Improved 93-11 Genome and Time-Course Transcriptome Expand Resources for Rice Genomics

Sen Wang1, Shenghan Gao2†, Jingyi Nie2†, Xinyu Tan2†, Junhua Xie1, Xiaochun Bi2, Yan Sun2, Sainan Luo2, Qianhui Zhu2, Jianing Geng2, Wanfei Liu1, Qiang Lin1, Peng Cui1*, Songnian Hu2* and Shuangyang Wu2*

1 Shenzhen Branch, Guangdong Laboratory for Lingnan Modern Agriculture, Genome Analysis Laboratory of the Ministry of Agriculture, Agricultural Genomics Institute at Shenzhen, Chinese Academy of Agricultural Sciences, Shenzhen, China, 2 State Key Laboratory of Microbial Resources, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China, 3 Gregor Mendel Institute, Austrian Academy of Sciences, Vienna, Austria

In 2002, the first crop genome was published using the rice cultivar 93-11, which is the progenitor of the first super-hybrid rice. The genome sequence has served as a reference genome for the indica cultivars, but the assembly has not been updated.

In this study, we update the 93-11 genome assembly to a gap-less sequence using ultra-depth single molecule real-time (SMRT) reads, Hi-C sequencing, reference-guided, and gap-closing approach. The differences in the genome collinearity and gene content between the 93-11 and the Nipponbare reference genomes confirmed to map the indica cultivar sequencing data to the 93-11 genome, instead of the reference. Furthermore, time-course transcriptome data showed that the expression pattern was consistently correlated with the stages of seed development. Alternative splicing of starch synthesis-related genes and genomic variations of waxy make it a novel resource for targeted breeding. Collectively, the updated high quality 93-11 genome assembly can improve the understanding of the genome structures and functions of Oryza groups in molecular breeding programs.

Keywords: time course transcriptome, alternative splicing, waxy, 93-11, chromosome level

INTRODUCTION

For decades, the rice cultivar 93-11, which is one of the main indica cultivars grown across China, has served as a reference genome for molecular breeding research (Yu et al., 2002). Recently, with the development of sequencing technology and the dramatic decrease in its cost, several chromosome-level assemblies of various rice cultivars have been reported, including telomere to telomere (T2T, telomeres to telomere). Transcriptome Expand Resources (TER) has served as a reference genome for molecular breeding research (Yu et al., 2002). Recently, with the development of sequencing technology and the dramatic decrease in its cost, several chromosome-level assemblies of various rice cultivars have been reported, including telomere to telomere. Transcriptome Expand Resources (TER) has served as a reference genome for molecular breeding research (Yu et al., 2002). Recently, with the development of sequencing technology and the dramatic decrease in its cost, several chromosome-level assemblies of various rice cultivars have been reported, including telomere to telomere.
telomere assemblies of ZS97 and MH63 (Du et al., 2017; Choi et al., 2020; Song et al., 2021). The assembly data reveals the complex genetic diversity that give rise to desirable domestication traits. As 93-11 is the parental cultivar of super-hybrid rice LYP9, which is widely grown across China, a high-quality reference genome of 93-11 is necessary for future molecular breeding of super-hybrid rice. Meanwhile, the 93-11 genome is the historical reference genome for indica; therefore, an upgrade may also facilitate evolutionary studies of these cultivars.

The amylase content (AC) is the most important factor that affects the taste and cooking quality of rice. The AC phenotype is mainly determined by the starch synthesis pathway. GBSSI (granule-bound starch synthase I), which is encoded by the waxy (Wx), is essential for the biosynthesis of amylase in rice (Vrinten and Nakamura, 2000; Tian et al., 2009). The evolution of the Wx largely contributed to improving the eating and cooking quality of rice, and its allelic variants (Wx^a, Wx^b, Wx^m, Wx^op, and Wx^mp) affected the AC of rice thereby affected consumer preferences (Zhang et al., 2019). The splicing form of the Wx and its related gene regulatory network contribute to the phenotypic selection in rice. Two Ser/Arg-rich proteins, Os-RSp29 and Os-RS23, can enhance, splice, and alter the acceptor splice site in the first intron of Wx^b (Ishiki et al., 2006). The Prp1 protein Du1 (dull endosperm1) affects the splicing efficiency of Wx^b and regulates starch biosynthesis (Zeng et al., 2007). In addition, the transcription factors OsBP-5, OsEBP and OsbZIP58 are involved in the regulation of Wx expression and starch biosynthesis (Wang et al., 2013). The metabolism involved in the amylase synthesis pathway is complicated and is a typical quantitative genetic trait. Recent studies have documented that the large structural variations between indica and japonica groups, which contribute to the cost of domestication of rice (Kou et al., 2020). Thus, it is necessary to genotype the agricultural traits of individual genomes.

In this study, we present the high-quality chromosome level genome sequence of the 93-11 cultivar using Illumina short pair-end reads, PacBio long reads, and Hi-C contact reads. This high-quality genome enabled us to investigate the structural variations between the indica and japonica groups. We also designed a continuous transcriptomic sampling experiment from 4 to 16 days after pollination (DAP) to investigate the gene expression pattern during seed development, as well as the gene regulation network of starch synthesis, which is conducive to the development of follow-up breeding work. This assembly and the related sequencing data provide a valuable complement to the rice study community and contribute to the subsequent breeding research.

**MATERIALS AND METHODS**

**Plant Material**

All cultivars, including 93-11, were collected from paddy fields in Beijing, China (Supplementary Table 1). Young leaves and seed samples collected at DAP 9 from Nipponbare and 12 other cultivars, as described in the Supplementary Table 1, were used to construct paired-end DNA-seq and RNA-seq libraries, respectively. For time course analysis, we collected samples during seed development until the full grain stage (from 4 to 16 DAP). All samples were washed using sterile physiological saline (37°C), snap-frozen in liquid nitrogen, and stored at −80°C.

**Sample Preparation**

Genomic DNA was extracted from young leaves using TRIzol to construct the paired-end library, single molecule real-time (SMRT) library, and Hi-C library, by following the manufacturer's protocol. The quality and quantity of the DNA were evaluated using a 0.8% agarose-gel electrophoresis, a NanoDrop micro-spectrophotometer (NanoDrop Technologies, Wilmington, DE, United States), and a Qubit® 3.0 Fluorometer (Thermo Fisher Scientific, Carlsbad, CA, United States), respectively.

Total RNA was extracted using the Huayueyang Plant RNA Extraction Kit (GK0416; Huayueyang, Beijing, China) according to the manufacturer's instructions. The RNA quality and quantity were examined using an electrophoresis agarose gel and a NanoDrop micro-spectrophotometer (NanoDrop Technologies), respectively.

**Genome Sequencing**

Sequel® Sequencing Kit 2.1 (PacBio) was used to construct an SMRT library, and high-quality long reads were generated using an RSII system. For Hi-C, libraries were performed according to a previously described method (Wang et al., 2020; Wu et al., 2020), via HindIII digestion and sequencing using a Hiseq X platform (Illumina) with an average depth of 69 ×. Poly (A) strand-specific libraries were constructed using the KAPA Stranded mRNA-Seq Library Preparation kit (KK8421; Roche, Pleasanton, CA, United States). All libraries were sequenced using an Illumina Hiseq 4000 sequencing system.

**Genome Assembly**

Subreads with an ultra-depth of 196 × that combined the dataset from a previous study (Zhang et al., 2018) were used to perform the de novo assembly using CANU (version 1.8) (Koren et al., 2017) with the parameters “minReadLength = 3,000, correctedErrorRate = 0.030, batOptions = -dg 3 -db 3 -dr 1 - ca 500 -cp 50.” The raw contigs were further corrected using the Illumina pair-end reads using Pilon (version 1.23) (Walker et al., 2014), and the haplotypic duplication contigs were removed using Purge Haplotigs (version 1.1.1) (Roach et al., 2018).

To construct a high-quality chromosome level assembly, we further carried out the following steps (Supplementary Figure 1): (I) All the contigs were anchored on the chromosome using the Hi-C reads by Juicer (parameters: -s HindIII) and 3d-dna pipelines (parameters: -r 0 -i 50000 -m haploid) (Durand et al., 2016; Dudchenko et al., 2017). (II) Misjoins of the interaction map were corrected in JBAT (Juicebox Assembly Tools). (III) The accurately identification of telomere and centromere sequences. The tandem repeats (TRs) in the contigs were identified using the TRF program (Benson, 1999). The ends of the contigs containing the tandem repeat motif AAACCCT or AGGGTTT were considered as telomere regions, and these contigs were located at the ends of the pseudo-chromosomes. Hmmsearch
(version 3.3) (Eddy, 1998) with the default settings was used to
detect the location of the centromere sequences in the
genome using an hmm file built from a file containing 155–
165 CentO satellite sequences. The contigs containing TRs
(unit: the satellite sequence) with a length of more than 10
kb were considered as centromere contigs and located in
the middle of the pseudo-chromosomes. (IV) Self-alignment
of all the contigs was performed by Mummer (version 4.0)
with the default settings to determine the contig overlap and
merge two adjacent contigs. (V) Long reads were aligned to
all the contigs using Minimap2 (version 2.17-r941) to detect
the adjacent contigs that shared the same long reads. If two
adjacent contigs overlapped with the two ends of long reads,
respectively, then the gap between the two contigs were closed
using the reads. (VI) Mummer (version 4.0) (Delcher et al.,
2003) with the default setting was used to align the contigs on
the three genomes [Nipponbare (Kawahara et al., 2013),
ZS97, and MH63 (Song et al., 2021)] to detect the potentially
adjacent contigs. If one unlocated contig is adjacent to a
contig in the interaction map along as supported by PacBio
reads, and the unlocated contig was put back into the map.
Steps IV and V were performed again to close gaps. (VII)
Long reads were aligned to the polished contigs to detect
sequence continuity. Finally, we manually checked the errors and
finalized the chromosome.

The completeness of the assembly was assessed using the
Benchmarking Universal Single-copy Orthologs (BUSCO)
pipeline (version 5.0.0) (Sepey et al., 2019) with the
embryophyta_odb10 dataset, which contains 1,614 highly
conserved genes. Merqury was used for the assembly consensus
accuracy estimation (Rhiie et al., 2020).

Repeat Annotation
The species-specific repetitive sequence library and transposable
element (TE) library were identified using RepeatModeler
(version 1.0.5) (Flynn et al., 2020) and Extensive de novo TE
Annotator (EDTA) (Su et al., 2021), respectively. Mummer
(version 4.0) the default parameters was used to align the
sequences from the species-specific repeat library (REL) to
transposable elements of the TE library (TEL). The sequences in
the REL with a coverage ratio of 0.5 to a sequence in the TEL
were filtered, and the TEL and REL with remained sequences
were merged to construct a non-redundant repetitive sequence
library. The dispersed repeats and TRs in the 93-11 genome
were annotated by RepeatMasker (version 4.0.5) (Tarailo-Graovac and
Chen, 2009) with the curated library.

Gene Annotation
Protein-coding gene annotation was performed using a strategy
integrating de novo prediction, transcriptome-based prediction,
and homology-based prediction. Hisat2 (version 2.1.0) and
StringTie (v2.1.2) were used to align the RNA-seq reads and
assemble the transcripts (Pertea et al., 2016). The complete
open reading frames (ORFs) of the transcripts from the 13
stages were identified to generate high-confidence gene models
using the TransDecoder pipeline (version 5.5.0) (Haas et al.,
2013). Augustus (version 3.2.2) (Keller et al., 2011) was used
to predict gene models based on the repeat-masked and
unmasked genomic sequences with the parameters “-s rice,”
respectively. The protein sequences obtained from Nipponbare
were aligned to the 93-11 genome using GenomeThreader
(version 1.7),1 with the parameters “-species rice.” The gene
models obtained from the three prediction methods were
integrated to generate a consensus gene set using GFFRead
(version 0.11.6) (Pertea and Pertea, 2020) and an in-house Perl
script. The gene structure was visualized via manual polishing
using Integrative Genomics Viewer (IGV) (Robinson et al.,
2011), and 3,215 genes were corrected based on the structural
information, completeness of genes, and similarities to proteins
of other species.

Gene functions were further annotated using protein-protein
basic local alignment search tool (BLASTP) (e-value: 1 × 10⁻⁵)
against the non-redundant protein database (NR) (Pruitt et al.,
2005) and Kyoto Encyclopedia of Genes and Genomes (KEGG)
database². Interproscan (v5.21) (Quevillon et al., 2005) was used
to identify the protein domains and annotate the gene ontology
(GO) protein function.

Structural Variation Identification
Mummer (version 4.0) (Delcher et al., 2003) with the default
parameters was used to align the 93-11 genome to the
Nipponbare genome. The alignment was filtered using delta-
filter implemented in Mummer with the parameters “-i 90 -l
10000,” and the filtered alignment was plotted and displayed
using mummerplot. The alignment was also filtered using delta-
filter with the parameters “-i 80 -l 1000” to analyze the syntenic
regions and genome structural variation between the 93-11 and
Nipponbare genomes.

Homolog Comparison
All-versus-all BLASTP (e-value: 1 × 10⁻⁵) was used to calculate
the pairwise similarities of the protein sequences between the
93-11 and Nipponbare genomes. The number of synonymous
substitutions per synonymous site (Ks) was calculated for the
orthologous gene pairs using codeml implemented in the PAML
program (version 4.9) (Yang, 2007), and the Ks distribution was
displayed using R (version 3.6.3). The syntenic gene regions
between 93 and 11 and Nipponbare were defined using MCloS
(Wang et al., 2012) with parameters: “-s 5 -m 10 -w 5,” and
the tandem duplication genes were obtained from the results.

Phylogeny Analysis
Protein sequences were generated obtained from six
domesticated cultivars (Oryza sativa indica cv. 93-11, O. sativa
indica cv. R498, O. sativa indica cv. Minghu63, O. sativa
indica cv. Zhenshan97, O. sativa japonica cv. Nipponbare,
and O. sativa japonica cv. Kitaake) and 10 wild species
(O. barthii, O. brachyantha, O. glaberrima, O. glumipatula,
O. longistaminata, O. meridionalis, O. nivara, O. punctata,
O. rufipogon, and Leersia perrieri)³. Then, OrthoFinder

¹https://genometheanalyzer.org/
²https://www.genome.jp/kegg/
³https://plants.ensembl.org/index.html
Identification of Rapidly Evolving Genes

The orthologous genes of the six species (O. sativa indica cv. 93-11, O. sativa indica cv. R498, O. sativa japonica cv. Nipponbare, O. sativa japonica cv. Kitaake, O. rufipogon, and O. nivara) were identified using OrthoFinder. Multiple protein alignments were performed using Clustal Omega, and the corresponding coding DNA sequence (CDS) alignments were converted using an in-house Perl script. Then, the trimAl (version 1.4) program (Capella-Gutierrez et al., 2009) was used to remove gaps in the CDS alignments. The branch model in PAML with a modified branch-site model, the null model (model = 0), and the alternative model (model = 2) were used to identify the rapidly evolving genes. A likelihood ratio test (LRT) with a df = 1 was performed based on the likelihood values obtained from the two models. Genes with a $P \leq 0.05$ were considered as rapidly evolving genes in the foreground branch.

Gene Ontology Enrichment Analysis

GO enrichment analysis and visualization were performed using BiNGO (version 3.04) (Maere et al., 2005) implemented in the Cytoscape (version 3.7.1) (Shannon et al., 2003) software, together with a hypergeometric test. The GO annotation profile of the 93-11 genome was constructed based on the InterPro result, and the ontology file was obtained from the GO website\(^1\). The GO terms with a $P \leq 0.05$ were considered significantly enriched.

Transcript Construction and Gene Expression Analysis

The high-quality RNA-seq reads for each sample were mapped to the 93-11 genome using Hisat2 (version 2.1.0) with the parameters, “--fr --rna-strandness RF” and the Sam files were converted to a Bam format and sorted using Samtools (version 1.6) (Li et al., 2009) with the default parameters. The transcripts of all samples were constructed using StringTie (version 2.1.2) with the parameter “–rf” and combined using TACO (version 0.7.3) (Niknafs et al., 2017). These transcripts were used to annotate the protein-coding genes. Salmon (version 1.4.0) (Patro et al., 2017) with the parameter “--SS_lib_type RF” was used to map the RNA-seq reads to the transcripts from the 93-11 genome and calculate the expression abundance (transcripts per million (TPM) and read count) of the genes and isoforms.

Gene Co-expression and Time-Course Analysis

The expression matrix was processed by removing the genes with a TPM $< 1$ of all samples to analyze the gene co-expression using the weighted correlation network analysis (WGCNA) (Langfelder and Horvath, 2008) package in R. Furthermore, the starch synthesis-related genes and transcription factors with a TPM $\geq 1$ were extracted to investigate the gene regulation network related to starch synthesis, and the highly correlated genes ($|r| \geq 0.9$ or weight $\geq 0.3$) were selected to construct the subsequent regulatory network. Finally, network visualization was performed using Cytoscape (version 3.7.1). Mfuzz (version 2.48.0) (Kumar and E Futschik, 2007) was used to cluster the time-series gene expression data from the 13 stages. The core genes with a membership coefficient value $\geq 0.9$ in each cluster were extracted and used for functional analysis.

Variant Calling

DNA-seq raw sequencing reads were trimmed to remove the low-quality and adaptor sequences using Trimmomatic (version 0.36) (Bolger et al., 2014) with the default parameters. Clean reads were mapped to the 93-11 genome using BWA (version 0.7.17-r1188) (Li and Durbin, 2009). Picard-tools (version 1.119)\(^2\) were used to remove the duplicate reads from the libraries. GATK (version 3.7) (McKenna et al., 2010) was used to identify the genomic variants: (1) RealignerTargetCreator and IndelRealigner were used to realign the reads to obtain reads with a lower mismatch rate; (2) HaplotyperCaller was used to identify and screen the genome variation (single nucleotide polymorphisms (SNPs and Indels); (3) GenotypeGVCFs was used to combine the variation sites of 14 cultivars.

RESULTS

Chromosome Level Genome and Annotation of the 93-11

To construct a high-quality chromosome level 93-11 genome, we used an integrated pipeline described in “Materials and Methods” section using ultra-high depth PacBio long reads ($\sim$196 $\times$) for assembly and Illumina reads for error correction (Supplementary Table 2). The total length of the assembly was 400 Mb with 281 contigs, and the contig N50 size was 17 Mb (Supplementary Table 3). Furthermore, a total of 24 contigs containing telomere sequences and 26 contigs containing CenO satellite sequences were identified. The assembled contigs were then anchored into 12 chromosomes using 27 Gb Hi-C (chromatin conformation contact) data and via manual correction, and the anchoring rate of the contig sequences was 98.25% (Supplementary Figure 2).

In total, the final assembled chromosome size was 393 Mb with a scaffold N50 size of 32 Mb (Figure 1 and Supplementary Table 3), which was larger than that of Nipponbare ($\sim$373 Mb). Compared with the previous assembly (Qin et al., 2021), we

\(^1\)http://geneontology.org/

\(^2\)http://picard.sourceforge.net
found good collinearity with fewer collapsed repeat regions between the pseudomolecules (Supplementary Figures 3, 4). In particular, our assembly corrected a potential assembly error of the previous version (Zhang et al., 2018) with a long insertion from chromosome 7 into chromosome 2. The LAI score (Ou et al., 2018), which evaluates the proportion of intact LTR-RT in this 93-11 assembly, was 26.02, indicating that the assembly had relatively high contiguity and quality. Assessment of the completeness of coding genes in the assembly using BUSCO, we observed that approximately 98.3% of the single copy orthologous genes (embryophytes_odb10) were complete (Supplementary Table 4). Additionally, approximately 99.34% of the Illumina reads, 97.02% of the PacBio long reads, and 95.87% of the RNA-seq reads could be mapped to the genome, further suggesting the completeness of the assembly (Supplementary Table 5). Further evaluation using Merqury revealed that the 93-11 genome had a high-quality score (QV) of 31.55, indicating a high (> 99.9%) assembly accuracy (Supplementary Figure 5). Overall, this is one of the most contiguous and complete assembly of rice genomes published to date.

This 93-11 assembly contained 190 Mb repetitive sequences, accounting for 48.28% of the genome, including 14.41% of DNA transposons, 28.11% of retrotransposons, and 1.12% of simple repeats. The total repeat length of the 93-11 genome was greater than that in the Nipponbare genome (Supplementary Table 6).

Protein-coding genes in the 93-11 genome were annotated using multi-evidence integrating strategy with manual correction based on pieces of evidence obtained from the transcriptome and Nipponbare gene models. A total of 40,345 genes were identified with an average length of 2,895 bp (Supplementary Table 7). Of all the annotated genes, 90.18% were aligned to the Nr database, 88.22% had functional annotations from the InterPro database, 55.07% (22,218) contained Pfam domains, and 42.28% (17,059) and 21.20% (8,555) were assigned to the GO terms and KEGG Orthology identifiers, respectively (Supplementary Table 8). Moreover, 1,616 transcription factors were detected and most of them belonged to the bHLH family (204).

The Genomes of 93-11 and Nipponbare Are Highly Similar, and Comparison of Gene Sets Showed Cultivar Specific Genes Accounting for Various Functions

To compare the 93-11 genome and Nipponbare genomes, we first cross-mapped the Illumina reads to each assembly. Approximately 98.21 and 97.88% of the reads were mapped to the 93-11 and Nipponbare genomes, respectively. Collinearity analysis between the 93-11 and Nipponbare genomes showed that approximately 91.12% of the 93-11 genome sequences had one-to-one syntenic blocks with the Nipponbare genome (nucleotide identity of 97.21%). These results confirmed that the two genomes were very similar.

A total of 32,961 genes in the 93-11 genome were homologous to those in Nipponbare, of which 27,729 genes were in the syntenic regions and 5,232 genes were in the non-syntenic regions (Figure 2A). Comparative genomics analysis showed that those genes in the syntenic region were more conserved, while the non-syntenic genes had higher evolution rates (Supplementary Figure 6). The non-syntenic genes was enriched in the following GO terms: telomere maintenance, telomere organization, and chromosome organization (Figure 2B and Supplementary Table 9), suggesting diversity in chromatin variation between the 93-11 and Nipponbare genome, which
FIGURE 2 | Comparative genomic analysis of 93-11 and Nipponbare. (A) Collinearity between the 93-11 and Nipponbare genomes. There are 27,729 homologous genes in the collinear regions, indicating that the two genomes have a high degree of collinearity. (B) Gene ontology (GO) enrichment of the genes in non-syntenic regions. (C) GO enrichment of the rapidly evolving genes in the 93-11 genome. These results indicate the species specificity of gene evolution after species differentiation. The color scale indicates the P-value, and the size of the circle indicates the number of genes.

Still need to be further validated with improved Nipponbare due to incompleteness of current Nipponbare genome. We further checked unaligned proteins specific to both genomes, and identified 1,026 genes and 743 in the 9,311 and Nipponbare genomes, respectively, most of which were hypothetical proteins.

Rapid evolution results in the generation of novel genes that enables better adaptation to the environment (Crow et al., 2006). We identified 498 and 481 rapidly evolving genes in the 93-11 and Nipponbare genome, respectively. Interestingly, the functions of these two gene sets were significantly different. The rapidly evolving genes in the 93-11 genome were enriched in functions related to response to endogenous stimulus, response to organic substance, methylation, and cellular nitrogen compound metabolic process (Figure 2C and Supplementary Table 10), whereas those in the Nipponbare genome were related to the organic substance biosynthetic process, translational initiation, and regulation of biological quality (Figure 2D and Supplementary Table 11). These genes may account for the high disease-resistance character of the 93-11 rice and the good cooking quality of Nipponbare rice.

Phylogeny Analysis Indicates the Different Origins of Indica and Japonica

Despite the high synteny and sequence similarity between the 93-11 and Nipponbare genome, the phylogenetic tree revealed that the 93-11 (representing indica) and Nipponbare (representing japonica) were clustered with two different wild species, O. nivara and O. rufipogon (Figure 3A and Supplementary Figure 7), which was agreed with the Ks results (Supplementary Figure 8). Nucleotide similarity analysis showed that the sequence similarity between 93-11 and O. nivara and between Nipponbare and O. rufipogon was 98.07 and 98.27%, respectively, which are both higher than that between 93-11 and Nipponbare (97.20%) (Supplementary Table 12). In conclusion, it is highly likely that the rice japonica and indica groups may arise separately from the progenitor O. rufipogon and O. nivara, rather than differentiating into subspecies, which confirms the results reported in a previous studies (Choi et al., 2017; Stein et al., 2018). This suggests that mapping sequencing data from the rice indica group to the Nipponbare genome might be incorrect. Therefore, it is always a good idea to map sequencing data of the indica cultivars to the 93-11 genome and vice versa.

Comparative Analysis Showed an Inversion Which Nestled Another Inversion Region

To discover the structural variations (SVs) that potentially shaped the 93-11 cultivar, compared with the Nipponbare genome, we identified 8,188 large fragment insertions (≥ 1 Kb) and 9,045 inversions in the 93-11 genome. The insertions and inversions account for 8.73% (34 Mb) and 9.80% (38 Mb) of the 93-11 genome, respectively. Notably, there was one remarkable pericentric inversion that was found to be specific for 93-11, which was located at 12994554–18810736 on chromosome...
6 (corresponding to chromosome 6:13119682–17632495 in Nipponbare) (Figure 3B and Supplementary Figure 9). Interestingly, we found a nestled inversion region of 700 Kb within the pericentric region, located at 14563302–15266093 in the short arm of chromosome 6 (corresponding to chromosome 6:15702521–16205397 in Nipponbare) (Figure 3B). It looks like a secondary inversion, but it is not clear whether this nestled inversion is reserved when the inter-arm inversion occurs or if it is inverted after the inter-arm inversion occurs. In addition, another paracentric inversion (93-11: 18812083–19050675; Nipponbare: 17659075–17900016) next to the remarkable inversion was identified. These inversions in the 93-11 genome were also confirmed by PacBio read coverage visualization (Supplementary Figure 10).

The inversion regions carried 367 genes, of which 187 were homologous to those of Nipponbare (Supplementary Table 13), and 224 were expressed during seed development. Genes encoding P450, NB-ARC, and multi-antimicrobial extrusion proteins were identified, which are related to disease resistance and seed development function (Supplementary Table 13). Functional enrichment was mainly included in response to stimulus/chemical/drug, transmembrane transporter activity, xenobiotic transmembrane transporter activity, oxidation-reduction process, organic cyclic compound binding, and cofactor binding (Figure 3C and Supplementary Table 14), which demonstrates the potential high disease resistance character of 93-11. Nine transcription factors were also found in this region, including one bHLH, one NAC, one ZF-HD, one CPP, one G2-like, one MIKC_MADS, one M-type_MADS, and two HB-other. In addition, the Ks result indicated that the genes in the inversion regions were evolving significantly faster than the other genes on chromosome 6 (Supplementary Figure 11).

Overall, these results demonstrate that large SVs may significantly contribute to cultivar formation and may facilitate the rapid adaptation or domestication of cultivars. The identified SVs could also be used as molecular markers to distinguish between indica and japonica for genetic traceability.

Continuous Sampling Provides Time-Course Insight on the Transcriptomic Pattern During Seed Development

The time-course RNA sequencing approach provides an opportunity to better evaluate gene expression patterns with sample collection in parallel over time (Spies and Ciaudo, 2015;
Thomas et al., 2020). We collected samples at 13 stages from 4 to 16 DAP to understand the seed developmental processes of rice using integrated multiple methods based on TPM values of the 13 stages, including Pearson correlation, hierarchical clustering (HC), WGCNA, and transcriptome-wide time series expression analysis. The results demonstrated that the number of expressed genes (EGs) and stage-specific expressed genes (SSEGs) increased from 4 to 5 DAP and decreased from 5 to 16 DAP. At 5 DAP, the most EGs and SSEGs were observed (19,255 and 441, respectively) (Supplementary Table 15 and Supplementary Figure 12). The functions of the stage-specific highly expressed genes (SSHEGs) and SSEGs at 5 DAP were mainly involved in defense response, response to stress, carbohydrate metabolic process, cell wall biogenesis, catalytic activity, hydrolase activity, peptidase regulator activity, etc. (Supplementary Table 16). All these results indicate that the rice seed development had entered an active state from this stage; the content of the seed had been synthesized quickly, and the resistance to external biological stimuli was enhanced.

A total of 28 expression modules (except for the gray module) were identified using WGCNA, and 13 modules were highly correlated with the 13 stages ($r > 0.9$) (Figure 4A). The analysis was consistent with the results from the transcriptome-wide time-series expression analysis (Supplementary Figure 13), which indicates that the gene expression was stage-specific, and the expression pattern of the core genes in each module could be used as a marker for each stage. Hierarchical clustering analysis results showed two main groups, corresponding to the milky and dough stages of rice seed development (Figure 4B and Supplementary Figure 14), which were also identified based on their morphology. Each main group was further divided into two sub-groups, corresponding to the early and late periods of their respective stages.

Functional analysis of the highly EGs in the first five stages (early stage of milky stage) (Figure 4C) showed that these genes were mainly involved in biosynthetic processes, genes expression, translation, binding, enzymatic activity (such as hydrolase, ATPase, ligase, oxidoreductase, pyrophosphatase, etc.), and
regulation of translation (Supplementary Table 17). This indicated that the material synthesis in the seed was active during these stages, and the content began to accumulate rapidly. We also found that 1,004 genes related to the above-mentioned biological processes (Supplementary Table 18) showed the expression levels in the late stages (Figure 4D), which further indicates that these biological processes, such as content synthesis, slowed down in the late stages of seed development, leading to a gradual accumulation of the content in the seeds.

The genes in cluster 10 showed high expression levels during the milky stage but low expression levels during the dough stage (Figure 4E). Functional enrichment analysis revealed that these genes were highly associated with vesicle-mediated transport, intracellular transport and localization, cellular nitrogen compound metabolism, cytoskeleton organization and regulation, protein-containing complex, organelle, cytoskeleton, binding, intramolecular transferase activity, peptidase activity, structural constituent of cytoskeleton, etc. (Supplementary Table 19). These results indicate that the stage-specific genes expressed at the milky stage may be highly related to the formation of macromolecular structures during seed development. The expression pattern of these genes could be used as a marker of the seed development transition from the milky to the dough stages.

Gene Regulation Network Related to Starch Synthesis During Endosperm Genesis

Starch is synthesized in amyloplasts from its initial substrates, glucose 1-phosphate (G1P) and glucose 6-phosphate (G6P). G1P and G6P are imported into an amyloplast and synthesized to amylose and amylopectin by several enzymes playing orchestrated roles, including phosphoglucomutase (PGM), ADP-glucose pyrophosphorylase (AGP), granule-bound starch synthase (GBSS), soluble starch synthase (SS), starch branch enzyme (SBE), starch debranching enzyme (DBE), starch phosphorylase (Pho), and dismutase (DPE) (Figure 5A). In total, 32 starch synthesis-related genes (SSRGs) were annotated in the 93-11 genome (Figure 5B and Supplementary Table 20). Gene expression analysis demonstrated that there were significant differences in the expression levels of the SSRGs, as well as in the expression patterns during seed development.

We further analyzed the co-expression network of the SSRGs and transcription factors (TFs) using WGCNA. A total of seven modules containing SSRGs were identified, and highly correlated connections ($|r| \geq 0.9$ or weight $\geq 0.3$) were selected to construct the subsequent regulatory network (Figure 5C and Supplementary Figure 16). Eight SSRGs and 54 TFs participate in the network. Among the TFs, 22 of them were directly involved in regulating the expression of SSRGs, of which the bZIP transcription factor $R9311C07g25353$ (corresponding to OsbZIP58) regulated starch synthesis in rice endosperm (Supplementary Tables 21, 22; Wang et al., 2013). The network showed that these TFs belonged mainly to the C3H and MYB families, including 5 and 4 genes, respectively. In addition to OsbZIP58, $R9311C01g03921$ (OsDSR1), $R9311C04g17870$ (OsC3H31), $R9311C09g31905$, and $R9311C06g22141$ (OsARF16) may also be involved in regulating the expression of Wx. $R9311C09g31905$ is a transcription initiation factor that showed a negative regulation ($r = -0.8439$) with Wx expression (Supplementary Table 22). In addition, three transcription termination factors ($R9311C06g22381$, $R9311C06g22279$, and $R9311C11g36090$) were detected in the network (Supplementary Table 21). Overall, the network indicated that these TFs may play a vital role as potential regulators of SSRGs expression and starch synthesis.

Variations in Alternative Splicing Reveals Isoform Preferences in Amylose Synthesis-Related Genes

Alternative splicing is an important post-transcriptional regulatory mechanism that increases proteome diversity by altering the mRNAs. Interestingly, based on the endosperm transcriptome data, we noticed a special type of alternative splicing event induced by splice acceptor “drift” at the acceptor splice site (ASS) of the first intron of three SSRGs, $BT1$ ($R9311C02g06322$), $AGPL2$ ($R9311C01g02861$), and Wx ($R9311C06g21642$) (Supplementary Figure 16). Briefly, the splice site variation was due to multiple occurrences of the AG sequences in the 3’-ends of the first intron. In $BT1$ and $AGPL2$, the sequence of the 3’-end was TAGCAGCAG and TAGTTGCAG, producing three and two acceptor splice sites, respectively. Two ASSs in Wx were generated using the CAGTGCAG. In summary, adjacent AGs can easily form alternative splice sites in the pre-mRNA.

In addition to the ASS in the first intron, three donor splice sites (DSSs) in the first intron and two alternative DSSs in the eighth intron were identified, located at 1631782, 1631874, 1631875, 1634516, and 1634522 in chromosome 6, respectively (Figure 6A). These alternative splice sites resulted in the formation of five isoforms (Wx-1/Wx-2/Wx-3/Wx-4/Wx-5) of the Wx gene in the 93-11 cultivar (Figure 6A). Wx-1 and Wx-4 corresponded to Os06t0133000-02 and Os06t0133000-01 of Nipponbare, respectively. It is known that an SNP (T/G) at the 5’-end of the first intron of the Wx, which is located exactly at the splice donor site of the first intron, determines the retention of the first intron, thus forming two isoforms Os06t0133000-02 (T: intron retention) and Os06t0133000-01 (G: intron skipping). In the 93-11 genome, the SNP (chromosome 6:1631875) was of the T type, which is the same as that of Nipponbare; however, surprisingly, both isoforms could be detected in the transcriptome. This indicates that there are still some pre-mRNAs that could use this splicing site to produce mature mRNA.

As reported in a previous study (Frances et al., 1998), the proper splicing of the first intron of the Wx is critical for activating the GBSSI enzyme. Two cryptic DSSs (1631782 and 1631874) were detected in the Wx of 93-11 to form Wx-2, Wx-3, and Wx-5, which excluded the first intron. Interestingly, the expression levels and patterns of these five transcripts were significantly different (Figure 6B and Supplementary Table 23). Notably, the expression level of the Wx-2 isoform was much...
higher than those of the other isoforms. This showed that the Wx-2 isoform was the preferred in 93-11 and might be the main source of active GBSSI enzyme during seed development.

The Isoform Preference Is Closely Related With the (CT)$_n$ Microsatellite Polymorphism in the Waxy

A previous study (Bao et al., 2002) has reported that there was a (CT)$_n$ polymorphism downstream of I1-DSS in the rice population. To investigate if the (CT)$_n$ variation results in a Wx preference by impacting the ASS selection of the first intron, we further expanded the transcriptome analysis to 13 rice cultivars (Supplementary Table 1). Gene expression analysis showed that Wx-2 was highly expressed only in the medium amylose content (AC) cultivars (Figure 6C). The genotyping of (CT)$_n$ demonstrated that the length of (CT)$_n$ is closely related to AC in rice cultivars (Figure 6D): (CT)$_{17}$, (CT)$_{18}$ and (CT)$_{11/12}$ were present in low, medium, and high AC cultivars, respectively. Interestingly, in comparison with the low AC cultivars, a slight extension of (CT)$_n$ from $n = 17$ to $n = 18$ could activate the expression of the Wx-2 isoform, standing out from the low robust Wx-1. This result suggests that in addition to splice site variations, (CT)$_n$ may also play an important role in the splicing efficiency and isoform preference in rice cultivars.

In addition, we investigated the potential regulators involved in pre-mRNA splicing of the Wx using Pearson correlation and WGCNA based on all the expressed transcripts at the 13 stages. A total of 52 identified genes had a high correlation coefficient ($|r| > 0.8$) with any of the five isoforms of the Wx. These genes included 14 genes with an RNA recognition motif domain (PF00076) and six splicing factors (Supplementary Table 2). Two Ser/Arg-rich proteins, R9311C04g14651 (Os-RSp29) and R9311C02g08144 (Os-RSZ23) and a dull endosperm1 R9311C10g34727 (Du1) were reported to be involved in the splicing of the Wx pre-mRNA. The analysis
revealed that these genes may be potential regulatory factors for the alternative splicing of the Wx pre-mRNA.

**DISCUSSION**

To date, over 150 rice genome assemblies have been deposited in NCBI, of which 44 are chromosome-level assemblies and one is a complete genome assembly (Benson et al., 2013). Over the past few decades, a complete telomere to telomere (T2T) assembly has always been the final goal of all genome projects. As sequencing technology evolves and its costs are being reduced, we are now closer to achieving this goal than ever before. Currently, the biggest challenge in constructing a T2T complete genome is the complexity of the centromeres and the highly duplicated transposable elements (Li et al., 2021; Rhie et al., 2021). That is mainly because the high read length and the low error rate do not converge with each other. In this study, we attempted to use long reads with an ultra-depth (∼196 ×) for assembly, with a much lower corrected error rate while maintaining a relatively high read length. The results from this study could help solve the problems associated with the low-complexity regions, such as the telomeres and centromeres. The results showed that this strategy outperformed those used for previous assemblies as it yielded higher continuity. Using Hi-C scaffolding, short-read polishing, and manual curation, we elevated the reference genome of 93-11 to a higher quality level. This assembly not only provides robust data support for genome variation analysis, but also provides a valuable reference for applying ultra-deep coverage long-read assembly for further improvement of other complex plant genomes.

Owing to the lack of sufficient transcriptome data and the false positive error of annotation tools, manual checking is an efficient way to strengthen the automatic annotation result. We manually confirmed around 3,000 doubtful genes and revised them, which provided a reliable annotation file compared with the reference annotation sets. Orthologous analysis between the 93-11 and Nipponbare genomes yielded many orphan genes (7,383 in 93-11 and 3,750 in Nipponbare). However, when these orphan genes were cross mapped to the 93-11 and Nipponbare genomes, the number of species-specific genes was significantly reduced (1,026 in 93-11 and 743 in Nipponbare). This phenomenon may be due to differences between the cultivars, a lack of a phased (diploid) genome, or errors in gene annotation, and
further in-depth research is required. Although the two genomes have high similarities, there are significant differences in the number of genes and genome structure between the two cultivars. Differences between the two genomes make it inappropriate to use Nipponbare as a reference genome to explore indica cultivars. Therefore, it is necessary to construct a high quality indica rice reference genome or pan-genome.

A previous study showed that the Wx and SSIIa seem to be the main trait loci for the domestication in rice (Kharabian-Masouleh et al., 2012). In this study, we found that the complexity of the Wx isoforms and their preferences could be used to distinguish the AC of cultivars. The variation in the Wx of rice offers several options for molecular breeding programs. Previous studies have shown that the polymorphism of simple repeat (CT)n occurring in the 5′UTR region of genes could influence the promoter activity and gene expression in plants. In this study, we found that even tiny changes in microsatellite (CT)n motif could influence the isoform preference of the Wx and resulted in an apparent change in AC, which indicates the sophisticated mechanism behind splicing factors and splicing-related proteins that mediates alternative splicing.

CONCLUSION

In conclusion, our study demonstrates that the near-complete genome assembly is feasible to efficiently decipher multi-omics data. Moreover, the reference genome of the rice cultivar 93-11 could be used as a model to study the AC-related traits and the complex genome variations between indica and japonica, which may be helpful for the future breeding programs.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: http://bigd.big.ac.cn/gsa, GSA: CRA00469 and CRA005384, https://bigd.big.ac.cn/gwh, GWH: GWHBEBR0000000.

AUTHOR CONTRIBUTIONS

SeW, PC, SH, and SYW conceived this study and wrote the manuscript. SeW, SG, and SYW designed the experiments, collected the samples, and performed the analyses. JN and JX collected the data and partially performed the analysis. XT, XB, YS, SL, QZ, JG, WL, and QL assisted in sample collection. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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REFERENCES

Bao, S., Corke, H., and Sun, M. (2002). Microsatellites in starch-synthesizing genes in relation to starch physicochemical properties in waxy rice (Oryza sativa L.). Theor. Appl. Genet. 105, 898–905. doi: 10.1007/s00122-002-1049-3
Benson, D. A., Cavanaugh, M., Clark, K., Karsch-Mizrachi, I., Lipman, D. J., Ostell, J., et al. (2013). GenBank. Nucleic Acids Res. 41, D36–D42. doi: 10.1093/nar/gks1195
Benson, G. (1999). Tandem repeats finder: a program to analyze DNA sequences. Nucleic Acids Res. 27, 573–580. doi: 10.1093/nar/27.2.573
Bolger, A. M., Lohse, M., and Usadel, B. (2014). Trimmomatic: a flexible trimmer for illinuma sequence data. Bioinformatics 30, 2114–2120. doi: 10.1093/bioinformatics/btu170
Capella-Gutierrez, S., Silla-Martinez, J. M., and Gabaldon, T. (2009). trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. Bioinformatics 25, 1922–1923. doi: 10.1093/bioinformatics/btp348
Choi, J. Y., Lye, Z. N., Groen, S. C., Dai, X., Rughani, P., Zaaijer, S., et al. (2020). Nanopore sequencing-based genome assembly and evolutionary genomics of circumbasmati rice. Genome Biol. 21:21. doi: 10.1186/s13059-020-1938-2
Choi, J. Y., Platts, A. E., Fuller, D. Q., Hsing, Y. I., Wing, R. A., and Purugganan, M. D. (2017). The rice paradox: multiple origins but single domestication in Asian rice. Mol. Biol. Evol. 34, 969–979. doi: 10.1093/molbev/msx049
Crow, K. D., Wagner, G. P., and Investigators, S. T.-N. Y. (2006). Proceedings of the SMSE tri-national young investigators’ workshop 2005. What is the role of genome duplication in the evolution of complexity and diversity? Mol. Biol. Evol. 23, 887–892. doi: 10.1093/molbev/msj083
Darriba, D., Taboada, G. L., Doallo, R., and Posada, D. (2011). ProTest 3: fast selection of best-fit models of protein evolution. Bioinformatics 27, 1164–1165. doi: 10.1093/bioinformatics/btr088
Delscher, A. L., Salzberg, S. L., and Phillippy, A. M. (2003). Using MUMmer to identify similar regions in large sequence sets. Curr. Protoc. Bioinformatics 10:13. doi: 10.1002/0471250953.bi100360
Du, H., Yu, Y., Ma, Y., Gao, Q., Cao, Y., Chen, Z., et al. (2017). Sequencing and de novo assembly of a near complete indica rice genome. Nat. Commun. 8:15324. doi: 10.1038/ncomms15324
Dudchenko, O., Batra, S. S., Omer, A. D., Nyquist, S. K., Hoeger, M., Durand, N. C., et al. (2017). De novo assembly of the Aedes aegypti genome using Hi-C yields chromosome-length scaffolds. Science 356, 92–95. doi: 10.1126/science.aaj3327
Durand, N. C., Shamim, M. S., Machol, I., Rao, S. S., Huntley, M. H., Lander, E. S., et al. (2016). Juicer provides a one-click system for analyzing loop-resolution Hi-C experiments. Cell Syst. 3, 95–98. doi: 10.1016/j.cels.2016.07.002
Eddy, S. R. (1998). Profile hidden markov models. Bioinformatics 14, 755–763. doi: 10.1093/bioinformatics/14.9.755

Emms, D. M., and Kelly, S. (2019). OrthoFinder: phylogenetic orthology inference for comparative genomics. *Genome Biol.* 20:238. doi: 10.1186/s13059-019-1832-y

Flynn, J. M., Hubley, R., Goubert, C., Rosen, J., Clark, A. G., Feschotte, C., et al. (2020). RepeatModeler2 for automated genomic discovery of transposable element families. *Proc. Natl. Acad. Sci. U.S.A.* 117, 9451–9457. doi: 10.1073/pnas.1912056117

Frances, H., Bligh, J., Larkin, P. D., Roach, P. S., Jones, C. A., Fu, H., et al. (1998). Use of alternate splice sites in granule-bound starch synthase mRNA from low-amylose rice varieties. *Plant Mol. Biol.* 38, 407–415. doi: 10.1007/s11103-013-084

Haas, B. J., Papanicolaou, A., Yassour, M., Grabherr, M., Blood, P. D., Bowden, J., et al. (2013). De novo transcript sequence reconstruction from RNA-seq using the Trinity platform for reference generation and analysis. *Nat. Protoc.* 8, 1949–1952. doi: 10.1038/nprot.2013.084

Ishikawa, Y., de la Bastide, M., Hamilton, J. P., Kanamori, H., McCombie, W. R., Ouyang, S., et al. (2013). Improvement of the *Oryza sativa* Nipponbare reference genome using next generation sequence and optical map data. *Rice* 6:4. doi: 10.1186/1939-8433-6-4

Keller, O., Kollmar, M., Stanke, M., and Waack, S. (2011). A novel hybrid gene prediction method employing protein multiple sequence alignments. *Bioinformatics* 27, 757–763. doi: 10.1093/bioinformatics/btr100

Kawahara, Y., ., Yamanaka, A., Watanabe, H., Minegishi, M., and Akiyama, N. (2020). Progress in the reference genome of *Oryza sativa* using a new mapping tool developed using optical maps. *Genome Res.* 30, 2105–2117. doi: 10.1101/gr.210557.119

Koren, S., Walenz, B. P., Berlin, K., Miller, J. R., Bergman, N. H., and Phillippy, A. M. (2017). Canu: scalable and accurate long-read assembly via iterative k-mer weighting and repeat separation. *Genome Res.* 27, 722–736. doi: 10.1101/ gr.210587.116

Kou, Y., Liao, Y., , Tian, X., Emerson, J. J., et al. (2020). Dynamics in transcriptomics: advancements in experimental design and analysis. *Comput. Struct. Biotechnol. J.* 18, 146–158. doi: 10.1186/s13059-019-2332-8

Langfelder, P., and Horvath, S. (2008). WGCNA: an R package for weighted correlation network analysis. *BMC Bioinformatics* 9:559. doi: 10.1186/1471-2105-9-559

Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with burrows-wheeler transform. *Bioinformatics* 25, 1754–1760. doi: 10.1093/bioinformatics/btp324

Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., et al. (2009). The sequence alignment/map format and SAMtools. *Bioinformatics* 25, 2078–2079. doi: 10.1093/bioinformatics/btp522

Li, K., Jiang, W., , Meng, D. Y., Guo, L. M., and Lu, J. Z., et al. (2021). Gapless genome assembly of 33 genetically diverse rice accessions reveals hidden genomic variations. *Cell* 184, 3542–3558 e16. doi: 10.1016/j.cell.2021.04.046

Lin, K., Jiang, W., Hui, Y., Kong, M., Feng, L. Y., Gao, L. Z., et al. (2021). Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. *Nat. Protoc.* 11, 1650–1667. doi: 10.1038/s41596-019-095

Pratt, K. D., Tatusova, T., and Maglott, D. R. (2005). NCBI reference sequence (RefSeq): a curated non-redundant sequence database of genomes, transcripts and proteins. *Nucleic Acids Res.* 33, D50–D54. doi: 10.1093/nar/gk025

Qin, P., Lu, H., Du, H., Wang, H., Chen, W., Chen, Z., et al. (2021). Pan-genome analysis of 33 genetically diverse rice accessions reveals hidden genomic variations. *Cell* 184, 3542–3558 e16. doi: 10.1016/j.cell.2021.04.046

Robinson, J. T., Thorvaldsson, H., Winckler, W., , Lander, E. S., Getz, G., et al. (2011). Integrative genomics viewer. *Nat. Biotechnol.* 29, 24–26. doi: 10.1038/nbt.1754

Shannon, P., Markiel, A., Ozier, O., Baliga, N. S., Wang, J. T., Ramage, D., et al. (2003). Cytoscape: a software environment for integrated models of biomolecular interaction networks. *Genome Res.* 13, 2498–2504. doi: 10.1101/ gr.1239303

Sievers, F., Wilm, A., Dineen, D., Gibson, T. J., , Li, W., et al. (2011). Fast, scalable generation of high-quality protein multiple sequence alignments using clustal omega. *Mol. Syst. Biol.* 7:539. doi: 10.1038/msb.2011.75

Song, J. M., Xie, W. Z., Wang, S., Guo, Y. X., Kudrna, D., et al. (2012). Two gap-free reference genomes and a global view of the centromere architecture in rice. *Mol. Plant* 14, 1757–1767. doi: 10.1016/j.molp.2016.01.018

Spies, D., and Ciaudo, C. (2015). Dynamics in transcriptomics: advancements in RNA-seq time course and downstream analysis. *Comput. Struct. Biotechnol. J.* 13, 469–477. doi: 10.1016/j.csbj.2015.08.004

Stamatakis, A. (2014). RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* 30, 1312–1313. doi: 10.1093/bioinformatics/btu333

Stein, J. C., Yu, Y., Copetti, D., Zweck, D. J., Zhang, L., Zhang, C., et al. (2018). Genomes of 13 domesticated and wild rice relatives highlight genetic conservation, turnover and innovation across the genus *Oryza*. *Nat. Genet.* 50, 285–296. doi: 10.1038/s41588-018-0040-0

Su, W., Ou, S., Hufford, M. B., and Peterson, T. (2021). A tutorial of EDTA: extensive de novo TE annotator. *Methods Mol. Biol.* 2250, 55–67. doi: 10.1007/ 978-1-0716-1134-0_4

Tarlo-Graovac, M., and Chen, C. (2009). Using repeatmasker to identify repetitive elements in genomic sequences. *Curr. Protoc. Bioinformatics* 4:10. doi: 10.1002/0471230993.hb0410s25

Thomas, J., and Kukushkin, S. (2009). Understanding of gene regulation during the formation of nodule-like structures in rice. *Plant Mol. Biol.* 103, 113–128. doi: 10.1007/s11103-009-0978-0

Tian, Z., Qian, Q., Liu, Q., Yan, M., Liu, X., Yan, C., et al. (2009). Allelic diversities in rice starch biosynthesis lead to a diverse array of rice eating and cooking qualities. *Plant Physiol.* 149, 184, 3542–3558 e16. doi: 10.1016/j.cell.2021.04.046

Walker, B. J., Abell, T., Ciaudo, C., and Kukushkin, S. (2014). Pilong: an integrated tool for comprehensive microbial variant detection
Wang et al. Improved Genome Assembly of 93-11 and genome assembly improvement. *PLoS One* 9:e112963. doi: 10.1371/journal.pone.0112963

Wang, J. C., Xu, H., Zhu, Y., Liu, Q. Q., and Cai, X. L. (2013). OsbZIP58, a basic leucine zipper transcription factor, regulates starch biosynthesis in rice endosperm. *J. Exp. Bot.* 64, 3453–3466. doi: 10.1093/jxb/ert187

Wang, P., Luo, Y., Huang, J., Gao, S., Zhu, G., Dang, Z., et al. (2020). The genome evolution and domestication of tropical fruit mango. *Genome Biol.* 21:60. doi: 10.1186/s13059-020-01959-8

Wang, Y., Tang, H., Debarry, J. D., Tan, X., Li, J., Wang, X., et al. (2012). MCScanX: a toolkit for detection and evolutionary analysis of gene synteny and collinearity. *Nucleic Acids Res.* 40:e49. doi: 10.1093/nar/gkr1293

Wu, S., Gao, S., Wang, S., Meng, J., Wickham, J., Luo, S., et al. (2020). A reference genome of bursaphelenchus mucronatus provides new resources for revealing its displacement by pinewood nematode. *Genes (Basel)* 11:570. doi: 10.3390/genes11050570

Yang, Z. (2007). PAML 4: phylogenetic analysis by maximum likelihood. *Mol. Biol. Evol.* 24, 1586–1591. doi: 10.1093/molbev/msm088

Yu, J., Hu, S., Wang, J., Wong, G. K., Li, S., Liu, B., et al. (2002). A draft sequence of the rice genome (*Oryza sativa* L. ssp. indica). *Science* 296, 79–92. doi: 10.1126/science.1068037

Zeng, D., Yan, M., Wang, Y., Liu, X., Qian, Q., and Li, J. (2007). Du1, encoding a novel Prp1 protein, regulates starch biosynthesis through affecting the splicing of Wab pre-mRNAs in rice (*Oryza sativa* L.). *Plant Mol. Biol.* 65, 501–509. doi: 10.1007/s11103-007-9186-3

Zhang, C., Zhu, J., Chen, S., Fan, X., Li, Q., Lu, Y., et al. (2019). Wxlv, the ancestral allele of rice waxy gene. *Mol. Plant* 12, 1157–1166. doi: 10.1016/j.molp.2019.05.011

Zhang, Q., Liang, Z., Cui, X., Ji, C., Li, Y., Zhang, P., et al. (2018). N6-Methyladenine DNA Methylation in Japonica and Indica rice genomes and its association with gene expression, plant development, and stress responses. *Mol. Plant* 11, 1492–1508. doi: 10.1016/j.molp.2018.11.005

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