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Year: 2017

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Abstract: Finger millet (Eleusine coracana (L.) Gaertn) is an important crop for food security because of its tolerance to drought, which is expected to be exacerbated by global climate changes. Nevertheless, it is often classified as an orphan/underutilized crop because of the paucity of sci- entific attention. Among several small millets, finger millet is considered as an excellent source of essential nutrient elements, such as iron and zinc; hence, it has potential as an alternate coarse cereal. However, high-quality genome sequence data of finger millet are currently not available. One of the major problems encountered in the genome assembly of this species was its polyploidy, which hampers genome assembly compared with a diploid genome. To over- come this problem, we sequenced its genome using diverse technologies with sufficient cover- age and assembled it via a novel multiple hybrid assembly workflow that combines next-generation with single-molecule sequencing, followed by whole-genome optical mapping using the Bionano IrysVR system. The total number of scaffolds was 1,897 with an N50 length >2.6 Mb and detection of 96% of the universal single-copy orthologs. The majority of the homeologs were assembled separately. This indicates that the proposed workflow is applicable to the as- sembly of other allotetraploid genomes.

DOI: https://doi.org/10.1093/dnares/dsx036

Posted at the Zurich Open Repository and Archive, University of Zurich ZORA URL: https://doi.org/10.5167/uzh-139393 Published Version



Originally published at:

Hatakeyama, Masaomi; Aluri, Sirisha; Balachadran, Mathi Thumilan; Sivarajan, Sajeevan Radha; Patrignani, Andrea; Grüter, Simon; Poveda, Lucy; Shimizu-Inatsugi, Rie; Baeten, John; Francoijs, Kees-Jan; Nataraja, Karaba N; Reddy, Yellodu A Nanja; Phadnis, Shamprasad; Ravikumar, Ramapura L; Schlapbach, Ralph; Sreeman, Sheshshayee M; Shimizu, Kentaro K (2017). Multiple hybrid de novo genome assembly of finger millet, an orphan allotetraploid crop. DNA Research:Epub ahead of print. DOI: https://doi.org/10.1093/dnares/dsx036 **Full Paper**

Multiple hybrid *de novo* genome assembly of finger millet, an orphan allotetraploid crop

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Edited by Dr. Sachiko Isobe

Received 6 May 2017; Editorial decision 16 August 2017; Accepted 23 August 2017

Abstract

Finger millet (*Eleusine coracana* (L.) Gaertn) is an important crop for food security because of its tolerance to drought, which is expected to be exacerbated by global climate changes. Nevertheless, it is often classified as an orphan/underutilized crop because of the paucity of scientific attention. Among several small millets, finger millet is considered as an excellent source of essential nutrient elements, such as iron and zinc; hence, it has potential as an alternate coarse cereal. However, high-quality genome sequence data of finger millet are currently not available. One of the major problems encountered in the genome assembly of this species was its polyploidy, which hampers genome assembly compared with a diploid genome. To overcome this problem, we sequenced its genome using diverse technologies with sufficient coverage and assembled it via a novel multiple hybrid assembly workflow that combines next-generation with single-molecule sequencing, followed by whole-genome optical mapping using the Bionano Irys® system. The total number of scaffolds was 1,897 with an N50 length >2.6 Mb and detection of 96% of the universal single-copy orthologs. The majority of the homeologs were assembled separately. This indicates that the proposed workflow is applicable to the assembly of other allotetraploid genomes.

Key words: hybrid de novo assembly, finger millet, allotetraploid genome, whole-genome optical mapping

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1. Introduction

After sorghum and pearl millet, finger millet (Eleusine coracana (L.) Gaertn) is the coarse cereal that is most widely cultivated in dry regions under rain-fed conditions. Finger millet is an allotetraploid (2n $4 \times = 36$) with an AABB genome that belongs to the family Poaceae and subfamily Chloridoideae. It was domesticated from the wild allotetraploid E. africana in Uganda to Ethiopia more than 5000 years ago. It is widely accepted that the A genome progenitor is the wild diploid species E. indica $(2n = 2 \times = 18)$ or one of its close relatives, including E. tristachya, while the B genome progenitor is unknown or most likely extinct.^{1,2,3,4} Finger millet is one of the richest sources of mineral elements such as calcium, iron, manganese, and zinc, in addition to being a fairly good source of vitamins and several essential amino acids.⁵ Among all millets, finger millet is often considered as the best source of lysine.⁶ With its high satiety value and low glycemic index, finger millet has excellent potential as a nutraceutical cereal. It is grown widely, including in India, many African countries, China, and, historically until the present day, even in Japan.⁷

Despite its importance, very little research emphasis has been given to finger millet; hence, this species is classified as an orphan or underutilized crop.^{7,8} The major reasons for this classification include a distant relationship to any of the major cereals, a large genome size, and particularly, its tetraploidy. Thus, limited efforts have been made toward the generation of genomic resources for this species, including sequencing of its genome. The genome size of finger millet was estimated to be 3.34-3.87 pg (2C),^{7,9} which corresponds to 1.63-1.89 Gb in haploid (1C) cells assuming that 1 pg DNA corresponds to 978 Mbp with a GC content of 50%. Several attempts have also been made to prospect novel genes linked to the droughtstress acclimation response,¹⁰⁻¹³ and a small number of SSR markers for the construction of genetic linkage maps have been reported.¹⁴ The EcNAC1 gene of E. coracana, which encodes a NAC family transcriptional factor, was shown to be induced by drought and to confer resistance to drought and other stresses when expressed in tobacco.¹⁰ However, compared with the progress made in generating genomic information in sorghum and pearl millet, efforts in finger millet remain scarce. The absence of such genomic resources has been the major impediment in improving finger millet through the adoption of molecular breeding approaches.

The genome assembly and annotation of polyploid species has been a major challenge in genomics, because homeologous genes (i.e. duplicated genes derived from genome duplication) are by definition highly similar and their separate assembly is difficult. About 35% of vascular plants are estimated to be recent polyploids,^{15,16} and it has long been known that important crop species are polyploids.¹⁷ However, the genome assemblies of only a handful of polyploid species have been reported. More importantly, the quality of the polyploid assembly, in particular the scaffold length, tends to be low in such reports. The presence of duplicated genes and polyploidy split the assembly, thus leading to shorter scaffolds and contigs.^{18,19}

Some polyploid species have, however, been assembled; e.g. *Nicotiana tabacum*,²⁰ *Gossypium hirsutum*,²¹ *Triticum aestivum*,²² and *Brassica juncea*.²³ The genome assembly of hexaploid wheat, *Triticum aestivum* (TGACv1),²² was performed via chromosome separation, albeit with an N50 of several kb.²² A combined approach of PacBio long reads and Bionano Genomics optical genome mapping succeeded in assembling the genome of *Brassica juncea*, which is an allotetraploid, with an N50 scaffold that was 1.5 Mb long.²³ Subsequently, pseudochromosomes were constructed using the diploid genome and genetic linkage maps. However, in previous polyploid assemblies, little was reported on the validation of the

separation of homeologs in cases with unknown parental species. Another potential assembly strategy that can be used when all the parental species are known is the assembly of the genome sequences based on the parental diploid species. This was employed for the allotetraploid Arabidopsis kamchatica using the two parental species A. halleri and A. lyrata;^{24,25} however, this cannot be simply applied to finger millet because one of the parental species is unknown. Recently, the rubber tree genome was assembled using Dovetail Chicago method²⁶ (Dovetail Genomics, LLC, Santa Cruz, CA, USA) with a scaffold N50 of 96.8 kb, and they revealed the paleotetraploidy of the genome. The Chicago library is a longer range technique than mate-pair library, which is based on chromosome conformation capture sequencing (Hi-C) protocol.27 The barley genome was assembled in chromosome length scale with Hi-C library.²⁸ NRGene Ltd (Ness-Ziona, Israel) has succeeded recently in assembling some crop genomes, e.g. the genome of maize with a very high scaffold N50 of 35.5 Mb, using the DeNovoMAGICTM software (http://nrgene.com/products-technology/denovomagic/).²⁹ They require some PCR-free paired-end libraries and mate-pair libraries for the DeNovoMAGICTM, but the company has not reported its algorithm and strategy in detail.

Here, we report the whole-genome draft sequencing and assembly of a tetraploid finger millet, cultivar PR202 (IC: 479099), using a novel polyploid genome assembly workflow. This cultivar was chosen for this work considering the extent of its adaptability. A selection from the Godavari regions of Southern India was chosen for this study based on its resilience to drought and high-temperature stresses. This cultivar is being used as the national check in all of the multilocation yield-evaluation trials performed by the All India Coordinated Small Millets Improvement Project (AICSMIP) (http:// millets.res.in/aicrp_small.php).³⁰

2. Materials and methods

Seeds of finger millet (variety name, PR202; IC number, 479099; the IC number is the indigenous collection number issued by the National Bureau of Plant Genetic Resources, India) that were stocked at the Project Coordinating Unit (Small Millets), University of Agricultural Sciences, GKVK, Bangalore, India) were used for germination. DNA was extracted from young leaves of finger millet plants grown in a plant chamber at 20 °C to 22 °C, 3 weeks later for sequencing and 2 months later for genome scanning. In this section, the methods used for DNA and RNA isolation and preparation of libraries for sequencing are described.

2.1. Preparation of DNA libraries for sequencing

Good-quality and high-molecular-weight genomic DNA (gDNA) was isolated using a combination of CTAB and Genomic tip 500/G column (Qiagen AG, Hombrechtikon, Switzerland) purification method, in accordance with the protocol 'Preparing Arabidopsis Genomic DNA for Size-Selected ~20 kb SMRTbellTM Libraries' (http://www.pacb.com/ wp-content/uploads/2015/09/Shared-Protocol-Preparing-Arabidopsis-DNA-for-20-kb-SMRTbell-Libraries.pdf). The DNA pellet obtained using the conventional CTAB method was dissolved in 1 ml of Dnaseand RNase-free distilled water (Life Technologies, Carlsbad, CA, USA) and purified by passing through Genomic tip 500/G columns, to obtain intact DNA with a size above 60 kb. For library preparation, the TruSeq DNA NanoSample Prep Kit v2 (Illumina Inc., San Diego, CA, USA) was used in the following steps. DNA samples (1 µg) were sonicated on a Covaris instrument (Covaris, Woburn, MA, USA)

using settings specific for a fragment size of 550 bp. The fragmented DNA samples were size selected using AMPure beads (Beckman Coulter, Inc., Brea, CA, USA), end-repaired, and polyadenylated. TruSeq adapters containing the index for multiplexing were ligated to the fragmented DNA samples. Fragments containing TruSeq adapters at both ends were selectively enriched by PCR. The quality and quantity of the enriched libraries were validated using Qubit[®] 2.0 Fluorometer (Life Technologies) and TapeStation system (Agilent, Waldbronn, Germany). The products were a smear with an average fragment size of approximately 700 bp. The libraries were normalized to 10 nM in Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20. The 3-8 and 20kb mate-pair libraries were prepared according to Lucigen's NxSeq® Long Mate Pair Kit protocol and NxSeq® 20kb Mate Pair Library kit protocol (Lucigen Corporation, Middleton, WI, USA), respectively. Briefly, genomic DNA was sheared using Covaris G-TUBES (Covaris, Woburn, MA, USA) to the desired size (3, 5, 8, and 20 kb), end-repaired, A-tailed, and ligated to adaptors. The insert was size selected and ligated to a unique coupler that contained encrypted Chimera CodeTM sequences. The circularized inserts were treated with exonuclease for the removal of unwanted linear DNA and digested with a selection of endonucleases to produce correctly sized di-tags (400-900 bp). Biotin capture was used to remove unwanted DNA fragments prior to the addition of the Junction CodeTM Reagent. The library was recircularized and PCR amplified and the quality was assessed using a Qubit® 2.0 Fluorometer (Life Technologies), and the TapeStation system (Agilent, Waldbronn, Germany). The final matepair libraries were normalized to 10 nM in Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20.

2.2. Preparation of RNA libraries for sequencing

RNA was extracted from young leaves of finger millet 1 month after germination. Freshly germinated finger millet plants were exposed to drought stress, in which the plants were not watered for 7 days. Tissue samples were taken from the young leaves before and after the exposure to drought conditions. RNA was extracted using an RNeasy plant kit (Qiagen AG, Hombrechtikon, Switzerland). The quantity and quality of the isolated RNA was determined using a Qubit® 2.0 Fluorometer (Life Technologies) and a Bioanalyzer 2100 (Agilent, Waldbronn). For library preparation, TruSeq Stranded mRNA Sample Prep Kit (Illumina) was used in the subsequent steps. Briefly, total RNA samples (1 µg) were ribosome depleted and then reverse transcribed into double-stranded cDNAs, with addition of actinomycin during first-strand synthesis. The cDNA samples were fragmented, end-repaired, and polyadenylated before the ligation of TruSeq adapters. The adapters contained the index for multiplexing. Fragments containing TruSeq adapters at both ends were selectively enriched with PCR. The quality and quantity of the enriched libraries were assessed using a Qubit® 2.0 Fluorometer and a Bioanalyzer 2100 (Agilent, Waldbronn). Products were a smear with an average fragment size of ~360 bp. The libraries were normalized to 10 nM in Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20.

2.3. Library preparation and sequencing for PacBio RS II

The SMRT Bell library was produced using a DNA Template Prep Kit 1.0 (PacBio p/n 100-259-100). The concentration of the input gDNA was measured using a Qubit Fluorometer dsDNA Broad Range assay (Life Technologies p/n 32850). gDNA (10μ g) was mechanically sheared to an average size distribution of 10 kb using a

Covaris gTube (Kbiosciences p/n 520079). A Bioanalyzer 210012K DNA Chip assay (Agilent p/n 5067-1508) was used to assess the fragment size distribution. Sheared gDNA (5 µg) was DNA-damage repaired and end-repaired using polishing enzymes. A blunt-end ligation reaction followed by exonuclease treatment was performed to create the SMRT Bell template. A Blue Pippin device (Sage Science, Inc., Beverly, MA, USA) was used to size select the SMRT Bell template and enrich for large fragments (>10 kbp). The size-selected library was quality inspected and quantified on Agilent Bioanalyzer 12kb DNA Chip and Qubit Fluorimeter (Life Technologies). A ready-to-sequence SMRT Bell-Polymerase Complex was created using a P6 DNA/Polymerase Binding Kit 2.0 (PacBio p/n 100-236-500), according to the manufacturer's instructions. The PacBio RS II instrument was programmed to load and sequence the sample on PacBio SMRT cells v3.0 (PacBio p/n 100-171-800), acquiring one movie of 360 min per SMRT cell. A MagBead loading (PacBio p/n 100-133-600) method was chosen to improve the enrichment of the longer fragments. After the run, a sequencing report was generated for each cell via the SMRT portal, to assess adapter dimer contamination, sample-loading efficiency, average read length, and number of filtered sub-reads.

2.4. Next-generation optical mapping

A Bionano PrepTM Plant Tissue DNA Isolation Kit was used according to the Bionano PrepTM Plant Tissue DNA Isolation Base Protocol (Document number: 30068, Document revision: B, Bionano Genomics, Inc., San Diego, CA, USA). Briefly, 1g of young leaves collected 2 months after germination was fixed in formaldehyde, to protect the nuclei and DNA against mechanical shearing. The leaves were then homogenized with a rotor-stator apparatus, to break the tough plant cell wall, followed by a density gradient purification for the recovery of intact nuclei. These samples were embedded in lowmelt agarose plugs for proteinase K (Qiagen AG, Hombrechtikon, Switzerland) and RNase A (Qiagen AG, Hombrechtikon) digestion and subsequent purification of high-molecular-weight (HMW) DNA. The resulting HMW DNA was processed further according to the IrysPrepTM Labeling-NLRS User Guide (Document number: 30024, Document revision: G, Bionano Genomics) using an NLRS Kit (Bionano Genomics). In brief, 300 ng of HMW DNA was nicked by Nt.BspQI (New England Biolabs, Ipswich, MA, USA) and labeled. These reactions provided sequence specificity for an optimal label density of 8-15 labels per 100 kbp. The nick enzyme Nt.BspQI was selected by predicting the label density as 11.9 labels per 100 kbp using Bionano Genomics Knickers (ver. 1.5.5.0, http://www. bnxinstall.com/knickers/Knickers.htm) beforehand. The labeled nicks were repaired to restore strand integrity, and the labeled DNA was stained for backbone visualization. Labeled nicks were subsequently detected as dots on a string when run on a Bionano Irys® system instrument.

2.5. Sequencing and optical mapping

Sequencing was performed on an Illumina NextSeq500, MiSeq instrument (Illumina) and a PacBio RS II system (Pacific Bioscience of California, Inc., CA, USA). Genome optical mapping was carried out on a Bionano Irys[®] system (Bionano Genomics). The three pairedend libraries with insertion lengths of 250, 550, and 700 bases were sequenced on an Illumina NextSeq500 apparatus (reagent version 2 kits 2×150 cycles). In contrast, the four mate-pair libraries with insert sizes of 3, 5, 8, and 20 kb were sequenced on an Illumina MiSeq instrument (reagent version 3 kits 2×300 cycles). Long reads were obtained from PacBio RS II using a total of 23 SMRT cells. The nicked-labeled HMW DNA was run on a Bionano IrysChip[®] flow-cell that scanned 6 times for $96.5 \times$ coverage.

2.6. Genome size estimation

Approximately 0.5 cm² of fresh leaf tissue of finger millet, together with an equal amount of leaf tissue of tomato (*Lycopersicon esculen-tum*; used as an internal standard [diploid genome size, 1.96 pg³¹]), was chopped with a sharp razor blade in extraction buffer (CyStain PI Absolut P; Sysmex Europe GmbH, Norderstedt, Germany). The suspension of nuclei was filtrated and incubated in staining solution containing propidium iodide (PI) and RNase A (CyStain PI Absolut P; Sysmex). The suspension of stained nuclei was analyzed using a flow cytometer (CyFlow Space, Sysmex).

2.7. Hybrid *de novo* assembly workflow and gene annotation

The hybrid *de novo* assembly workflow is shown in Fig. 1. Adapters that contaminated the sequenced paired-end reads and low-quality bases were trimmed using Trimmomatic (ver. 0.33)³² on the SUSHI framework³³ after sequencing. Regarding the mate-pair libraries, first, chimera reads were filtered out, and chimera code and linker sequences from the 5' end were trimmed using the script



Figure 1. Hybrid de novo assembly workflow. In the figure, the white box represents a process, the gray box represents a data entity, the blue solid arrow shows the process flow, and the black dashed arrow shows the data flow. The hybrid de novo assembly was executed as follows: (1) contig assembly using only paired-end reads (Platanus ver. 1.2.4), (2) hybrid contig assembly using contigs >1 kb from the first step and long reads (DBG2OLC commit id: 7f1712ae76b015b1f4efee81a5d4e1305701197a), (3) polishing of the assembled contig with paired-end reads (Pilon ver. 1.21) (NGS contigs), (4) using the polished contigs in the two subsequent independent steps, (4-1) scaffolding of the polished contigs with mate-pair reads (NGS scaffolds), (4-2) after converting scanned molecules into a CMAP file and optical genome mapping to the NGS contigs, (5) de novo assembly of scanned molecules based on the parameters of the molecular-quality report and autonoise (autodetection of appropriate noise parameters) with the NGS contigs (BNG contigs), (6) hybrid scaffolding with both NGS scaffolds (step 4-1 output) and BNG contigs (step 5 output), and (7) gap closing (GMclose ver. 1.6) of the scaffolds with all the contigs generated in the first step (step 1 output). Finally, the workflow produced the super scaffolds. The raw data preprocesses, such as adapter trimming and removing chimera code from matepair reads, are not shown in this workflow.

IlluminaChimera-Clean5.py provided by Lucigen Co., Wisconsin, USA, followed by another script, IlluminaNxSeqJunction-Split9.py, to segregate the reads into proper mate-pair reads and unsplit reads via the detection of junction sequences. Subsequently, Platanus (ver. 1.2.4)³⁴ was executed using the preprocessed paired-end reads to assemble contigs (Fig. 1, step 1), which were then used as anchors to connect the PacBio long reads on DBG2OLC (commit id: 7f1712ae76b015b1f4efee81a5d4e1305701197a)³⁵ (Fig. 1, step 2). Contigs from the Platanus assembly that were longer than 1 kb were used for the input of the DBG2OLC hybrid assembly. Note that many short contigs were filtered out to adopt a conservative approach aimed at avoiding misassembly, as short and fragmented contigs stemmed most probably from complex regions, such as repeats and duplicates. The contigs from the DBG2OLC hybrid assembly were polished using Pilon (ver. 1.21)³⁶ with the accurate Illumina short reads, to correct misassemblies (NGS contigs) (Fig. 1, step 3). We used only half amount of the paired-end reads (250 base insert size) for the polishing. Subsequently, the polished contigs were scaffolded with the mate-pairs using SSPACE standard (ver. 3.0)³⁷ (NGS scaffolds) (Fig. 1, step 4-1). We only used the proper mate-pair reads for scaffolding. Independently, optical genome mapping was performed on a Bionano Irys[®] system (Fig. 1, step 4-2), followed by *de novo* assembly with the detected molecules using the IrysView® and Bionano IrysSolve® software (ver. 2.1) (BNG contigs) (Fig. 1, step 5). The de novo assembly was performed twice: the first de novo assembly used the estimated nose parameters by mapping the detected molecules to the NGS contigs, and the second assembly used autonoise (autodetection of appropriate noise parameters). The hybrid scaffolding was performed using Bionano IrysSolve® (ver. 2.1) with both the NGS scaffolds and the BNG contigs. Finally, GMclose (ver. 1.6)³⁸ filled the gaps in the super-scaffolds using all the contigs assembled by Platanus. The actual executed commands as a shell script with configuration files and parameters are provided in Supplementary File S1.

The genes were predicted and annotated using MAKER (ver. 2.31.8)³⁹ based on the rice genome (Oryza sativa ssp. japonica IRGSP-1.0, ENA Assembly: GCA_001433935.1) and the RNAseq data that were obtained from young leaves. The rice cDNAs and RNAseq data for the EST evidence and the rice proteins for the protein homology evidence were used for the MAKER job. To render the GFF file, the input of MAKER, the RNAseq data were mapped to the assembled reference sequences using STAR aligner (ver. 2.5.2b)⁴⁰ and the GFF annotation file was constructed via transcriptome construction using StringTie (ver. 1.3.3b).⁴¹ The rice-specific repeat-making library obtained from Repbase database⁴² was used for RepeatMasker (ver. 4.0.5).⁴³ The rice model was used for Augustus (ver. 3.2.0)⁴⁴ gene prediction in the MAKER pipeline. The detail options are written in the Supplementary File S1. For assessing the quality of the genome assembly and annotation, we executed BUSCO (ver. 2.0.1)⁴⁵ using the published plant universal single-copy orthologs. We also performed an RNAseq data alignment to the final assembled genome using STAR $(\text{ver. } 2.5.2\text{b})^{40}$ and checked the mapping statistics.

3. Results and discussion

In this section, we report the genome assembly and discuss the results of its validation.

3.1. Sequencing and assembly statistics

Using the flow cytometry experiment, we estimated that the size of the finger millet genome is 1.5 Gb. In total, short reads amounting to

140× coverage and long reads amounting to 16× coverage were sequenced. We scanned the DNA molecules on a Bionano Irys[®] system and detected 144,761 Mb molecules in a total of six runs (Supplementary Table S1). Based on the estimated genome size, the sequenced coverage depth for each library is reported in Table 1 (PacBio subreads length distribution and calculated insert size of distribution are shown in Supplementary Figs S1 and S2, respectively).

The statistics of each step in the hybrid assembly are shown in Table 2. The total assembled size was 1,188,784,944 bases, which is 78.2% of the genome size estimated by flow cytometry. The hybrid contig assembly by DBG2OLC and the scaffolding by SSPACE enhanced the continuity from the Platanus contig assembly, although the mate-pair sequenced coverage was less than $10\times$, and the total assembly size decreased compared with the Platanus contig assembly. This could be explained by the low sequencing coverage of the PacBio long reads, rather than the filtering out of the shorter Platanus contigs. At the final hybrid scaffolding and gap closing, the N50 of the final super-scaffolds increased to 2.68 Mb, which is approximately twice as long as the NGS scaffolds. The hybrid scaffolding generated super-scaffolds that concatenated the NGS scaffolds based on the BNG contigs that were assembled from the optical genome mapping. Some conflicts were detected between NGS scaffolds and BNG contigs, which split the conflicting NGS scaffolds into smaller pieces. This had the major advantage of correcting the polyploid genome assembly, as scaffolding with only mate-pair short

reads may fail to solve the long-range homeologous complexity and may sometimes lead to mis-scaffolding.^{46,47}

Currently, the so-called de novo assembly method based on nextgeneration and third-generation sequencing technologies is classified mainly into two approaches: (i) de Bruijn graph (DBG) assembly and (ii) overlap-layout consensus (OLC) assembly.48,49 Although the DBG method is more efficient computationally regarding both memory and calculation time, the fragment size parameter, usually called k-mer length, critically influences the results and often causes a misassembly because of the small fragment k-mers on homologous or repetitive genome regions. Some assemblers, such as Platanus,³⁴ solve this problem by using multiple k-mer sizes for the construction of the DBG. As another solution for the homeologous complexity, longer reads produced from a single-molecule sequencer combined with the OLC method may be beneficial. However, high-coverage sequencing of long reads may solve and correct the wrongly sequenced bases, albeit at a high cost. The hybrid de novo assembly described in this article, which uses both accurate short reads and inaccurate long reads with a low coverage depth, represents another solution regarding assembly cost and calculation efficiency; moreover, the beneficial feature of long reads can be applied to solve the homeologous complexity encountered during polyploid genome assembly. The classification of reads via homeolog phasing⁵⁰ based on long reads or the parental diploid genome may improve the polyploid assembly further. In addition, further longer molecule information from such

Table 1. Raw sequence data

Library type	DNA RNA	Sequencer	Insert length (bases)	Number of bases	Coverage depth
Paired-end	DNA	Illumina NextSeq	250	93,783,186,606	63×
Paired-end	DNA	Illumina NextSeq	550	63,519,282,198	42×
Paired-end	DNA	Illumina NextSeq	700	50,551,135,834	34×
Mate-pair	DNA	Illumina MiSeq	3,000	789,059,46	$0.053 \times$
Mate-pair	DNA	Illumina MiSeq	5,000	364,070,088	$0.24 \times$
Mate-pair	DNA	Illumina MiSeq	8,000	1,138,183,671	$0.76 \times$
Mate-pair	DNA	Illumina MiSeq	20,000	1,366,910,280	$0.91 \times$
PacBio CLR	DNA	PacBio RS II	-	25,086,179,201	$16.7 \times$
Paired-end	RNA	Illumina HiSeq4000	_	148,856,244,082	99.2×

The number of sequenced raw data in each library and the coverage depth based on the genome size, 1.5 Gb as estimated by flow cytometry, are shown.

Table 2. Assembly statistics at each assembly step

Statistics	Contig assembly	Hybrid assembly	Scaffold	Hybrid scaffold
Total sequences	2,812,919	6,374	2,387	1,897
Total bases	1,307,217,455	1,067,045,564	1,069,478,541	1,188,784,944
Min sequence length	115	3,408	3,911	1,244
Max sequence length	27,802	2,072,336	5,237,708	13,553,037
Average sequence length	464.72	167,405.96	448,042.96	626,665.76
Median sequence length	154.00	102,084.50	237,045.00	92,178.00
N25 length	3,419	520,874	1,581,824	5,029,714
N50 length	1,410	285,549	905,318	2,683,090
N75 length	311	145,770	457,768	1,232,573
N90 length	133	76,519	216,067	419,121
(G + C)s	43.15%	43.83%	43.57%	40.98%
Ns	0.00%	0.00%	0.73%	6.63%

Assembly statistics at each step are summarized. Contig assembly represents the contigs that were assembled using Platanus with preprocessed paired-end reads. Hybrid assembly represents the contigs that were generated using DBG2OLC with the Platanus contigs (> 1 kb long) and PacBio long reads. Scaffold is the result of the SSPACE standard scaffolding with mate-pair reads, and Hybrid scaffold is the hybrid scaffolding result with both Bionano Genomics de novo assembled contigs and the SSPACE scaffolds, followed by GMclose gap closing. as Bionano Irys[®] system is very promising, not only for superscaffolding, but also for solving the long-range homeologous complexity in polyploid genome assembly, as demonstrated by the results reported in this article.

3.2. Gene annotation and assembly validation

After the hybrid assembly, we conducted gene annotation prediction using MAKER and 62,348 genes were predicted. The distribution of Annotation Edit Distance (AED)³⁹ tagged by MAKER is shown in Fig. 2A, in which more than 91% of the annotated genes (57,066 genes among a total of 62,348 genes) show an AED <0.5. This result indicates that the annotation is well supported by the evidence. For the assessment of the quality of the assembly and annotation, the assembled scaffolds and predicted genes were analyzed on BUSCO (Fig. 2B). In the assembled scaffolds (shown in Fig. 2B, 'genome'), 96.5% of the universal single-copy genes (1,389 genes among a total of 1,440 genes) were identified, supporting the quality of the assembly. Among the 1,389 single-copy genes detected, 606 genes were detected as being complete single-copy genes, 783 genes were found to be duplicates, and 766 of the 783 duplicates were exactly twice found from two different scaffolds. An NCBI BLAST search of the complete single-copy genes (606) that were detected only once in the assembled genome found an additional 556 homologous partial genes with a shorter alignment length (40% of the 1,389 complete single-copy genes detected). This indicates that the majority of the homeologous genes were assembled separately (see section 3.3). In tetraploid species, two copies may be found for each 'single-copy gene', although duplicated genes may be gradually lost because of redundancy (called nonfunctionalization or homeolog loss).51,52 Similar results were obtained from the BUSCO analysis of proteins and transcripts (Fig. 2, 'proteins' and 'transcripts'), but the missing and fragmented proportion was increased compared with the 'genome' analysis. It is well known that one of the homeologs may not be expressed (or silenced) after polyploidization, because of redundancy; thus, some genes were not annotated and the missing and fragmented proportion increased among transcripts and proteins because of the lack of evidence. Moreover, we searched the predicted proteins in the Pfam database⁵³ and found that 74% of predicted proteins were present in this database (Supplementary Fig. S3). RNAseq data obtained using other tissues and conditions may improve the gene annotation process.

Furthermore, we aligned the data from six RNAseq sequenced libraries to the assembled genome for validation: 95.9% of the reads were mapped, which indicates that the assembled genome may cover more than 95% of the coding region (STAR alignment statistics is shown in Supplementary Table S2).

3.3. Polyploid assembly validation and phylogenetic analysis

To validate the quality of the polyploid assembly, i.e. to determine whether the homeologs were assembled in different scaffolds, we analyzed our assembly data of three genes for which Sanger sequencing data of E. coracana had been already reported. First, we performed an NCBI BLAST (ver. 2.2.31+)⁵⁴ search of the published low-copynumber marker phosphoenolpyruvate carboxylase 4 $(Pepc4)^1$ and granule-bound starch synthase I-like $(waxy)^2$ in the assembled scaffolds. The reciprocal BLAST best-hit search with a very low E-value threshold identified exactly two scaffolds with two different completely matching sequences from different clades (the NCBI BLAST search result is shown in Supplementary File S2). Second, we conducted a molecular phylogenetic analysis using MEGA4 (ver. 7.0.26)⁵⁵ based on the detected low-copy-number homeologs with the published sequences of close relatives¹ (Supplementary Fig. S4). As shown in these figures, the homeologs detected were clearly separated in different clusters. This result indicates that, in these gene regions, the homeologs were successfully assembled, regardless of the



Figure 2. Annotation edit distance (AED) and benchmark of universal single-copy orthologs. (A) The distribution of AED tagged as a gene by MAKER. (B) Benchmark result of the plant set of universal single-copy orthologs produced by BUSCO. In each bar, the pale-blue color indicates the number of genes that were detected completely as single-copy genes in the genome, the dark-blue color indicates the number of genes that were detected as single-copy genes but were counted more than twice, the yellow color indicates the number of genes that were detected as single-copy genes but not completely, and the red color indicates the number of single-copy genes that were not detected among the plant universal single-copy orthologs. The 'genome' is the BUSCO result that was obtained by searching the assembled genome FASTA file, and the 'transcripts' and 'proteins' constitute the BUSCO result that was obtained by searching the transcript FASTA file and amino acid FASTA file produced by MAKER.

Next, we analyzed EcNAC1, which is the best-studied gene of E. coracana related to the drought response.¹⁰ NAC is one of the largest families of plant-specific transcription factors and has been shown to be involved in diverse processes, such as growth, development, and abiotic stresses. The EcNAC1 of the E. coracana GPU28 variety was induced by drought stresses and conferred tolerance to various abiotic stresses, such as simulated osmotic stress by polyethylene glycol (PEG) and mannitol and salinity stresses in the overexpression lines of tobacco.¹⁰ Based on our assembly of PR202, we found a sequence that was similar to, and clustered with, the EcNAC1 of GPU28, together with another sequence from the Trichy1 variety (GenBank accession ID: KU500625.1) (Supplementary Fig. S5). The assembly also had another copy, which is likely to be a homeolog in a different scaffold. It would be valuable to study this copy, which may well have drought-related functions.

In addition, we searched homologous genes as 'duplicated genes' in the predicted transcripts using NCBI BLAST and found 57,913 genes that had more than two copies in the genome (Supplementary File S3). Among these duplicates, we found 34,678 genes as reciprocal BLAST best-hit genes located in different scaffolds. Assuming that these genes are homeologs, we found that 56% of all 62,348 genes were considered homeologs, which is consistent with the ratio (55% (766/1389)) of duplicated single-copy genes detected exactly twice in BUSCO (the analytical process and results are shown as a shell script in Supplementary File S3). To check the location of these homeologs, we calculated the gene density (the number of genes per Mb) and the homeolog density (the number of homeologs per Mb) in each scaffold (Supplementary Fig. S6 A1-2, B1-2). Panels A1 and B1 of Supplementary Fig. S6 show the density against the scaffold length of all genes and homeologs, respectively, and panels A2 and B2 show the density distribution of all genes and homeologs, respectively, as well. The genome-wide gene density was 52.4 genes per Mb and the genome-wide homeolog density was 29.2 homeologs per Mb. As shown in these figures, all genes and homeologs were distributed similarly over the genome. Nonetheless, there was a small number of scaffolds (>1 kb long) with high homeolog density (Supplementary Fig. S6, B1), which indicates that there are areas that are dense in homeologs compared with the genome-wide homeolog distribution.

3.4. Summary of the assembly of the genome of polyploid finger millet

Over half of the world's freshwater resources are currently being used for irrigating agricultural crops. The domestic and industrial demands of the burgeoning population have led to an unprecedented withdrawal of freshwater.⁵⁶ Combined with the erratic pattern of rainfall caused by the ensuing global climate change, agriculture production is expected to be severely affected in the years to come.⁵⁷ This scenario urgently necessitates the cultivation of more water-productive crops and/or the development of improved cultivars with a superior water-use efficiency. Millets, with their C₄ photosynthetic metabolism, are generally considered to possess higher levels of drought tolerance and are generally cultivated under rain-fed conditions. Because of their higher resilience to harsh climatic conditions, millets exhibit a wider adaptation, and hence can be cultivated in a

wider range of agro-ecosystems. Therefore, millets currently represent an integral component of human diets in most impoverished areas, especially in the tropical and subtropical regions of Africa and Southeast Asia.

We assembled the genome of the allotetraploid finger millet (Eleusine coracana (L.) Gaertn, variety: PR202, IC: 479099) via a novel pipeline using various types of sequencers. The size of the assembled genome was about 1.2 Gb, while the genome size measured by flow cytometry was 1.5 Gb. Considering that the majority of the grass genomes are likely to be composed of repetitive elements, the coverage of the genome is sufficiently high. MAKER predicted 62,348 genes, which is roughly the double of the 35,679 genes identified in diploid rice (Oryza sativa Japonica IRGSP-1.0); this is also consistent with a good coverage of the genome. Three lines of evidence supported the good coverage. First, we provided a polyploid assembly with a long scaffold size (N50 length >2.5 Mb), while the assembly of polyploid genomes tends to be fragmented. It is true that the N50 may not mean correct assembly and scaffolding, but we avoided too-greedy scaffolding. Second, 96.5% of the 1,440 universal single-copy orthologs were identified, and 95.9% of RNAseq reads were mapped to the assembled genome, which supported the adequateness of the assembly. Third, the majority of these universal sets had two copies, suggesting the separation of homeologs was adequate in our assembly strategy. Concomitantly, however, we found a significant number of genes that had a single copy. There are several likely explanations for this finding. (i) One of the duplicated genes tends to be lost because of redundancy. Polyploidization was estimated to have occurred well before the domestication of millet (0.50-2.71 mya), in contrast with other crops, in which polyploidization occurred through domestication during the last thousand years.^{58,59} Thus, it is possible that many duplicated genes were lost. (ii) The published plant universal single-copy gene set is not collected by random sampling from the entire plant genes in terms of the rate of gene loss. Despite the frequent occurrence of genome duplication events throughout angiosperms, these genes tend to be lost and become single copy, potentially because of dosage balance or redundancy. (iii) The assembly size is less than the genome size estimated by flow cytometry. Most of the lacking sequences are likely to be repetitive sequences, but it is also likely that parts of the genic regions were not assembled. (iv) The assembler could not deal with two homeologs; thus, one copy was missing or two copies were combined. We verified that the two copies of the Pepc4 gene, waxy and NAC, were assembled separately. Currently, not many genome sequences from diploid E. indica are available, preventing the systematic validation of homeologs. Further improvement of assembly algorithms and validation would be valuable in this field of research. It is still possible to improve the assembly and annotation of this genome. Nevertheless, the allotetraploid finger millet genome information reported here is expected to be of much value for further genomic research of this millet at the molecular level, as well as for studies aimed at improving finger millet productivity and bioavailability. We have successfully shown the applicability of the multiple hybrid de novo assembly workflow to a complex allotetraploid genome.

Recently, the genome assembly of the *E. coracana* ML-365 variety was elucidated using Illumina and SOLiD sequencing data.⁶⁰ We emphasize that our data and analyses were targeted at the analysis of complex polyploid genomes. The N50 of our assembly (>2.5 Mb) was two orders of magnitude higher than that of ML-365 (about 24 kb) and would allow RNAseq analysis of each homeolog separately, as well as resequencing analyses. Our results illustrate the strength of single-molecule sequencing and optical genome mapping in assembling polyploid genomes.

Acknowledgements

We wish to thank Dr. Scott Monsma at Lucigen Co., Wisconsin, USA, for mate-pair library preparation, Carmen Näf at the University of Zurich for RNA extraction and library preparation, Aki Morishima for plant growth and genome size estimation by flow cytometry, Dr. Misako Yamazaki and Dr. Lucas Mohn at the University of Zurich for sample handling, and Dr. Dario Copetti at the University of Arizona for advice on gene annotation, and Dr. Jun Sese and Dr. Tony Kuo at the National Institute of Advanced Industrial Science and Technology for discussion about polyploid genome assembly. We also thank Dr. Lilian Gilgen and Damaris O'Brien at the Indo-Swiss Collaboration in Biotechnology (ISCB) office at the Federal Institute of Technology in Lausanne (EPFL), Nangsa Kamtzi and Isavel Schöchli at the University of Zurich, and Lydia Imhof and Jana Ploszaj at the Functional Genomics Center Zurich for the administrative support. Moreover, we would like to thank the members of the Evolutionary and Ecological Genomics Group at the University of Zurich and the members of Genomics/ Transcriptomics Group at the Functional Genomics Center Zurich for useful discussions and suggestions.

Availability

All sequenced raw data in FASTQ format, scanned molecular data in BNX format, *de novo* assembly results by Bionano IrysSolve[®] in CMAP format, and the final scaffolds and annotation in FASTA and GFF format are available in the DDBJ Sequence Read Archive (DRA) database with the BioProject accession number: PRJDB5606 and BioSample number: SAMD00076255.

Supplementary data

Supplementary data are available at DNARES online.

Conflict of interest

John Baeten and Kees-Jan Francoijs are employees of Bionano Genomics.

Accession numbers

BioProject accession number: PRJDB5606 BioSample number: SAMD00076255, DRA accession number: DRA005897

Funding

This project is funded by the Indo-Swiss Collaboration in Biotechnology (ISCB), a JST CREST Grant (number JPMJCR16O3), and a KAKENHI Grant (number 16H06469, 16K21727), Japan.

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