#### 4.1. RNA-Seq analysis of *C. heterospera* and *C. japonica*

Since next-generation sequencing (NGS) technologies appeared on the market ten years ago, dramatic increases have been made in terms of speed, read length and throughput, along with a sharp reduction in per-base cost, which paved the way for the wide application of NGS technologies in basic science (Dijk et al., 2014). Illumina-based transcriptome sequencing technology has proved to be a powerful tool for obtaining large amounts of transcriptome data, and is widely used in gene discovery (Clark et al., 2010; Mao et al., 2016), molecular marker development (Huang et al., 2014; Ai et al., 2015), phylogenomic analysis (Teasdale et al., 2016), differential gene expression (Akashi et al., 2016) and so on. In this study, we presented the first reference transcriptome (*CjTranscriptome*) in *Croomia* and the first comparative analysis of flower transcriptomes in Stemonaceae, for which, no genomic resources have been reported to date.

Through Illumina transcriptome sequencing analysis of *CjFlower*, *CjLeaf*, and *CjStem*, we obtained 57,193—64,448 unigenes with an average length of 1042—1106 bp and an N50 of 1575—1737 bp for these three samples (Table 1). Our results closely resembled the transcriptome assembly outcome of other monocots using similar technologies (Huang et al., 2015, 2016). By pooling the *CjFlower* and *CjLeaf* unigenes into a *CjTranscriptome*, we obtained the longer unigenes of *CjTranscriptome* than those of *CjFlower* and *CjLeaf*, possibly due to a relative increase of read coverage. For each transcriptome, more than 67% of the unigenes had significant Blast similarities.
hits against the protein databases, however, a fraction of unigenes from our study had no BLAST matches to these databases. This phenomenon is common in many other plants and these “non-BLASTable” genes were thought to be Croomia-specific genes, rapidly evolving genes, or untranslated regions (UTRs) (Parchman et al., 2002; Rudall et al., 2005). However, the detailed picture of evolutionary biology, and phylogenomic studies.

4.2. Orthologous genes and SCNGs between ChFlower and CjFlower

To evaluate the effects of selection on these two Croomia species with different flower symmetry types, the Ka/Ks ratio, an indicator of selective pressure on protein-coding genes, was calculated between ChFlower and CjFlower orthologs, only six orthologs had a Ka/Ks ratio significantly > 1, and an additional 13 pairs were above the relaxed threshold of 0.5 (Table 3), suggesting that these genes are likely under the effect of positive selection and may be related to the adaptive divergence of these two species. According to the N annotation of these positively selected genes, more than half were involved in several biological functions (e.g., disease resistance, RNA-binding), while the remaining seven were recognized as uncharacterized or hypothetical proteins. Thus, further studies on expression analysis, cellular localization, molecular interactions and three-dimensional structures are needed on these uncharacterized proteins (Schlupean et al., 2013).

The identification of orthologous genes is a crucial prerequisite for reliable phylogenetic investigation (Teasdale et al., 2016), and the advantages of single or low copy nuclear genes for resolving deep evolutionary relationships have been proven at the species level to the order level (Duarte et al., 2010; Ai et al., 2015; Xiang et al., 2016; Zeng et al., 2017). Previous phylogenetic analyses using sequence data from a few chloroplast genes and nrDNA ITS, as well as morphological characters, revealed that Stemona was monophyletic; Pentastemonata firstly diverged from the remaining genera with Stemona sister to (Croomia + Stichonereon) (Caddick et al., 2002; Rudall et al., 2005). However, the detailed picture of species relationships within each genus has remained unclear due to too few genetic markers employed in previous studies. In this study, the 429 putative SCNGs identified in the transcriptomes of Croomia species may represent more desirable choices for species-level phylogenetic reconstruction for Croomia and its related genera. Moreover, since multiple unlinked genetic loci provide independent realizations of divergence history, accounting for mutational and coalescent stochasticity (McCormack et al., 2011; Leavitt et al., 2012), these putative SCNGs are suitable candidates for studies aimed at understanding molecular phylogeography and population genetics in Croomia.

4.3. EST-SSR markers

Unlike SSRs, EST-SSR markers are easier to obtain, more widely transferable among species, and related to phenotypic variation (Andersen and Lübbertstedt, 2003; Li et al., 2004; Duran et al., 2009). Therefore, in recent years, EST-SSR markers have been developed in many species for genetic analysis (Poncet et al., 2006; Qiu et al., 2010; Chen et al., 2015). However, prior to this study, neither ESTs nor EST-SSR markers were available on public databases for medicinally important, endangered species of Croomia. This lack of genomic resources has impeded our understanding of Croomia conservation, phylogeography, and population genetics. Of the 21,460 EST-SSRs identified in our work, dinucleotide (47.4%) repeats were the most frequent SSR repeat type, followed by trinucleotide (30.2%), and mononucleotide (13.4%) repeats (Fig. 2, Table S5). This result is in contrast to most results observed in previous studies, in which tri-nucleotide repeats were the most abundant EST-SSR repeat motif (Varshney et al., 2005; Koikinda et al., 2012; Guo et al., 2014; Mao et al., 2016). Morgante et al. (2002) investigated the distribution of SSRs across different genomic fractions and found that, excluding tri-nucleotides, SSRs rarely occurred in coding regions compared with non-coding regions. Therefore, the dominance of di-nucleotide repeats detected here may be caused by an over-representation of untranslated regions (UTRs) compared with open reading frames (Kumpatla and Mukhopadhyay, 2005; Chen et al., 2015). Recently, 126 genomic SSR primer pairs were designed and only 11 (8.7%) primer pairs showed interspecies transferability among the three Croomia species (Fang et al., 2013). In this study, of the 100 primer pairs, 46 (46%) primer pairs successfully produced PCR fragments of expected size, suggesting EST-SSRs are more transferable than genomic SSRs in Stemonaaceae. Moreover, another important advantage of EST-SSRs is the possibility to track adaptive divergence processes for their functional involvement (Ai et al., 2015). Thus, these newly identified EST-SSR markers will be powerful tools for future population genetics across a wide range of taxa in Stemonaaceae.

4.4. Analysis of CYC/TB1-like genes in Stemonaaceae species

Zygomorphy has evolved repeatedly from actinomorphy throughout angiosperms (Zhang et al., 2010). During the evolution of angiosperms, floral zygomorphy has evolved independently at
least 130 times, while reversal to actinomorphy may have evolved at least 69 times. Among the origins of floral zygomorphy, 29 cases have been in monocots (Reyes et al., 2016). CYC/TB1-like genes, which are associated with the evolution and maintenance of flower monosymmetry have not been reported in Stemonaceae. Previous phylogenetic analyses of the reduced CYC/TB1 dataset resolve two clades of TBL genes from monocots, named TBL1 and TBL2 (Bartlett and Specht, 2011). This ancient duplication in the TBL gene lineage is found to predate the divergence of the commelinid monocots (Bartlett and Specht, 2011). The first two strategies of our study (see Materials and Methods) failed to obtain CYC/TB1-like genes in both flower and leaf transcriptomes, suggesting that these genes are not expressed, or are expressed at an extremely low level at the stages of anthesis and leaf. The expression patterns of CYC/TB1-like genes are found to vary across different plant species. For example, asymmetric expression of TB1a in Commelina communis and C. diandrophila (zygomatic flowers) is initiated in early flower development (Preston and Hileman, 2012), while asymmetric expression of CYC/TB1-like genes in the only zygomatic magnolid, Aristolochia, is only detected after the establishment of perianth zygomorphy at later stages of floral development (Horn et al., 2015). Future studies should detect and compare the expression patterns at different stages (primordium, 0.5 cm buds, 1.0 cm buds etc.), and using dissected tissues (lateral petals, carpels and stamens etc.). Even though we have not detected CYC/TB1-like genes from the transcriptomes of Croomia, based on PCR-based methods using different sets of TBL primers (Howarth and Donoghue, 2005), we have isolated a single copy of the CYC/TB1-like gene from each of the Stemonaceae taxa surveyed. The resulting tree topology (Fig. 3) strongly supports the monophyly of the 7 SteTBL sequences (PP = 1.00), which clearly belong to the TBL1 clade (Bartlett and Specht, 2011). This phylogenetic pattern provides additional evidence for the homology of these sequences by representing orthologous genes in Stemonaceae that descended from a single ancestral copy. The association between CYC-like copy number and floral symmetry has been discovered in the Dipsacales (Howarth and Donoghue, 2005) and Plantaginaceae (Reardon et al., 2009). In fact, the TBL1 gene is also found to be single copy in a large number of grasses, e.g., Oryza, Sorghum, and Brachypodium (Lukens and Doebley, 2001). The single TBL gene in Stemonaceae and Poaceae differs from the inferred homologs of TB1 in the Zingiberaleans, ZntTB1a and ZntTB1b, which have both been maintained in the Zingiberaleans genome following a duplication event in the TBL1 gene lineage (Bartlett and Specht, 2011). Taken together, the shifts of floral symmetry in Stemonaceae do not appear to be correlated with TBL copy number. Within the TBL1 clade, multiple sequences are clustered in different subclades, and generally grouped according to taxonomy (e.g., Stemonaceae, Poaceae, Asteroermeriaceae). This phylogenetic pattern suggests that a common ancestral copy of TB1 has undergone significant diversification following the diversification of these families. On the other hand, these results imply that functional diversification of the TBL genes may play an important role in the diversification of these families, resulting in a new protein function that was maintained through extreme purifying selection acting on the TBL gene (Bartlett and Specht, 2011).

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Appendix A. Supplementary data

Supplementary data for this article can be found online at https://doi.org/10.1016/j.pld.2018.11.006.

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