Genome analysis of *Hibiscus syriacus* provides insights of polyploidization and indeterminate flowering in woody plants

Yong-Min Kim¹,†, Seungill Kim²,†, Namjin Koo¹,†, Ah-Young Shin³,†, Seon-In Yeom⁴,†, Eunyoung Seo², Seong-Jin Park¹, Won-Hee Kang⁴, Myung-Shin Kim², Jieun Park², Insu Jang¹, Pan-Gyu Kim¹, Iksu Byeon¹, Min-Seo Kim¹, JinHyuk Choi¹, Gunhwan Ko¹, JiHye Hwang⁵, Tae-Jin Yang², Sang-Bong Choi⁶, Je Min Lee⁷, Ki-Byung Lim⁷, Jungho Lee⁹, Ik-Young Choi⁹, Beom-Seok Park⁵, Suk-Yoon Kwon³, Doil Choi², and Ryan W. Kim¹,*

1Korean Bioinformation Center, Korea Research Institute of Bioscience and Biotechnology, Daejeon 34141, Korea, 2Department of Plant Science, College of Agriculture and Life Sciences, Seoul National University, Seoul 08826, Korea, 3Plant Systems Engineering Research Center, Korea Research Institute of Bioscience and Biotechnology (KRIIBB), Daejeon 34141, Korea, 4Department of Agricultural Plant Science, Institute of Agriculture and Life Science, Gyeongsang National University, Jinju 52828, Korea, 5National Institute of Agricultural Sciences, Rural Development Administration, Jeonju 54875, Korea, 6Division of Bioscience and Bioinformatics, Myongji University, Yongin 17058, Korea, 7Department of Horticultural Science, College of Agriculture and Life Science, Kyungpook National University, Daegu 41566, Korea, 8Green Plant Institute, Yongin 446-908, Korea, and 9Department of Agriculture and Life Industry, Kangwon National University, Chuncheon 24341, Korea

*To whom correspondence should be addressed. Tel: 82-42-870-8500. Fax: 82-42-870-8519. Email: rwkim@kirbb.re.kr

†These authors contributed equally to this work.

Abstract

*Hibiscus syriacus* (L.) (rose of Sharon) is one of the most widespread garden shrubs in the world. We report a draft of the *H. syriacus* genome comprised of a 1.75 Gb assembly that covers 92% of the genome with only 1.7% (33 Mb) gap sequences. Predicted gene modeling detected 87,603 genes, mostly supported by deep RNA sequencing data. To define gene family distribution among relatives of *H. syriacus*, orthologous gene sets containing 164,660 genes in 21,472 clusters were identified by OrthoMCL analysis of five plant species, including *H. syriacus*, *Arabidopsis thaliana*, *Gossypium raimondii*, *Theobroma cacao* and *Amborella trichopoda*. We inferred their evolutionary relationships based on divergence times among Malvaceae plant genes and found that gene families involved in flowering regulation and disease resistance were more highly divergent and expanded in *H. syriacus* than in its close relatives, *G. raimondii* (DD) and *T. cacao*. Clustered gene families and gene collinearity analysis revealed that two recent rounds of whole-genome duplication were followed by diploidization of the *H. syriacus* genome after speciation. Copy number variation and phylogenetic divergence indicates that WGDs and subsequent diploidization led to unequal duplication and deletion of flowering-related genes in *H. syriacus* and may affect its unique floral morphology.
1. Introduction

*Hibiscus syriacus* (*L.*) (rose of Sharon) is a fast-growing deciduous shrub of the Malvaceae family, which includes species such as *Gossypium raimondii* and *Theobroma cacao*. Although its name indicates this species was first identified in Syria, *H. syriacus* likely originated from the Korean peninsula and southern China and has since spread to Western countries. In temperate zones, *H. syriacus* is a commonly grown ornamental species with attractive white, pink, red, lavender, or purple flowers displayed over a long blooming period, though individual flowers last only a day. Its Korean name, *Mugunghwa*, literally means ‘flowering forever’. In addition to its ornamental value, *H. syriacus* acts as an ozone bioindicator, and its dried flowers and root bark are used in Oriental herbal medicines. Specifically, a novel cyclic peptide (Hisheptin A) and three naphthalene compounds (syriacins A-C) isolated from the plant’s root bark have been used as anti-pyretic, anti-helminthic and anti-fungal agents.

Polyplody is a well-established influence on plant genome evolution but is now recognized as a common phenomenon in diverse eukaryotes, as signs of whole-genome duplication (WGD) have been detected in many sequenced genomes. Recent genome analysis demonstrated that most eudicot plants descended from an ancient hexaploid ancestor and followed lineage-specific polyploidization and that two rounds of WGD occurred in ancestral vertebrates. In general, changes in ploidy are expected to be deleterious and an ‘evolutionary dead end’ for many species. However, polyploidization of plants mediated their survival during the Cretaceous-Tertiary extinction event by increasing their genetic diversity. Each round of polyploidization was followed by many gene deletions (homeolog gene loss), interchromosomal rearrangements, neo-functionalization, and sub-functionalization. In Malvaceae plants, *Gossypium* includes five tetraploid taxa (*AD*; AD, 2n = 4x) and 45 diploid taxa (2n = 2x). Among them, *G. raimondii* (DD, D-genome), *G. arboreum* (AA, A-genome) and *G. hirsutum* (AD, D-genome) genomes were reported. *Hibiscus* also includes many polyploid species, such as *H. syriacus* (2n = 4x = 80), *H. aspera* (2n = 8x = 72), and *H. rosa-sinensis* (2n = 16x = 144) and diploid species (*H. pedunculatus* (2n = 2x = 30) and *H. phoenceus* (2n = 2x = 22)).

Here, we report the genome sequence of *H. syriacus* and the possible correlation between polyploidization and its phenotypes. Comparative genomic analysis of Malvaceae species, including *H. syriacus*, *T. cacao*, and *G. raimondii* (DD), provides clues of the recent polyploidization in *H. syriacus* by WGDs and unequal regulation of gene dosage by subsequent paleopolyploidy. Our investigation of copy number variations of *H. syriacus* were reported. *Hibiscus* also includes many polyploid species, such as *H. syriacus* (2n = 4x = 80), *H. aspera* (2n = 8x = 72), and *H. rosa-sinensis* (2n = 16x = 144) and diploid species (*H. pedunculatus* (2n = 2x = 30) and *H. phoenceus* (2n = 2x = 22)).

2. Materials and methods

2.1. Plant materials and whole genome sequencing

Leaves of *H. syriacus* plants >100-years-old and nominated as National Monument of Korea trees (serial number 520) were harvested and frozen immediately in liquid nitrogen. Genomic DNA for Paired-end (PE) and Mate-pair (MP) libraries was extracted, and libraries for next-generation sequencing were constructed according to the manufacturer’s instructions (Illumina, San Diego, CA, USA). The quality of each library was validated using the KAPA SYBR FAST Universal 2× qPCR Master Mix (Kapa Biosystems, Boston, MA, USA). Each library was sequenced with the Illumina HiSeq 2000 platform.

2.2. Genome assembly, scaffolding and gap-closing

Genome assembly was performed using both Platanus v1.2.14 and SSPACE v2.0. To generate longer initial contigs, single reads merged using FLASH v1.2.26 and reads from the PE libraries were assembled using Platanus with parameters to resolve heterozygosity in the *H. syriacus* genome (-u 0.2 -c 5 -d 0.3 -m 460). The scaffolding process was performed with Platanus and SSPACE. We first determined mapping seed length for scaffolding and then generated longer scaffolds using optimized Platanus parameters (-1.5 -s 41 -u 0.3). To extend scaffold length, SSPACE fulfilled serial scaffolding with hash parameters for the scaffolds generated by Platanus. Lastly, remaining gaps were filled with Platanus and GapCloser version 1.10 (http://soap.genomics.org.cn/down/GapCloser_release_2011.tar.gz) using reads from the PE and MP libraries.

2.3. Genome annotation

Annotation of the *H. syriacus* genome was performed using the KOBIC annotation pipeline (modified PGA pipeline) consisting of repeat masking, mapping of different protein sequence sets, and *ab initio* prediction performed by AUGUSTUS v3.0.3. The protein sequences of *A. thaliana* (TAIR10, http://www.arabidopsis.org), *T. cacao* and *G. raimondii* were mapped using GeneWise v2.1 to generate protein-based gene models for consensus modeling. AUGUSTUS was used for gene prediction in *H. syriacus*. Then predicted gene models from AUGUSTUS were validated using BLASTp with protein sequences from the three genomes (*T. cacao, G. raimondii* and *A. thaliana*) as queries and erratic gene models were filtered with a BLASTp cut-off value of query coverage ≥ 0.3. The predicted gene models from GeneWise were also filtered using query coverage ≥ 0.3. Remaining gene models of GeneWise depicted as GeneWise format were reformatted as GFF3 data and used to determine the consensus gene model via EVidenceModeler (EVM), which combines *ab initio* gene predictions with protein alignments into weighted consensus gene structures (*ab initio* predictions = 1, protein alignment = 5, transcript alignment assemblies = 7). Biological functions of the final gene models were assigned using InterPro, plant protein sequences in the RefSeq23 and UniProt databases, which includes SWISS-PROT and TrEMBL data as described in previous study. For functional annotation, three quality criteria were concerned: (i) bit score of the BLAST result is >50 and e-value is <e−10; (ii) subject coverage of the BLAST result is >60%; and (iii) top token score from lexical analysis is >0.5. To infer function for the protein-coding genes, we used InterProScan version 5.4 to scan protein sequences against the protein signatures from InterPro.

Key words: *Hibiscus syriacus*, Whole Genome Duplication, Diploidization, Multivoltinism, Homeolog
2.4. RNA sequencing and de novo transcriptome assembly

Total RNA was extracted from plant leaves, petals, ovaries, and roots using TRriol reagent (Invitrogen, CA, USA) following the manufacturer’s instructions. RNA-Seq libraries were generated using purified total RNA and sequenced using an Illumina HiSeq 2000 system. Thirty-six gigabases of raw reads were generated and preprocessed using DynamicTrim and LengthSort in SolexaQA.26 The preprocessed raw reads were then used for transcriptome assembly and DEG analysis. Velvet v1.2.07 was used to assess k-mer sizes and assembled contigs, which were then merged using Oases v0.2.08. Assembled transcripts were validated using BLASTx (e-value < e−10, best hit) against 1,917,424 protein sequences from 39 plant genomes selected from each family including Arabidopsis thaliana, Brassica rapa, Solanum lycopersicum, Solanum phureja, T. cacao, G. raimondii, Oryza sativa, Zea mays, Cucumis sativus, Vitis vinifera.

2.5. Evaluation of genome assembly

For validation of the assembled genome sequence, CEGMA (Core Eukaryotic Genes Mapping Approach) v2.527 and BUSCO (Benchmarking Universal Single-Copy Orthologs) v1.2.28 were used in H. syriacus genome using default parameters. The CEGMA mapped a gene structure to new genomic sequence using a set of highly conserved protein family in eukaryotes by Hidden Markov Model. We evaluated 248 core eukaryotic genes defined by CEGMA to our genome sequence. The BUSCO provides completeness assessment of assembled genome based on orthologous gene sets with single copy from OrthoDB (http://www.orthodb.org) using hidden Markov model for profile of amino acid alignment. For BUSCO assessments, we used 429 gene sets of conserved orthologs in eukaryotes.

2.6. Detection of gene families in the H. syriacus genome

OrthoMCL v2.0.219 was used to identify gene family clusters in H. syriacus and the other four sequenced genomes which are G. raimondii, T. cacao, A. trichopoda and A. thaliana (In the first step, a set of high quality of gene models was obtained by rejecting low-quality sequences based on default parameters in OrthoMCL. The default parameters of rejecting low quality protein sequences were (i) shorter than 10 amino acids (ii) >20% stop-codons (iii) >20% non-standard amino acids. Pairwise sequence similarities between all input protein sequences were calculated by all-by-all BLASTp with an e-value cut-off of 1e−05 and a minimum match length of 50%. To define ortholog cluster structure, a Markov clustering algorithm was applied with an inflation value (−I) of 1.5 (default value in OrthoMCL). Putative splice variants were removed from the data set; longest protein sequences were kept and subsequently filtered for premature stop codons and incompatible sequences.

2.7. Detection of collinearity blocks in Malvaceae plants

MCScanX30 was used to construct synteny and collinearity blocks between H. syriacus and G. raimondii against T. cacao.19 First, homologous gene pairs were identified using protein sequences from the three genomes and then scanned inter- and intra-species by BLASTp (options with −e 1e−10 −b 5 −v 5). The BLASTp output was used with merged GFFs of three species to perform MCscanX with default parameters. We generated gene synteny and collinearity data to align proteins of the two species against reference chromosome of T. cacao. Collinearity blocks containing fewer than five proteins were excluded. To search a candidate of duplicated regions, we made the groups of collinear block from multiple collinear blocks which have similar protein members (>80%), and the same chromosome in T. cacao. Then, each block in H. syriacus and G. raimondii was counted by overlap against the cluster blocks of T. cacao. The duplicated regions in H. syriacus and G. raimondii were identified, if the number of blocks was more than two.

2.8. Estimation of speciation time in Malvaceae plants

To construct a phylogenetic tree of the five species (A. trichopoda, A. thaliana, G. raimondii, T. cacao and H. syriacus), we extracted 941 single-copy gene sets from all genomes in the OrthoMCL clusters. We performed multiple alignments of the CDSs of each gene set using Prank (−f = nexus -codon).31 The alignment file was used to construct a phylogenetic tree based on calculations of divergence time for the five species.32 For accurate tree construction, we assigned taxon sets based on previously calculated speciation of A. thaliana, G. raimondii and T. cacao. The Bayesian software package BEAST12 (v1.8.2) was used to estimate divergence times and construct the final tree. The Markov chain Monte Carlo (MCMC) analyses in BEAST was conducted for 10 million generations with samples every 1,000 steps and the effective sampling size was over 150 for all of parameters. We used SRD0633 as a substitution model and the Yule process34 as a traditional speciation model.

2.9. Identification of TF candidates

We identified TF candidates as previously described.17 Briefly, predicted proteins containing TF domains were screened by InterProScan32 search against Pfam35 databases. The TF candidates were classified based on rules as indicated at PlnTFDB (http://plntfdb.bio.uni-potsdam.de/v3.0/downloads.pl). InterProScan search against Pfam (v1.8.2) was used to detect TFs, followed by HMMER.37 The assigned TF candidates were confirmed by BLASTp against plant TF protein sequences downloaded from PlnTFDB and Hidden Markov Model profiles were built and screened using HMMER.37 The assigned TF candidates were confirmed by BLASTp against plant TF protein sequences downloaded from PlnTFDB (http://plntfdb.bio.uni-potsdam.de/v3.0/downloads.php).

2.10. Identification of genes encoding nucleotide-binding site proteins

To identify nucleotide-binding site (NBS)-encoding genes, representative genes from each plant genome were screened using the raw Hidden Markov Model (HMMER3.0)38 to search for the Pfam NBS family PF00931 domain (e-value cut-off of 1.0). All putative NBS protein sequences were analysed and manually curated by a BLASTp search against known R gene sequences in GenBank. To further identify TIR homologs and sequences encoding CC and LRR motifs, candidate NBS-LRR protein sequences were characterized using SMART,39 the Pfam database40 and the COILS programme41 with a threshold of 0.9 to detect CC domain specifically.

3. Results

3.1. Genome sequencing and assembly

H. syriacus plants over 100-years-old were selected for genome sequencing. Illumina whole-genome shotgun sequencing generated 233.3 Gb (122.8× coverage) of genomic sequences (Supplementary
Table 1. Summary of *H. syriacus* genome assembly

| Parameter                                      | Value               |
|------------------------------------------------|---------------------|
| Number of scaffolds                            | 77,492              |
| Total length of scaffolds                      | 1,748 Mb            |
| N50 of scaffolds                               | 140 kb              |
| Longest (shortest) length of scaffolds         | 1.54 Mb (500 bp)    |
| Number of contigs                              | 172,672             |
| Total length of contigs                        | 1,715 Mb            |
| N50 of contigs                                | 30.0 kb             |
| Longest (shortest) length of contigs           | 643 kb (87 bp)      |
| Number of gap sequences                        | 33 Mb (1.9%)        |
| GC content                                     | 34.04%              |
| Total size of TEs                              | 1,095 Mb (57.6%)    |

Table S1. PE libraries (250–500 bp) were generated, and 2 kb and 5 kb MP libraries were sequenced with a read length of 101 or 151 bp. Pre-processing analysis of raw sequences was performed to remove extraneous sequences for accurate genome assembly as described in the previous report. After filtering, 156.6 Gb (82.4× coverage) of *H. syriacus* genome sequences were used for further analysis (Supplementary Table S2). K-mer distribution analysis, which provides information related to low frequencies, sequencing depth, level of heterozygosity, and genome size was then applied using Jellyfish (Supplementary Fig. S1). The estimated genome size of 1,901 Mb was calculated by dividing the total volume by the peak of distribution as described previously.

Validation of the assembled genome was performed using 128,888 representative transcripts derived from de novo assembly of a combined transcriptome from all libraries (Supplementary Table S3). To confirm sequence alignment between transcriptome assembly and scaffolds, we performed BLAST comparisons for the transcriptome assembly and scaffolds as queries and subjects, respectively. We found that 117,431 (91.1%) assembled transcripts as query sequences matched to scaffolds with 98% identity. In addition, 93,688 (79.8%) transcripts matched to genome sequence with query coverage over 80%, and 82,394 (70.2%) transcripts matched over 90% coverage by the assembled scaffolds (Supplementary Fig. 2). The quality of the assembly was also evaluating using CEGMA (Core Eukaryotic Genes Mapping Approach) and BUSCO (Benchmarking Universal Single-Copy Orthologs). These analyses showed 92.74% of completeness (230 of 248 CEGs) from CEGMA and 92% of complete BUSCOs. These results suggested that *H. syriacus* genome assembly was high quality. As a result, 1,748 Mb (91.9% of 1,901 Mb) of genomic sequences were assembled into 77,492 scaffolds. The assembled genome was comprised of 33 Mb (1.9%) gap sequences and 1,715 Mb of contigs with N50 = 30 kb (Table 1 and Supplementary Table S4).

3.2. Genome annotation

Annotation of the *H. syriacus* genome was performed using the KOBIC Genome Annotation pipeline (Supplementary Fig. S3), including masking repetitive sequences, transcriptome mapping, reference protein mapping using GeneWise, *ab initio* gene prediction, and determination of consensus gene models using EVM.

Before masking repetitive sequences, repeat annotation was performed by RepeatModeler and RepeatMasker (http://www.repeatmasker.org) for the assembled genome. Due to a lack of repeat sequence information for this genome, we constructed a *de novo* repeat library using RepeatModeler, and RepeatMasker was applied for annotation of the constructed repeat library. Repeat sequences, except for unknown transposable elements (TEs), were masked so we could identify essential gene families, such as those that encode receptor-like kinases and nucleotide-binding proteins. TEs comprised 1,095.8 Mb (57.6%) of the genome (Supplementary Table S5) and mostly included long terminal repeats (LTRs), which accounted for approximately 30% of total TEs. Gypsy and Copia retrotransposons were the most common LTRs detected.

Transcripts mapping was performed using TopHat and Cufflinks, and protein alignment was performed by GeneWise. The protein sequences of *A. thaliana* (TAIR10), *T. cacao* and *G. raimondii* were mapped to generate protein-based gene models. For annotation of duplicated genes or gene families, mapping regions of a reference protein in the *H. syriacus* genome were determined from tBLASTn (default e-value 10) results using custom Perl scripts. These steps prevent mis-annotation of duplicated genes due to lack of mapping data for reference proteins from parsing single best-matched regions in the *H. syriacus* genome. We annotated 87,603 genes using KOBIC annotation prediction with an average CDS length of 1,188 bp, similar to that for *G. raimondii* (Table 2). Consensus gene models were evaluated using 88.4 Gb of Illumina-derived RNA-seq data. Overall, 91.76% of the predicted coding sequences were supported by Illumina data, demonstrating the high accuracy of KOBIC annotation prediction. The *H. syriacus* genome contains two times more genes than *G. raimondii* and three times more genes than *T. cacao* (Table 2), suggesting a polyploid genome as first indicated in a previous report.

We also performed OrthoMCL analysis to detect orthologous genes among Malvaceae plants, *A. thaliana*, and *Amborella trichopoda*. We identified 21,472 orthologous gene sets containing 164,660 genes, 5,300 of which were *H. syriacus*-specific (Fig. 1). Interestingly, these genes were the number of gene was three times larger than those of *G. raimondii* and *T. cacao*, further indicating *H. syriacus*’ polyploidy. In addition, relatively large numbers of singletons in the genomes of *A. thaliana* and *A. trichopoda* suggest that the Malvaceae lineage diverged long ago and now shows a high degree of evolutionary distance from other eudicots. For further analysis, estimation of speciation time and comparison of genome structures among Malvaceae plants were performed using paired gene sets.

3.3. Genome structure and polyploidization of *H. syriacus*

To compare genome structures among Malvaceae plants, collinearity blocks were detected using M CSca nX. Two WGDs or a triplication event have occurred in *G. raimondii* (DD), while none have occurred in *T. cacao*. Therefore, the *T. cacao* genome was used as a template to detect collinearity blocks in *G. raimondii* and *H. syriacus* (Fig. 2A). We detected *T. cacao* collinearity blocks in *G. raimondii* and *H. syriacus* with frequencies ranging from 2 to 7. The *H. syriacus* genome contains four times as many collinearity blocks than *G. raimondii* and blocks two times larger, indicating WGD events in *H. syriacus*. Duplication patterns were identified using phylogenetic analyses, which revealed single-copy flowering regulator genes in the diploid genomes of *A. thaliana*, *A. trichopoda* and *T. cacao* (Fig. 2B and Supplementary Fig. S4). Duplication of the GIGANTEA (GI) gene indicated WGDs occurred three times in *H. syriacus* but that many descendant genes since the first WGD have been lost (Fig. 2B), such as the CONSTANS and SOC1 genes (Supplementary Fig. S4). Thus, diploidization and homeolog loss in *H. syriacus*, first proposed
in previous studies, included duplication of distinct, individual gene families stemming from random homeolog gene loss after each WGD. Paleohexaploidy has occurred in the G. raimondii genome, and duplication patterns we observed were consistent with these previous results.

To estimate divergence time among Malvaceae plants, we calculated synonymous substitution rates (Ks) and constructed phylogenetic trees via the BEAST package using single-copy genes in OrthoMCL clusters. The trees revealed that the Malvaceae family diverged from a Brassicae-Malvaceae common ancestor approximately 91.91 MYA (Fig. 2C) and that H. syriacus, G. raimondii and T. cacao belong to a common subclade that diverged from a common ancestor approximately 30.88 MYA, which corroborates earlier studies. Occurrence of duplications in G. raimondii genes ranged from 24.46 to 45.46 MYA (Fig. 2B), while H. syriacus individual gene duplications before speciation and WGD events ranged from 25.23 to 48.23 MYA and from 4.61 to 21.15 MYA, respectively (Fig. 2B and C). These results suggest that one WGD occurred in H. syriacus before speciation and two WGDs occurred after speciation.

Previous reports indicate transcriptional factors (TFs) were retained as duplicated genes, while other genes remained singletons. We investigated the duplication status of TFs in H. syriacus, and other Malvaceae plants and identified 9,642 TFs and transcriptional regulators in 81 families in the H. syriacus genome. Eighteen H. syriacus TF gene families, including AP2-ERF, AUX/IAA, and FAR1, contained more genes than those in diploid genomes (Supplementary Table S6). In particular, the H. syriacus genome contains 10 times more FAR1 family genes than the other genomes we analysed, although 19 TF genes showed convergent evolution patterns, and the proportions of other major TF family genes were similar across species. Thus, complex WGD events followed by diploidization led to unequal regulation of gene dosage and caused gene family copy number variations in H. syriacus.

### 3.4. Flowering-time and disease-resistance genes in H. syriacus

Genetic and molecular mechanisms of floral development in different plant species is highly conserved and include four major flowering pathways (photoperiod, autonomous, vernalization and gibberellin) well-characterized in A. thaliana. Main flowering signals are regulated by the FLOWERING LOCUS T (FT) in the photoperiod pathway, while the vernalization pathway acts via removal of an FT repressor after exposure to certain stimuli. H. syriacus is a long-day flowering plant with a long blooming period and can express a multivoltinism phenotype with 20–30 blossoms per day. However, the flowers of H. syriacus open daily and last for only one day. To uncover the genetic mechanisms controlling these phenotypes, we investigated genes involved in the four major flowering pathways of

| Protein-coding loci | Total CDS length (bp) | Avg. CDS length (bp) | Avg. Exon length (bp) | Avg. Intron length (bp) |
|---------------------|-----------------------|----------------------|-----------------------|------------------------|
| H. syriacus         | 87,603                | 104,087,809          | 1,188                 | 239                    | 383                    |
| T. cacao*           | 28,798                | 33,494,538           | 1,857                 | 231                    | 502                    |
| G. raimondiib       | 40,976                | 45,237,504           | 1,104                 | 244                    | 339                    |
| A. thalianac        | 27,206                | 24,861,465           | 1,212                 | 265                    | 164                    |

* Cacao genome paper
b Cotton genome paper
c TAIR10 annotation (http://www.arabidopsis.org)

Figure 1. Distribution of orthologous gene families of H. syriacus, G. raimondii, T. cacao, A. trichopoda and A. thaliana, from which 169,570 sequences were clustered into 9,076 groups. The number of clustered groups and genes in each species are shown on the left and center, and total gene numbers are shown on the right.
flowering time is frequently dependent on gene copy number, we identified genes from *H. syriacus* that have over-represented copy numbers compared to those of *S. lycopersicum* and *A. thaliana*. These genes are divided into two subclasses based on the presence of tandem leucine-rich repeats (LRR) and the NBS domain. Our results suggest that *H. syriacus* may have undergone several polyploidization events in its evolutionary history.

**4. Discussion**

Polyplody is an important mechanism of plant speciation. It is characterized by increased genetic diversity, gene duplication, and altered chromosome pairings. The newly formed polyploids may experience rapid homeolog gene loss, genome reconstruction post-polyploidization, and altered patterns of gene expression.
Table 3. Comparison of flowering-time gene copy numbers

| Regulators Arabidopsis locus | H. syriacus | T. cacao | G. raimondii | A. trichopoda |
|-----------------------------|-------------|----------|---------------|---------------|
| CO  AT5G15840 6             | 1           | 1        | 0             |               |
| ELF4  AT2G40080 7           | 1           | 4        | 1             |               |
| FCA  AT4G16280 2            | 1           | 1        | 1             |               |
| FKF1  AT1G68050 4           | 1           | 2        | 1             |               |
| FLK  AT3G04610 3            | 1           | 2        | 1             |               |
| FT   AT1G65480 2            | 1           | 1        | 1             |               |
| GI   AT1G22770 5            | 1           | 2        | 0             |               |
| LFY  AT3G61850 4            | 1           | 1        | 1             |               |
| LHY  AT1G01060 7            | 1           | 3        | 1             |               |
| VIN3 AT5G37380 7            | 1           | 2        | 1             |               |
| SOCI AT2G45660 4            | 1           | 2        | 1             |               |
| TFL  AT5G03840 4            | 2           | 2        | 0             |               |
| SVP  AT1G24260 3            | 2           | 2        | 0             |               |
| PHYA AT1G09570 4            | 1           | 1        | 1             |               |
| PHYB AT2G18790 4            | 0           | 1        | 1             |               |
| PHYC AT5G35840 2            | 1           | 1        | 1             |               |
| PHYE AT4G18130 4            | 1           | 0        | 0             |               |

The flowering phenotype of *H. syriacus* is characterized by multivolinitism, a long blooming period, and high blossom turnover. We found that the copy numbers of most flowering-related genes, such as GIGANTEA, CONSTANS, and ELF4 (but not FT), were higher in *H. syriacus* than in the diploid genomes of *T. cacao*, *A. trichopoda*, and *A. thaliana*. In addition, FAR1 genes, which modulate phytochrome A signaling by directly activating transcription of FHY1 and FHL and lead to accumulation of nuclear phytochrome A, were significantly increased in *H. syriacus*. FAR1 regulates the circadian clock, and its high copy number could directly affect the flowering phenotype of *H. syriacus* as seen in plants with spike inflorescence.
The investigation of duplication event timing in *H. syriacus* genome showed that two recent WGDs occurred after speciation. In the past 50 million years, the average global temperature has been $<20^\circ C$, which is far below the optimal flowering temperature for *H. syriacus*. Lower temperatures, especially between 5 and 20 MYA, could have been an environmental suppressor of *H. syriacus* pollination that prompted polyploidization to overcome these unfavourable conditions and unreduced gamete formation. Furthermore, low temperatures could have exerted selective pressure on *H. syriacus* to extend its blooming period for increased chance of pollination.

Perennial plants are prone to invasion by pathogens before reproduction, and many fungal and bacterial diseases often threaten the life cycle of *H. syriacus*. Aside from primary defenses, such as thickened cell walls and secondary metabolites, plants have numerous disease resistance (*R*) genes that confer protection against various pathogens. In *H. syriacus*, *R* genes account for $\sim0.53\%$ of its total predicted genes, which is lower than other plant genomes studies, whose *R* gene proportions ranged from 0.63 to 1.35%. However, subsets of these genes, including those with TIR- and RPW8-encoding motifs, are markedly over-represented in the *H. syriacus* genome compared to those of other plants. Genes in the RPW8-NBS-LRR subclass provide broad-spectrum resistance against powdery mildew pathogens in *Arabidopsis*, and genes in TIR-NBS subclasses are conserved in basal angiosperms and eudicots, such as *A. trichopoda* (Supplementary Table S8 and Supplementary Fig. S5), but are absent in most monocots. Their

### Table 4. Comparative NBS-LRR gene family numbers

| Predicted domain | Class   | *H. syriacus* | *G. raimondii* | *T. cacao* | *S. lycopersicum* | *A. thaliana* | *V. vinifera* | *O. sativa* | *A. trichopoda* |
|------------------|---------|---------------|----------------|-----------|-------------------|--------------|--------------|-------------|----------------|
| **TIR group**    | TIR-NBS-LRR | TNL 68 | 26 | 14 | 19 | 87 | 19 | 0 | 9 |
|                  | TIR-NBS  | TN 9 | 1 | 3 | 6 | 17 | 4 | 2 | 2 |
|                  | % on NBS genes | 17 | 9 | 6 | 9 | 61 | 7 | 0.4 | 10 |
| **Non-TIR group** | CC-NBS-LRR | CNL 183 | 220 | 202 | 116 | 52 | 138 | 337 | 27 |
|                  | CC-NBS | CN 77 | 24 | 25 | 37 | 3 | 19 | 104 | 27 |
|                  | NBS-LRR | NL 81 | 28 | 34 | 39 | 8 | 110 | 70 | 18 |
|                  | NBS | N 54 | 4 | 9 | 50 | 3 | 32 | 14 | 29 |
|                  | % on NBS genes | 84 | 91 | 94 | 91 | 39 | 93 | 99.6 | 90 |
| Total NBS genes | 472 | 303 | 287 | 267 | 170 | 322 | 527 | 112 |
| % on total genes | 0.53 | 0.81 | 0.97 | 0.77 | 0.63 | 1.22 | 1.35 | 0.41 |
| Total no. of genes | 87,603 | 37,505 | 29,452 | 34,727 | 27,206 | 322 | 527 | 112 |

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greater dominance in the *H. syriacus* genome indicates divergent evolution of TIR- and RPW-containing NBS genes from an ancestral origin may have led to more extensive expansion of this gene family. Moreover, the long life cycle of woody plants makes it difficult for them to adapt to pathogens undergoing more rapid evolution, thus favouring R gene maintenance and expansion for the plants' survival.

Polyplody in plants is a common mechanism for their adaptation to environmental change. After divergence from the *H. syriacus*-*G. raimondii* common ancestor, two WGDs and subsequent diploidization occurred in the *H. syriacus* genome to promote the plants' survival in unfavourable environments. During the diploidization events, low temperatures may have selected for the maintenance of duplicate flowering-related genes whose high copy numbers led to the multivoltinism and long blooming period phenotypes expressed by *H. syriacus*. Further analyses *H. syriacus*, *T. caacao* and *G. raimondii* (DD) genomes with another diploid genome, *G. arboresum* (AA) and allotetraploid genome, *G. hirsutum* (AD) will provide more information of evolution of Malvaceae plants.

**Availability**

This Whole Genome Shotgun project has been deposited at DDBJ/ENA/GenBank under accession MGJ00000000. The version described in this article is version MGJ01000000. The raw sequence reads have been deposited at DDBJ/ENA/GenBank under accession SRP087036 (PRJNA341341). In addition, the genome data of *H. syriacus* are accessible at https://hibiscus.kobic.re.kr/hibiscus.en.

**Conflict of interest**

None declared.

**Supplementary data**

Supplementary data are available at www.dnaresearch.oxfordjournals.org.

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Genome data including genome assembly, annotation and transcriptome data were also provided through webpage (https://hibiscus.kobic.re.kr/hibiscus.en).

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