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Chromosome-scale genome assembly of kiwifruit Actinidia eriantha with single-molecule sequencing and chromatin conformation capture

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**Abstract:**

Background: Kiwifruit (Actinidia spp.) is a dioecious plant with fruits containing abundant vitamin C and minerals. A handful of kiwifruit species have been domesticated, among which the A. eriantha is increasingly favored in breeding due to its superior commercial traits. Recently, elite cultivars from A. eriantha have been successfully selected and further studies on their biology and breeding potential require genomic information which is currently unavailable.

Findings: Here, we assembled a chromosome-scale genome sequence of A. eriantha cv. White using single-molecular sequencing and chromatin conformation capture. The assembly has a total size of 690.6 Mb and an N50 of 21.7 Mb. Approximately 99% of the assembly were in 29 pseudomolecules corresponding to the 29 kiwifruit chromosomes. Forty-three percent of the A. eriantha genome are repetitive sequences, and the non-repetitive part encodes 42,850 protein-coding genes, of which 39,075 have homologues from other plant species or contain protein domains. The divergence time between A. eriantha and its close relative A. chinensis is estimated to be 3.3 million years, and after diversification, 1,740 and 1,345 gene families are expanded or contracted in A. eriantha, respectively.

Conclusions: We generate a high-quality reference genome of kiwifruit A. eriantha. This chromosome-scale genome assembly is substantially better than two published kiwifruit assemblies from A. chinensis in terms of genome contiguity and completeness. The availability of A. eriantha genome provides a valuable resource for facilitating kiwifruit breeding and the studies of kiwifruit biology.

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Chromosome-scale genome assembly of kiwifruit *Actinidia eriantha* with single-molecule sequencing and chromatin conformation capture

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**Findings:** Here, we assembled a chromosome-scale genome sequence of *A. eriantha* cv. White using single-molecular sequencing and chromatin conformation capture. The assembly has a total size of 690.6 Mb and an N50 of 21.7 Mb. Approximately 99% of the assembly were in 29 pseudomolecules corresponding to the 29 kiwifruit chromosomes. Forty-three percent of the *A. eriantha* genome are repetitive sequences, and the non-repetitive part encodes 42,850 protein-coding genes, of which 39,075 have homologues from other plant species or contain protein domains. The divergence time between *A. eriantha* and its close relative *A. chinensis* is estimated to be 3.3 million years, and after diversification, 1,740 and 1,345 gene families are expanded or contracted in *A. eriantha*, respectively.

**Conclusions:** We generate a high-quality reference genome of kiwifruit *A. eriantha*. This chromosome-scale genome assembly is substantially better than two published kiwifruit assemblies from *A. chinensis* in terms of genome contiguity and completeness. The availability of *A. eriantha* genome provides a valuable resource for facilitating kiwifruit breeding and the studies of kiwifruit biology.

Key words: Kiwifruit; *Actinidia eriantha*; Genome assembly; single molecular sequencing; Hi-C
Data description

Introduction

Kiwifruit is well known as the king of fruits due to its remarkably high vitamin C content and abundant minerals [1, 2]. Native to China, kiwifruit belongs to the genus *Actinidia* which contains 54 species and 75 taxa [3]. All species in this genus are perennial, deciduous and dioecious with a climbing or scrambling growth habit, and they also have many common morphological features including the characteristic radiating arrangement of styles of the female flower and the structure of the fruit [4]. Despite rich germplasm resources in kiwifruit, only a few *Actinidia* species have been domesticated, such as *A. chinensis* var. chinensis, *A. chinensis* var. deliciosa and *A. eriantha*, whose fruit size are close to commercial standard [5-7].

Owing to its strong resistance to *Pseudomonas syringae* pv. *Actinidiae*, long shelf-life, enriched ascorbic acid and peelable skin [7-11], *A. eriantha* (2n=58) has been favored in kiwifruit breeding. Recently, new cultivars have been selected either from the wild germplasm of *A. eriantha* such as ‘White’ (Fig. 1) or from the interspecific hybridization between *A. eriantha* (♂) and *A. chinensis* (♀) such as ‘Jinyan’ [7, 12]. The ‘White’ has particularly large fruits (96 g on average) with green flesh and favorable flavor and has been widely cultivated in China [7].

*Actinidia eriantha* has also been used for genetic and genomic studies thanks to its high efficiency in genetic transformation and relatively short phase of juvenility [13]. The flowering and fruiting of *A. eriantha* can be accomplished within two years in greenhouse conditions with a low requirement for winter chilling [13]. In addition, roots of *A. eriantha* which contain many bioactive compounds such as triterpenes and polysaccharides are employed as a traditional Chinese medicine for the treatment of gastric carcinoma, nasopharyngeal carcinoma, breast carcinoma, and hepatitis [12, 14].
Previously, two kiwifruit genomes were published and both are from A. chinensis (‘Hongyang’ and ‘Red 5’) [15, 16]. These short-read based assemblies are very fragmented, possibly due to the high complexity and heterozygosity of the kiwifruit genomes as well as technical limitations. Here, we used single-molecular sequencing combined with the high-throughput chromosome conformation capture (Hi-C) technology to assemble the genome of the elite kiwifruit cultivar ‘White’ of A. eriantha. The availability of this high-quality chromosome-scale genome sequence not only provides fundamental knowledge regarding kiwifruit biology but also presents a valuable resource for kiwifruit breeding programs.

**Sample collection and genome sequencing**

Fresh young leaves were collected from a female individual of A. eriantha cv. White. High molecular weight (HMW) genomic DNA was extracted using the CTAB method as described in the protocol (https://www.pacb.com/wp-content/uploads/2015/09/Shared-Protocol-Preparing-Arabidopsis-DNA-for-20-kb-SMRTbell-Libraries.pdf). To construct genomic libraries (SMRTbell libraries) for PacBio long-read sequencing, HMW genomic DNA was sheared into fragments of approximately 20 kb using a Covaris g-Tube (KBiosciences p/n520079), enzymatically repaired and converted to SMRTbell template following the manufacturer’s instruction (DNA Template Prep Kit 1.0, PacBio p/n 100-259-100). The templates were size-selected using a BluePippin (SageScience, Inc.) to enrich large DNA fragments (> 10 kb) and then sequenced on a PacBio Sequel system. A total of nine SMRT cells were sequenced, yielding 3,889,480 reads with a mean and median length of 10,065 and 15,661 bp, respectively, and a total of 39.1 Gb sequences, about 51.6× coverage of the kiwifruit genome with an estimated size of 758 Mb based on the flow cytometry analysis [17] (Table S1).
Three paired-end Illumina libraries with insert sizes of 180, 220 and 500 bp, and seven mate-pair libraries with insert sizes of 3, 4, 5, 8, 10, 15, 17 kb, were prepared using Illumina’s Genomic DNA Sample Preparation kit and the Nextera Mate Pair Sample Preparation kit (Illumina, San Diego, CA), respectively. All libraries were sequenced on an Illumina HiSeq 2500 system, which yielded about 80.1 and 97.3 Gb of raw sequence data for paired-end and mate-pair libraries, respectively (Table S1). The raw Illumina paired-end reads were processed to remove adaptors and low-quality bases using Trimmomatic [18] (v0.35), and the mate-pair reads were cleaned using NextClip [19] (v1.3.1) with default parameters. Finally, we obtained 76.6 and 46.2 Gb high-quality cleaned sequences for paired-end and mate-pair libraries, respectively (Table S1).

To construct the Hi-C library, ‘White’ plants were grown in a greenhouse, and approximately 4–6 grams young leaves were then harvested and subsequently fixed in the formaldehyde (1% v/v) for 10 min at room temperature. The fixation was terminated by adding glycine to a final concentration of 0.125M. The fixed samples were ground into powder in liquid nitrogen and then lysed with the addition of Triton X-100 to a concentration of 1% (v/v). The nuclei were isolated and prepared for Hi-C library construction according to a previously published protocol [20]. The library was sequenced on an Illumina HiSeq 2500 system using the paired-end mode, which yield a total of approximately 118 million read pairs.

Transcriptome sequencing

To improve gene prediction, we generated transcriptome sequences from a pool of mixed tissues of ‘White’ including root, stem, leaf, flower, and fruits at 7, 30, 60, 90 and 120 days after anthesis. Total RNA was extracted from these tissues using an RNA extraction kit (BIOFIT, China), treated with DNase I and further purified with an RNA clean kit (Promega, USA). RNA-Seq libraries
were constructed with the NEBNext® Ultra™ RNA Library Prep Kit (Illumina, USA), and sequenced on an Illumina HiSeq 2500 system using the paired-end mode. A total of ~19.5 million raw read pairs were obtained, which were processed with Trimmomatic to remove adaptors. The cleaned reads were assembled de novo with Trinity [21] (version 2.4.0). Mapping of RNA-Seq reads to the genome assembly was performed with STAR [22] (version 020201), and read counting on the coding regions was performed with HTSeq [23] (version 0.6.0.).

Chromosome-scale assembly of the A. eriantha genome

We employed a strategy which took into account the unique advantage of different assemblers to construct the ‘White’ genome using PacBio long reads. First, PacBio long reads were corrected and assembled using the Canu program [24] (v1.7), which is a modularized pipeline consisting of three primary stages - read correction, trimming and assembly. The Canu-corrected reads were also assembled independently with the wtdbg program (https://github.com/ruanjue/wtdbg), a fast assembler for long noisy reads. Subsequently, the two independent assemblies (one with Canu and another with wtdbg) were merged by Quickmerge [25] (v0.2) to improve the contiguity. The merged assembly was further processed to correct errors using Pilon [26] (version 1.22) with high-quality cleaned Illumina reads from all paired-end and mate-pair libraries representing a total genome coverage of 171× (Table S1). This yielded 2,818,370 nucleotides, 2,495,388 insertions and 1,691,495 deletions being corrected. The resulting final assembled A. eriantha cv. ‘White’ genome contained 4,076 contigs with a N50 length of 539,246 bp and a cumulative size of 690,376,929 bp (Table 1). The contiguity and completeness of this assembly far exceeds that of two published kiwifruit A. chinensis genomes (Table 1).
Table 1 Assembly statistics

|                      | A. eriantha | A. chinensis |
|----------------------|-------------|--------------|
|                      | White       | Hongyang    | red5         |
| Contigs              |             |              |              |
| Total contig number  | 4,076       | 26,721       | 39,868       |
| (#)                  |             |              |              |
| Total contig length  | 690.4       | 604.2        |              |
| (Mb)                 |             |              |              |
| Contig N50 (kb)      | 539.2       | 58.9         |              |
| Contig N90 (kb)      | 50.7        | 11.6         |              |
| Longest contig length (kb) | 3,260.20 | 423.5        |              |
| Scaffolds            |             |              |              |
| Total scaffold number (#) | 1,735      | 7,698        | 3,887        |
| Total scaffold length (Mb) | 690.6      | 616.1        | 550.5        |
| Scaffold N50 (kb)    | 23,583.9    | 646.8        | 623.8        |
| Scaffold N90 (kb)    | 20,112.1    | 122.7        | 140.7        |
| Longest scaffold length (Mb) | 28.6      | 3.4          | 4.43         |
| Anchored to chromosome (Mb/%) | 682.4 / 98.84 | 452.4 / 73.4 | 547.9 / 98.9 |
| Anchored with order and orientation (Mb/%) | 634.4 / 91.90 | 333.6 / 54.1 |              |

To scaffold the contigs based on chromatin interaction maps inferred from the Hi-C data, we first used HiC-Pro [27] to evaluate and filter the cleaned Hi-C reads. The Hi-C data usually contains a considerable part of invalid interaction read pairs which are non-informative and need to be filtered out beforehand. Among the 51 million read pairs that were uniquely aligned to the A. eriantha assembly, 33 million (64.1%) were valid interaction pairs and their insertion size spanned predominantly from dozens to hundreds of kilobases, therefore providing efficient information for scaffolding. As a part of error correction of the assembly, we also used valid Hi-C reads to identify potential misassembled contigs. In principle, a genuine contig should display a continuous Hi-C interaction map whereas the discrete distribution of an interaction map likely indicates a misassembly. We examined the interaction map for each contig and broke 51 that were possibly misassembled. Subsequently, the corrected PacBio assembly was used for scaffolding using the LACHESIS program [28] with parameters “CLUSTER_MIN_RE_SITES=48, CLUSTER_MAX_LINK_DENSITY=2, CLUSTER_NONINFORMATIVE_RATIO=2, ORDER_MIN_N_RES_IN_TRUN=14, ORDER_MIN_N_RES_IN_SHREDS=15”. LACHESIS
assigned 3,666 contigs with a total size of 682,355,494 bp (98.84% of the assembly) into 29 groups corresponding to the 29 kiwifruit chromosomes (Fig. 2 and 3a), among which 634,430,648 bp (91.90%) had defined order and orientation (Table 1 and S2). The final chromosome-scale assembly had a total length of 690,781,529 bp and an N50 of 23,583,865 bp.

**Evaluation of the genome assembly**

We first evaluated the quality of the assembled *A. eriantha* ‘White’ genome by mapping Illumina genomic and RNA-Seq reads to the assembly. Reads from the three paired-end genomic libraries had very high mapping rates, ranging from 98.6% to 98.8%, and the properly paired read mapping rates were between 76.9% and 90.4%. For the RNA-Seq reads, 91.7% could be mapped to the genome and 87.1% were uniquely mapped. The high mapping ratio of both genomic and RNA-Seq reads suggest a high quality of the *A. eriantha* ‘White’ assembly.

We then identified synteny between the *A. eriantha* ‘White’ assembly and the assembly of *A. chinensis* ‘red5’ using MUMMER [29] (version 4.0.0beta2). In general, the two assemblies showed a high macro-collinearity, with only a few inconsistencies (Fig. 3b). Detailed check of the inconsistent regions using mate-pair read alignments supported the correct assemblies in the *A. eriantha* ‘White’ genome, and therefore the inconsistencies could be due to errors in the ‘red5’ assembly or structure variations between ‘White’ and ‘red5’ (Fig. S1).

**Repeat annotation**

Repeats were annotated following a protocol described in Campbell et al [30]. The customized repeat library was built to include both known and novel repeat families. We first searched the assembly for miniature inverted transposable elements (MITEs) using MITE-Hunter [31] with
default parameters. The long terminal repeat (LTR) retrotransposons were then identified from the A. eriantha ‘White’ genome using LTRharvest and LTRdigest wrapped in the GenomeTools package [32]. The LTR identification pipeline was run iteratively to collect both recent (sequence similarity ≥99%) and old (sequence similarity ≥85%) LTR retrotransposons. Candidates from each run were filtered based on the elements typically encoded by LTR retrotransposons. The default parameters (-minlenltr 100 -maxlenltr 6000 -mindistltr 1500 -maxdistltr 25000 -mintsd 5 -maxtsd 5 -motif tgca) were used in LTR calling according to Campbell et al. [30]. An initial repeat masking of A. eriantha ‘White’ genome was performed with the repeat library derived by combining the identified MITEs and LTR transposons. The repeat masked genome was fed to RepeatModeler (http://www.repeatmasker.org/RepeatModeler/) to identify novel repeat families. Finally, all identified repeat sequences were combined and searched against a plant protein database where transposons encoding proteins were excluded. Elements with significant similarity to plant genes were removed. The final repeat library contained 1,670 families, and 526 of them were potentially novel repeat families. We used this species-specific repeat library to mask the A. eriantha ‘white’ genome. Approximately 43.3% of the A. eriantha ‘White’ genome was masked, and the largest family of repeats was LTR transposons (Table S3). Repeat content identified in A. eriantha ‘White’ was much higher than that in A. chinensis (e.g. 36% in Hongyang [15]), and this difference could be largely due to the improvement of the repeat region assembly with PacBio long reads. In addition, variations between the two kiwifruit species could also contribute to this difference.

Prediction and functional annotation of protein-coding genes

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Protein-coding genes were predicted from the repeat-masked A. eriantha ‘White’ genome with the MAKER-P program [30] (version 2.31.10), which integrates evidence from protein homology, transcripts and ab initio predictions. The homology-based evidence was derived by aligning proteomes from 20 plant species to the ‘White’ genome assembly with exonerate (v2.26.1; https://www.ebi.ac.uk/about/vertebrate-genomics/software/exonerate). SNAP [33], AUGUSTUS [34] (version 3.3), and GeneMark-ES [35] (version 4.35) were used for ab initio gene predictions.

RNA-Seq data generated in this study were assembled de novo with Trinity and the assembled contigs were aligned to the ‘White’ genome assembly to provide transcript evidence. Predictions supported by the three different sources of evidence were finally integrated by MAKER-P, which resulted in a total of 52,514 primitive gene models. We then filtered and polished these gene models by two steps. First, we combined our RNA-Seq data with others collected from a previous study [36], and mapped the reads to the ‘White’ genome using the STAR program [22], and a total of 266 million read pairs were mapped. Based on the mapping, raw count for each predicted gene model was derived and then normalized to CPM (counts per million mapped read pairs). Gene models with ultra-low expression (CPM < 0.1) were less likely to be real genes. Furthermore, we found that these lowly expressed genes had relatively high annotation edit distance (AED) score, an indication of low-confidence as defined by the MAKER-P program. Therefore, for gene models with CPM < 0.1, we only kept those containing both pfam domains and homologous sequences in the NCBI nr protein database. After this filtering process 42,613 gene models were kept. Second, the predicted protein-coding genes of kiwifruit A. chinensis ‘red5’ have been manually curated [16], and therefore these gene models should have relatively higher accuracy and could be used to modify A. eriantha ‘White’ gene models whose predictions were not consistently supported by the different types of evidence. To this end, we performed another two ab initio predictions using
BRAKER2 [37] and GeMoMa [38] (version 1.5.2) with ‘red5’ proteome as the sole evidence. These two predictions were compared with the gene models predicted by MAKER-P. Consequently, a total of 237 gene models not predicted by MAKER-P were added and another 415 gene models which had better predictions by BRAKER2 or GeMoMa were used to replace the corresponding gene models predicted by MAKER-P. Finally, we obtained a total of 42,850 protein-coding genes in the *A. eriantha* ‘White’ genome, with a mean coding sequence (CDS) size of 1,004 bp and containing an average of five exons.

The predicted genes were functionally annotated by blasting their protein sequences against TAIR, Swiss-Prot and TrEMBL databases with an E-value cutoff of 1e-5. Functional descriptions of the protein hits were assembled with the AHRD program (https://github.com/groupschoof/AHRD) and transferred to *A. eriantha* genes. Protein domains were identified using InterProScan [39] (version 5.29-68.0) by searching the protein sequences against domain databases including PANTHER, Pfam, SMART, and PROSITE. The Gene Ontology (GO) terms were assigned to the *A. eriantha* ‘White’ predicted genes using the Blast2GO program [40] with entries from NCBI protein database and InterProScan. Collectively, 91.2% (N=39,075) of the predicted genes contain at least one annotation from the above databases (Table S4).

**Evolutionary and comparative genomic analysis**

To infer the divergence time between *A. eriantha* and *A. chinensis*, we identified gene orthology between the two species using MCScanX [41] and calculated synonymous substitution rate (Ks) between each orthologous pair. Three additional species, cultivated tomato (*Solanum lycopersicum*), wild tomato (*S. pennellii*) and potato (*S. tuberosum*), were also included in the
analysis. The Ks distribution (Fig. 4a) suggested that the divergence between the two kiwifruit species was earlier than that between the two tomato species. We dated the divergence by assuming a strict molecular clock [42], and the time when *A. eriantha* and *A. chinensis* diverged from the common ancestor was estimated to be ~3.3 million years ago (Mya), compared to ~1.9 Mya between *S. lycopersicum* and *S. penellii* and ~6.0 Mya between *S. lycopersicum* and *S. tuberosum*. Gene family evolution was analyzed by comparing genomes of *A. eriantha*, *A. chinensis*, *S. lycopersicum*, *S. tuberosum*, *Vitis vinifera*, *Arabidopsis thaliana* and *Oryza sativa*. A total of 17,593 orthogroups were defined by OrthoFinder [43] (version 2.2.6), among which 1,246 were single-copy gene families (Fig. 4b). The single-copy family genes were aligned and concatenated to build a species phylogenetic tree using IQ-TREE [44] (version 1.5.5) with a best-fitting model (Fig. 4c). Gene family expansion/contraction along the branches of the phylogenetic tree was analyzed by CAFÉ [45] (version 4.1). Finally, a total of 1,740 and 1,345 gene families were found apparently expanded and contracted, respectively, in *A. eriantha* (Fig. 4c).

**Conclusion**

Here, we report a high-quality reference genome of kiwifruit *A. eriantha* cv. White. The assembly from single-molecular sequencing combined with Hi-C scaffolding yielded a much more continuous and complete genome than the two previously published kiwifruit genomes. This genome will provide a valuable source for exploration of genetic basis of unique traits in kiwifruit and also facilitate the studying of sexual determination loci in the dioecious plants.

**Availability of supporting data**
This Whole Genome Shotgun project has been deposited at DBJ/ENA/GenBank under the accession QOVS00000000. The version described in this paper is version QOVS01000000. Raw sequencing reads have been deposited in the Sequence Read Archive (SRA) database under the accession number SRP155011. The Actinidia eriantha ‘White’ genome sequence and the annotation are also available at Kiwifruit Information Resource (http://bdg.hfut.edu.cn/kir/).

Competing interests
The authors have no competing interests to declare.

Abbreviation
CTAB: Cetyl trimethylammonium bromide;
NCBI: National Center for Biotechnology Information;
RNA-Seq: RNA sequencing;
SMRT: Single Molecule Real-Time;
MITE: miniature inverted transposable element;
LTR: long terminal repeat;
CPM: counts per million mapped read pairs;
Mya: million years ago

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Author contribution

W.T., X.S. and J.Y. contributed equally to this work. W.T., J.Y., X.T., Y.Y., X.N., M.M., D.Z., S.H., W.S., C.F. and M.L. collected plant samples, extracted DNA/RNA, and performed transcriptome sequencing and gene expression analyses; W.T., X.S., J.Y., X.T., C.J., Z.F. and Y.L. performed DNA sequencing, genome assembly, gene annotation, evolution and comparative genomic analyses, and website construction; X.S., W.T., Z.F. and Y.L. wrote and revised the manuscript; Y.L. and Z.F. conceived strategies, designed experiments and managed projects. All authors read and approved the manuscript.

Figure legends

Figure 1. Tree and fruits of *A. eriantha* cv. White.

Figure 2. Chromatin interaction map of *A. eriantha* derived from Hi-C data. Each group represents an individual chromosome.

Figure 3. Genome of *A. eriantha* and synteny between the two kiwifruit species. (a) Genome landscape of *A. eriantha* cv. White. Track A: gene density, Track B: repeat density, Track C: GC
content; all were calculated in a 500-kb window; (b) Genome synteny between A. eriantha cv. White and A. chinensis cv red5.

**Figure 4.** Evolutionary and comparative genomic analyses. (a) Distribution of synonymous substitution rate (Ks) between A. eriantha and A. chinensis, S. lycopersicum and S. penellii, and S. lycopersicum and S. tuberosum; (b) Orthogroups shared by selected species; (c) Species phylogenetic tree and gene family evolution. Numbers on the branch indicate counts of gene family that under either expansion (red) or contraction (green).

**Figure S1.** An example of genome assembly inconsistency between A. eriantha cv. White and A. chinensis cv red5. (a) A chromosomal segment assembled into the Chr23 in ‘red5’, is syntenic to the region located at the terminus of Chr19 in ‘White’ (b) Snapshots of Illumina mate-pair reads mapped to the junctions of the break point as well as nearby regions supporting the assembly of ‘White’.
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Figure 1
Figure 2.
Figure 4

(a) Distribution of synonymous substitution rates (Ks) for different species pairs. Red line: S. lycopersicum vs. S. pennellii; Green line: A. eriantha vs. A. chinensis; Blue line: S. lycopersicum vs. S. tuberosum. The peaks at 0.026, 0.046, and 0.084 represent the percentage of paralogous pairs.

(b) Phylogenetic tree showing the number of species and their evolutionary relationships. The tree branches indicate the number of species at each node: A. chinensis (14355), A. eriantha (14355), S. lycopersicum (14355), S. tuberosum (14355), V. vinifera, and O. sativa.

(c) Time distribution of species divergence in million years ago (mya). The numbers on the branches represent the divergence times. For example, A. chinensis diverged from A. eriantha at +2999 / -608 mya, and S. lycopersicum diverged from S. tuberosum at +1095 / -1289 mya.
Supplementary Tables

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**Supplementary Material**

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