1 Twelve Platinum-Standard Reference Genomes Sequences (PSRefSeq) that

2 complete the full range of genetic diversity of Asian rice

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55 Abstract

56

57 As the human population grows from 7.8 billion to 10 billion over the next 30 years, 58 breeders must do everything possible to create crops that are highly productive and 59 nutritious, while simultaneously having less of an environmental footprint. Rice will play 60 a critical role in meeting this demand and thus, knowledge of the full repertoire of genetic 61 diversity that exists in germplasm banks across the globe is required. To meet this 62 demand, we describe the generation, validation and preliminary analyses of transposable 63 element and long-range structural variation content of 12 near-gap-free reference genome 64 sequences (RefSeqs) from representatives of 12 of 15 subpopulations of cultivated rice. When combined with 4 existing RefSeqs, that represent the 3 remaining rice 65 66 subpopulations and the largest admixed population, this collection of 16 Platinum 67 Standard RefSeqs (PSRefSeq) can be used as a pan-genome template to map 68 resequencing data to detect virtually all standing natural variation that exists in the pan-69 cultivated rice genome.

70 Background & Summary

Asian cultivated rice is a staple food for half of the world population. With the planet's 71 72 population expected to reach 10 billion by 2050, farmers must increase production by at 73 least 100 million metric tons per year (Seck et al 2012; Merrey et al. 2018). To address 74 this need, future rice cultivars should provide higher yields, be more nutritious, be 75 resilient to multiple abiotic and biotic stresses, and have less of an environmental 76 footprint (Wing et al. 2018; 3K RGP 2014). To achieve this goal, a comprehensive and 77 more in-depth understanding of the full range of genetic diversity of the pan-cultivated 78 rice genome and its wild relatives will be needed (Stein et al. 2018).

79 With a genome size of ~390 Mb, rice has the smallest genome among the 80 domesticated cereals, making it particularly amenable to genomic studies (Kawahara et 81 al. 2013) and the primary reason why it was the first crop genome to be sequenced 15 82 years ago (International Rice Genome Sequencing 2005). To better understand the full-83 range of genetic diversity that is stored in rice germplasm banks around the world, 84 several studies have been conducted using microarrays (Thomson et al. 2017; McNally et 85 al. 2009) and low coverage skim sequencing (Huang et al. 2012; Zhao et al. 2018). In 86 2018, a detailed analysis of the Illumina resequencing of more than 3,000 diverse rice 87 accessions (a.k.a. 3K-RG), aligned to the O. sativa v.g. japonica cv. Nipponbare 88 reference genome sequence (a.k.a. IRGSP RefSeq), showed how the high genetic 89 diversity present in domesticated rice populations provides a solid base for the 90 improvement of rice cultivars (Wang et al. 2018). One key finding from a population 91 structure analysis of this dataset showed that the 3,000 accessions can be subdivided into 92 nine subpopulations, where most accessions from close sub-groups could be associated to 93 geographic origin (Wang et al. 2018). 94 One critical piece of information missing from these analyses is the fact that single

95 nucleotide polymorphisms (SNPs) and structural variations (SVs) present in 96 subpopulation specific genomic regions have yet to be detected because the 3K-RG data 97 set was only aligned to a single reference genome. Therefore, the next logical step, to 98 capture and understand genetic variation pan-subpopulation-wide is to map the 3K-RG 99 dataset to high-quality reference genomes that represent each of the subpopulations of 100 cultivated Asian rice. At present, only a handful high-quality rice genomes for cultivated 101 rice are publicly available (Kawahara et al. 2013, Zhang et al. 2016a, Zhang et al. 2016b 102 and Stein et al. 2018), thus, there is an immediate need for such a comprehensive 103 resource to be created, which is the subject of this Data Descriptor.

104 Here we present a reanalysis of the population structure analysis discussed above 105 (Wang et al. 2018) and show that the 3K-RG dataset can be further subdivided into a total 106 of 15 subpopulations. We then present the generation of 12 new and near-gap-free high-107 quality PacBio long-read reference genomes from representative accessions of the 12 108 subpopulations of cultivated rice for which no high-quality reference genomes exist. All 109 12 genomes were assembled with more than 100x genome coverage PacBio long-read 110 sequence data and then validated with Bionano optical maps (Udall and Dawe 2018). The 111 number of contigs covering each of the twelve 12 assemblies, excluding unplaced 112 contigs, ranged from 15 (GOBOL SAIL (BALAM)::IRGC 26624-2) to 104 (IR 64). The 113 contig N50 value for the 12 genome data set ranged from 7.35 Mb to 31.91 Mb. When 114 combined with 4 previously published genomes (i.e. Minghui 63 (MH 63), Zhenshan 97 115 (ZS 97) (Zhang et al. 2016a, b), N 22 (Stein et al. 2018; updated in 2019) and the IRGSP 116 RefSeq (Kawahara et al. 2013)), this 16 genome dataset can be used to represent the 117 K=15 population/admixture structure of cultivated Asian rice.

118 Methods

119 Ethics statement

- 120 This work was approved by the University of Arizona (UA), the King Abdullah
- 121 University of Science and Technology (KAUST), Huazhong Agricultural University
- 122 (HZAU), the International Rice Research Institute (IRRI) and the International Center for
- 123 Tropical Agriculture (CIAT). All methods used in this study were carried out following
- 124 approved guidelines.

125 **Population structure**

- 126 We extracted 30 subsets of 100,000 randomly chosen SNPs out of the 3K-RG Core SNP
- 127 set v4 (996,009 SNPs, available at https://snp-seek.irri.org/_download.zul). For each
- subset, we ran ADMIXTURE (Alexander et al. 2009) with the number of ancestral
- 129 groups K ranging from 5 to 15. We then aligned the resulting Q matrices using CLUMPP
- 130 software (Jakobsson and Rosenberg 2007). Since different runs at a given value of K
- 131 often give rise to different refinements (splits) of the lower level grouping, we first
- 132 clustered the runs for each K according to similarity of Q matrices using hierarchical
- 133 clustering, thus obtaining several clusters of runs for each K. We discarded one-element
- 134 clusters (outlier runs), and averaged the Q matrices within each remaining cluster. Figure
- 135 S1 shows the admixture proportions taken from the averaged Q matrices of the final

136 clusters for K=9 to 15. The columns of these averaged Q matrices, representing

137 admixture proportions for groups discovered in different runs, were then used to define

the "K15" grouping. At K=9, 12, and 13, the Q matrices converged to two different

139 modes according to whether XI-1A or GJ-trop is split (these are labeled as K=9.1, 12.1

140 and 13.1).

141 The group membership for each sample was defined by applying the threshold of 0.65
142 to admixture components. Samples with no admixture components exceeding 0.65 were

143 classified as follows. If the sum of components for subpopulations within the major

144 groups cA (*circum*-Aus), XI (*Xian*-indica), and GJ (*Geng*-japonica) was \ge 0.65, the

145 samples were classified as cA-adm (admixed within cA), XI-adm (within XI) or GJ-adm

146 (within GJ), respectively, and the remaining samples were deemed 'fully' admixed.

147 The newly defined groups were mostly either aligned with the previous K=9 grouping, or

refined those groups, and they were named accordingly (e.g. XI-1B1 and XI-1B2 are newsubgroups within XI-1B).

150 The phenogram shown in Figure 1 was constructed with DARwin v6

151 (http://darwin.cirad.fr/, unweighted Neighbor-joining) using the identity by state (IBS)

152 distance matrix from Plink on the 4.8M Filtered SNP set (available at https://snp-

153 seek.irri.org/_download.zul). Colors were assigned to subpopulations based on K15

154 Admixture results. One entry, MH 63 (XI-adm) represents the admixed types among the

155 XI group.

156 Sample selection, collection and nucleic acid preparation

157 To select accessions to represent the 12 subpopulations of Asian rice that lack high-

- 158 quality reference genome assemblies, the following strategy was employed. The IBS
- 159 distance matrix was used for a principlal component analysis (PCA) analysis in R to

160 generate 5 component axes. Then, for each of the 12 subpopulations, i.e. *circum*-Aus2 =

161 cA2, *circum*-Basmati = cB, *Geng*-japonica (GJ) subtropical (GJ-subtrp), tropical1 (GJ-

162 trop1) and tropical2 (GJ-trop2), and Xian-indica (XI) subpopulations XI-1B1, XI-1B2,

163 XI-2A, XI-2B, XI-3A, XI-3B1, XI-3B2, the centroid of each group in the space spanned

164 by first 5 principal components was determined from the eigenvectors, and the entry

165 closest to the centroid for which seed was available was chosen as the representative for

166 that subpopulation (Table 1).

Single seed decent (SSD) seed from IR 64 and Azucena were obtained from the RiceGenetics and Genomics Laboratory, CIAT, in Cali, Colombia, and SSD seed from the

remaining 10 accessions (Table 1) were obtained from the International Rice Genebank,

- 170 maintained by IRRI, Los Baños, Philippines. All seed were sown in potting soil and
- 171 grown under standard greenhouse conditions at UA, Tucson, USA for 6 weeks at which
- 172 point they were dark treated for 48-hours to reduce starch accumulation. Approximately
- 173 20-50 grams of young leaf tissue was then harvested from each accession and
- 174 immediately flash frozen in liquid nitrogen before being stored at -80°C prior to DNA
- 175 extraction. High molecular weight genomic DNA was isolated using a modified CTAB
- 176 procedure as previously described (Porebski et al. 1997). The quality of each extraction
- 177 was checked by pulsed-field electrophoresis (CHEF) on 1% agarose gels for size and
- 178 restriction enzyme digestibility, and quantified by Qubit fluorometry (Thermo Fisher
- 179 Scientific, Waltham, MA).

180 Library construction and sequencing

181 Genomic DNA from all 12 accessions were sequenced using the PacBio single-molecule 182 real-time (SMRT) platform, and the Illumina platform for genome size estimations and 183 sequence polishing. High molecular weight (HMW) DNA from each accession was gently sheared into large fragments (i.e. 30Kb - 60Kb) using 26-gauge needles and then 184 185 end-repaired according to manufacturer's instructions (Pacific Biosciences). Briefly, 186 using a SMRTbell Express Template Prep Kit, blunt hairpins and sequencing adaptors 187 were ligated to HMW DNA fragments, and DNA sequencing polymerases were bound to 188 the SMRTbell templates. Size selection of large fragments (above 15Kb) was performed 189 using a BluPippin electrophoresis system (Sage Science). The libraries were quantified 190 using a Qubit Fluorometer (Invitrogen, USA) and the insert mode size was determined 191 using an Agilent fragment analyzer system with sizes ranging between 30Kb - 40Kb. The 192 libraries then were sequenced using SMRT Cell 1M chemistry version 3.0 on a PacBio 193 Sequel instrument. The number of long-reads generated per accession ranged from 2.01 194 million (LIMA::IRGC 81487-1) to 5.40 million (Azucena). The distribution of subreads 195 is shown in Figure S2 and the average lengths ranged from 10.58 Kb (Azucena) to 20.61 196 Kb (LIMA::IRGC 81487-1) (Table 2). According to the estimated genome size of the 197 IRGSP RefSeq, the average PacBio sequence coverage for each accession varied from 198 103x (LIMA::IRGC 81487-1) to 149x (IR 64) (Table 2). 199 For Illumina short-read sequencing, HMW DNA from each accession was sheared to 200 between 250-1000bp, followed by library construction targeting 350bp inserts following

standard Illumina protocols (San Diego, CA, USA). Each library was 2 x 150bp paired-

- 202 end sequenced using an Illumina X-ten platform. Low-quality bases and paired reads
- 203 with Illumina adaptor sequences were removed using *Trimmomatic* (Bolger et al. 2014).
- 204 Quality control for each library data set was carried out with *FastQC* (Brown et al. 2017).
- Finally, between 36.52-Gb and 51.05-Gb of clean data from each accession was
- 206 generated and used for genome size estimation (Table S1) by Kmer analysis (Figure S3)
- and the Genome Characteristics Estimation (GCE) program (Liu et al. 2013).

208 Bionano optical genome maps

- 209 Bionano optical maps for each accession were generated as previously described (Ou et
- al. 2019), except that ultra-HMW DNA isolation, from approximately 4g of flash-frozen
- 211 dark-treated (48 hour) leaf tissue per accession, was performed according to a modified
- version of the protocol described by Luo and Wing (Luo and Wing, 2003). Prior to
- 213 labeling, agarose plugs were digested with agarase and the starch and debris removed by
- short rounds of centrifugation at 13,000 X g. DNA samples were further purified and
- 215 concentrated by drop dialysis against TE Buffer. Data processing, optical map assembly,
- 216 hybrid scaffold construction and visualization were performed using the Bionano Solve
- 217 (Version 3.4) and Bionano Access (v12.5.0) software packages
- 218 (https://bionanogenomics.com/).

219 De novo genome assembly

- Genome assembly for each of the 12 genomes followed a five-step procedure as shown in(Figure 2):
- 222 Step 1: PacBio subreads were assembled *de novo* into contigs using three genome
- assembly programs: FALCON (Chin et al. 2016), MECAT2 (Xiao et al. 2017) and
- 224 Canu1.5 (Koren et al. 2017). The number of *de novo* assembled contigs obtained varied
- from 51 (e.g. NATEL BORO::IRGC 34749-1 and KETAN NANGKA::IRGC 19961-2)
- to 1,473 (CHAO MEO::IRGC 80273-1) for the 12 genomes (Table S2).
- 227 Step 2: Genome Puzzle Master (GPM) software (Zhang et al. 2016c) was used to merge
- the *de novo* assembled contigs from the three assemblers, using the high-quality O. sativa
- vg. indica cv. Minghui 63 reference genome MH63RS2 (Zhang et al. 2016a,b) as a guide.
- 230 GPM is a semi-automated pipeline that is used to integrate logical relationship data (*i.e.*
- contigs from three assemblers for each accession) based on a reference guide. Contigs
- were merged in the 'assemblyRun' step, with default parameters (minOverlapSeqToSeq
- 233 was set at 1 Kb and identitySeqToSeq was set at 99%). Redundant overlapping sequences

were also removed for each assembled contig. In addition, we gave contiguous contigs a

higher priority than ones with gaps to be retained in each assembly. After manual

- checking, editing, and redundancy removal, the number of contigs in each assembly
- 237 ranged from 26 (NATEL BORO::IRGC 34749-1) to 588 (LIU XU::IRGC 109232-1)

238 (Table S3).

239 Step 3: The sequence quality of each contig was then improved by "sequence polishing":

240 twice with PacBio long reads and once with Illumina short reads. Briefly, PacBio

- subreads were aligned to GPM edited contigs using the software *blasr* (Chaisson and
- 242 Tesler 2012). All default parameters were used, except minimum align length, which was
- set to 500-bp. Secondly, the tool arrow as implemented in SMRTlink6.0 (Pacific
- 244 Biosciences of California, Inc) was used for polishing the GPM edited contigs. The *bwa*-
- 245 *mem* program (Li 2013) was then used for mapping short Illumina reads onto assembled
- 246 contigs, and the tool *pilon* (Walker et al. 2014) was used for a final polishing step with
- 247 default settings.
- 248 Step 4: The polished contigs for each accession were arranged into pseudomolecules
- using *GPM*, using MH63RS2 (Zhang et al. 2016a,b) as the reference guide. The program
- 250 *blastn* (Altschul et al. 1997) with a minimum alignment length of 1 Kb and an e-value <
- $1e^{-5}$ as the threshold was used to align the corrected contigs to the reference guide. In
- doing so, the corrected contigs were assigned to chromosomes, as well as ordered and
- 253 orientated in the GPM assembly viewer function. The number of contigs after step 4
- ranged from a minimum of 15 contigs (GOBOL SAIL (BALAM)::IRGC 26624-2) to a
- 255 maximum of 104 contigs (IR64) (Table 3). The assembly size for the 12 accessions
- ranged from 376.86 Mb (CHAO MEO::IRGC 80273-1) to 393.74 Mb (KHAO YAI
- 257 GUANG::IRGC 65972-1) (Table 3) and the length of individual chromosome varied
- 258 from 23.06 Mb (chromosome 9 of CHAO MEO::IRGC 80273-1) to 44.96 Mb
- 259 (chromosome 1 of LIMA::IRGC 81487-1) (Table S4). The average N50 value was 23.10
- 260 Mb, with the highest and the lowest values being 30.91 Mb (LIU XU::IRGC 109232-1)
- and 7.35 Mb (IR 64), respectively. The average number of gaps among the 12 new
- 262 genome assemblies was 18, with 8 assemblies containing less than 10 gaps (Table 3).
- 263 Step 5: To independently validate our assemblies, we generated and compared Bionano
- optical maps to each assembly. In total, 17 (Azucena) to 56 (LIU XU::IRGC 109232-1)
- 265 Bionano optical maps were constructed for all 12 rice accessions, which yielded contig
- 266 N50 values of between 22.75 Mb (CHAO MEO::IRGC 80273-1) to 31.45 Mb (KHAO
- 267 YAI GUANG::IRGC 65972-1) (Table S5). As shown in Figure 3 and Figure S4, the

268 chromosomes and/or chromosome arms of all 12 *de novo* assemblies were highly

supported by these ultra-long optical maps. Although rare, a few discrepancies between

- the optical maps and genome assemblies can be seen and are likely due to small errors
- and chimeras that can be produced through both the optical mapping and sequence
- assembly pipelines (Udall and Dawe 2018).
- Following these five steps, we were able to produce 12 near-gap-free *Oryza sativa*
- 274 platinum standard reference genome sequences (PSRefSeqs) that represent 12 of 15
- 275 subpopulations of cultivated Asian rice.

276 **BUSCO evaluation**

- 277 The Benchmarking Universal Single-Copy Orthologs (BUSCO3.0) software package
- 278 (Simao et al. 2015) was employed to evaluate the gene space completeness of the 12
- 279 genome assemblies. These genomes captured, on average, 97.9% of the BUSCO
- reference gene set, with a minimum of 95.7% (IR64) and a maximum of 98.6% (LARHA
- 281 MUGAD::IRGC 52339-1 and KHAO YAI GUANG::IRGC 65972-1) (Table 3).
- 282 Of note, when performing this analysis, we observed that on average 30 out of the 1,440 conserved BUSCO genes tested (https://www.orthodb.org/v9/index.html) were 283 284 missing from each new assembly, 16 of which were not present in all 12, plus the IRGSP, 285 ZS 97, MH 63 and N 22 RefSeqs (Figure S5). This result suggested that these 16 286 "conserved" genes may not exist in rice, or other cereal genomes, thereby artificially 287 reducing the BUSCO gene space scores for our 12 assemblies. To test this hypothesis, we 288 searched for all 16 genes missing in maize, which diverged from rice about 50 million 289 years ago (MYA) (Wolfe et al., 1989, Gale et al., 1998 and Guo et al., 2019). We found 290 that 13 of the 16 genes in question could not be found in 3 high-quality recently 291 published maize genome assemblies (Figure S5) and therefore, concluded that 13 of the 292 16 "conserved" genes in the BUSCO database are not present in cereals, and should be 293 excluded from our gene space analysis. Taking this into account, we recalculated the 294 BUSCO gene space content for each of 12 assemblies and found that 10 of 12 assemblies 295 captured more than 98% of the BUSCO gene set (Table 3).

296 Transposable element (TE) prediction

297 To determine the pan-transposable element content of cultivated Asian rice we analyzed

- the 12 new reference genomes, presented here, along with the MH 63, ZS 97, N 22
- 299 PacBio reference genomes. In addition, we also included a reanalysis of the IRGSP

300 RefSeq as it is conventionally considered the standard rice genome for which all

- 301 comparisons are conducted. This 16 genome data set was used to represent the K=15
- 302 population structure of cultivated Asian rice.
- 303 A search for sequences similar to TEs was carried out using RepeatMasker (Smit
- 304 AFA et al, 2013) run under default parameters with the exception of the options: -
- 305 no_is -nolow. RepeatMasker was run using the library "rice 7.0.0.liban", which is
- an updated in-house version of the publicly available MSU_6.9.5 library (Ou et al. 2019),
- 307 retrieved from
- 308 https://github.com/oushujun/EDTA/blob/master/database/Rice_MSU7.fasta.std6.9.5.out.
- 309 The average TE content of this 16 genome data set was 47.66% with a minimum value of
- 310 46.07% in IRGSP RefSeq and a maximum of 48.27% in KHAO YAI GUANG::IRGC
- 311 65972-1 (Table 4). The major contribution to this fraction was composed of long terminal
- 312 repeat retrotransposons (LTR-RTs, min: 23.55%, max: 27.27% and average: 25.96%)
- followed by DNA-TEs (min:14.87%, max, 16.18% and average: 15.26%). Long
- 314 interspersed nuclear elements (LINEs) and short interspersed nuclear elements (SINEs)
- were identified as on average 1.43% and 0.39% of the 16 genomes, respectively.

316 Structural Variants

Each genome assembly (n=16), as described above, was fragmented using the EMBOSS tool *splitter* (Rice et al. 2000) to create a 10x genome equivalent redundant set of 50kb reads. These reads were then mapped onto every other genome assembly using the tool *NGMLR* (Sedlazeck et al. 2018). Finally, the software *SVIM* (Heller and Vingron 2019) was run under default parameters to parse the mapping output. Only insertions, deletions and tandem duplications up to a maximum length of 25 Kb were considered in this analysis.

324 The results of this analysis identified several thousand insertions and deletions 325 whenever an assembly was compared to any other. Greater variability was found between 326 varieties belonging to different major groups (e.g. Geng-japonica vs. Xian-indica) than 327 occurred between those within these groups. The amount of genome sequences with 328 structural variation between any two varieties ranged from 17.57 Mb to 41.54 Mb for 329 those belonging to the indica (XI, *Xian*-indica) varietal group (avg: 31.75 Mb) and from 330 18.55 Mb to 23.07 Mb (avg: 21.00 Mb) for those in the japonica (GJ, Geng-japonica) 331 varietal group. When all 16 genomes are considered together, the range is between 17.57 332 Mb and 41.54 Mb, with an average value of 33.70 Mb (Table S6). The total unshared

- fraction collected out of all pairwise comparisons was composed for 89.89% by TE
- related sequences.

335 Data Records

- 336 Data for all 12 genome shotgun sequencing projects have been deposited in Genbank
- 337 (https://www.ncbi.nlm.nih.gov/), including PacBio raw data, Illumina raw data, Bionano
- 338 optical maps and the twelve PSRefSeqs. The BioProjects, BioSamples, Genome
- 339 assemblies, Sequence Read Archives (SRA) accession and supplementary files (Bionano
- optical maps) of 12 genomes are listed in Table 3.
- 341 Technical Validation
- 342 DNA sample quality
- 343 DNA quality was checked by pulsed-field gel electrophoresis for size and restriction
- 344 enzyme digestibility. Nucleic acid concentrations were quantified by Qubit fluorometry
- 345 (Thermo Fisher Scientific, Waltham, MA).
- 346
- 347 Illumina libraries
- 348 Illumina libraries were quantified by qPCR using the KAPA Library Quantification Kit
- 349 for Illumina Libraries (KapaBiosystems, Wilmington, MA, USA), and library profiles
- 350 were evaluated with an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara,
- 351 CA, USA).
- 352
- 353 Gene Space Completeness
- 354 Benchmarking Universal Single-Copy Orthologs (BUSCO3.0) was executed using the
- embryophyta_odb9.tar.gz database to assess the gene space of each genome, minus 13
- 356 genes that do not appear to exist in the cereal genomes tested (Figure S5).
- 357

358 Assembly accuracy

- Bionano optical maps were generated and used to validate all 12 genome assemblies.
- 360
- 361 This paper is the first release of 12 PSRefSeqs, optical maps and all associated raw data
- 362 for the accessions listed in Table 3.

363	Code Availability
364	The population re-analysis of 3K-RG dataset and 12 genome assemblies were obtained
365	using several publicly available software packages. To allow researchers to precisely
366	repeat any steps, the settings and the parameters used are provided below:
367	
368	Population structure
369	ADMIXTURE (Alexander et al. 2009) was run with default options. The R scripts for
370	further population structure analysis, including setting up CLUMPP files, can be found in
371	Github repository https://github.com/dchebotarov/Q-aggr.
372	
373	Genome size estimation:
374	The K-mer and GCE program (Liu et al. 2013) were employed for genome size
375	estimation. Command line:
376	kmer_freq_hash -k (13-17) -l genome.list -a 10 -d 10 -t 8 -
377	i 400000000 -o 0 -p genom_kmer(13-17) &> genome_kmer(13-
378	17)_freq.log, and gce -f genom _kmer(13-17).freq.stat -c
379	\$peak -g #amount -m 1 -D 8 -b 1 -H 1 > genome.table 2>
380	genom_kmer(13-17).log
381	
382	Genome assembly:
383	(1) MECAT2: all parameters were set to the defaults. Command line:
384	<pre>mecat.pl config_file.txt, mecat.pl correct config_file.txt</pre>
385	and mecat.pl assemble config_file.txt
386	(2) Canu1.5: all parameters were set to the defaults. Command line:
387	canu -d canu -p canu genomeSize=400m -pacbio-raw
388	rawreads.fasta
389	(3) FALCON: all parameters were set to the defaults. Command line:
390	fc_run.py fc_run.cfg &>fc_run.out
391	(4) GPM: manual edit with merging de novo assemblies from MECAT2, Canu1.5, and
392	FALCON.

393	Polishing:
394	(1) <i>arrow</i> : all parameters were set to the defaults except alignment length = 500
395	bp. The arrow polish was carried out by the SMRT Link v6.0 webpage
396	(https://www.pacb.com/support/software-downloads/).
397	(2) <i>pilon1.18</i> : all parameters were set to the defaults and <i>pilon</i> polish was carried out as
398	recommended at the SMRT Link v6.0 (https://www.pacb.com/support/software-
399	downloads/).
400	
401	BUSCO:
402	The BUSCO3.0 version was employed in this study. Command line: run_BUSCO.py
403	-i genome.fasta -o genome -l embryophyta_odb9 -m genome -c
404	16
405	
406	RepeatMasker:
407	The repeat sequences were employed with the library rice7.0.0_liban in-house. Command
408	line:RepeatMasker -pa 24 -x -no_is -nolow -cutoff 250 -lib
409	rice7.0.0.liban.txt genome.fasta

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- 419 Perpignan, France)
- 420 Author contributions

421 J.Z., K.M., D.C., M.L., N.A., N.R.S.H., H.L., R.M, and R.A.W. designed and conceived

- 422 the research. D.C. and K.M. perform the population structure analysis. K.M., M.L.,
- 423 L.J.A., N.L. generated and provided SSD seed 12 O. sativa accessions. D.K., S.L., S.R.,

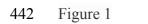
- 424 N.M prepared DNA and performed PacBio and Illumina sequencing. C.S.-S. managed all
- 425 PacBio and Illumina sequence data processing and transfer. P.P. and V.L. generated all
- 426 Bionano optical maps. J.Z. and Y.Z. performed sequence assembly. Y.Z. carried out
- 427 genome size estimation, GPM editing, assembly polishing and data submission. V.L. and
- 428 Y.Z. analyzed the Bionano optical maps and the validation of 12 PSRefSeqs. A.Z. and
- 429 Y.Z. carried out TE prediction and structural analysis. Y.Z., N.A., A.Z., J.Z., D.C., M.L.,
- 430 K.M., N.M. and R.A.W. wrote and edited the paper. All authors read and approved the
- 431 final manuscript.

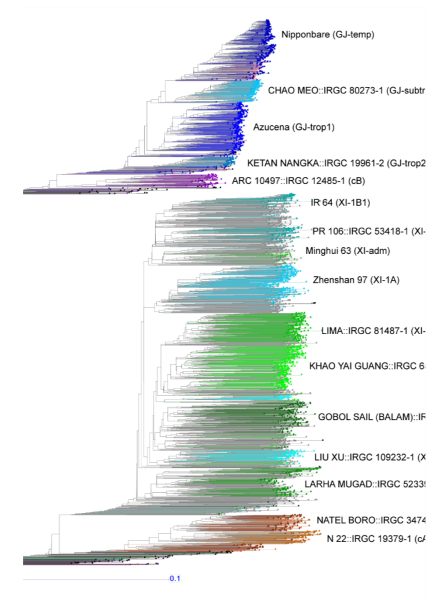
432 Competing interests

- 433 The authors declare that there is no conflict of interest regarding the publication of this
- 434 article.

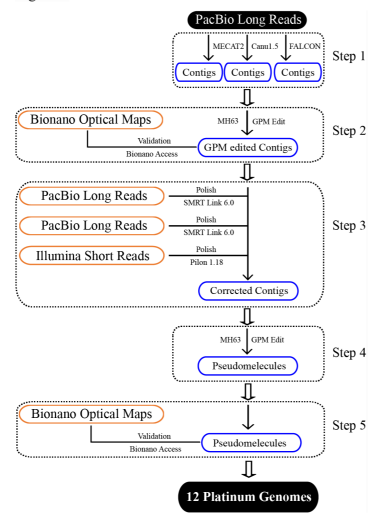
435 Figure legends

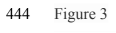
- 436 **Figure 1.** Phylogenetic tree with the accession selected for PSRefSeq sequencing for
- 437 each of the K=15 subpopulations and a single admixture group. Groups are colored
- 438 according to the assignment from Admixture analysis. The subpopulation designation is
- 439 in parentheses following the name.
- 440 **Figure 2.** Genome assembly and validation pipeline.
- 441 **Figure 3.** Bionano optical map validation of chromosome 1 for 12 *de novo* assemblies.

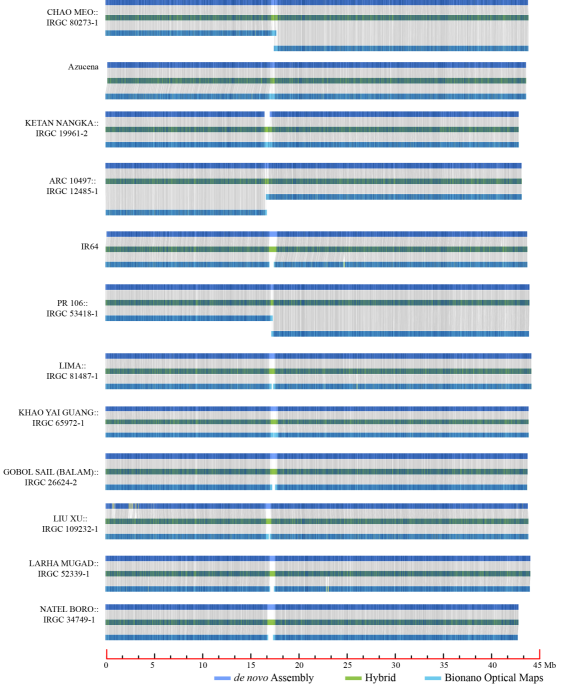




443 Figure 2







445 Tables

- 446 Table 1. Sample collection information for 12 *Oryza sativa* accessions.
- 447 Table 2. Sequencing platforms used and data statistics for the 12 *Oryza sativa* genomes.
- 448 Table 3. *de novo* assembly, BUSCO evaluation and accession numbers in GenBank of the
- 449 12 Oryza sativa genomes.
- 450 Table 4. Abundance of the major TE classes in the 16 *Oryza sativa* genomes.

451 Table 1.

Variety Name	Genetic Stock ID	Country Origin	12 subpopulations	
CHAO MEO::IRGC 80273-1	IRGC 132278	Lao PDR	GJ-subtrp	
Azucena	I1A41685	Philippines	GJ-trop1	
KETAN NANGKA::IRGC 19961-2	IRGC 128077	Indonesia	GJ-trop2	
ARC 10497::IRGC 12485-1	IRGC 117425	India	cB	
IR 64	I1A42114	Philippines	XI-1B1	
PR 106::IRGC 53418-1	IRGC 127742	India	XI-1B2	
LIMA::IRGC 81487-1	IRGC 127564	Indonesia	XI-3A	
KHAO YAI GUANG::IRGC 65972-1	IRGC 127518	Thailand	XI-3B1	
GOBOL SAIL (BALAM)::IRGC 26624-2	IRGC 132424	Bangladesh	XI-2A	
LIU XU::IRGC 109232-1	IRGC 125827	China	XI-3B2	
LARHA MUGAD::IRGC 52339-1	IRGC 125619	India	XI-2B	
NATEL BORO::IRGC 34749-1	IRGC 127652	Bangladesh	cA2	

Subpopulations: GJ = *Geng*-japonica where trop = tropical, subtrp = subtropical; cB = *circum*-Basmati; XI = *Xian*-indica; cA = *circum*-Aus

452 Table 2.

Variety Name	Sequencing platform	Raw data (Gb)	Depth	Number of subreads (M)	Mean subread length (Kb)
CHAO MEO::IRGC 80273-1	PacBio Sequel	49.1	123X	4.26	11.526
Azucena	PacBio Sequel	57.1	143X	5.40	10.581
KETAN NANGKA::IRGC 19961-2	PacBio Sequel	49.8	125X	2.78	17.876
ARC 10497::IRGC 12485-1	PacBio Sequel	44.7	112X	4.06	11.026
IR 64	PacBio Sequel	59.7	149X	5.24	11.393
PR 106::IRGC 53418-1	PacBio Sequel	42.2	105X	2.08	20.317
LIMA::IRGC 81487-1	PacBio Sequel	41.4	103X	2.01	20.612
KHAO YAI GUANG::IRGC 65972-1	PacBio Sequel	42.5	106X	2.37	17.954
GOBOL SAIL (BALAM)::IRGC 26624-2	PacBio Sequel	42.2	105X	2.13	19.777
LIU XU::IRGC 109232-1	PacBio Sequel	55.3	138X	3.66	15.109
LARHA MUGAD::IRGC 52339-1	PacBio Sequel	45.1	113X	3.22	14.011
NATEL BORO::IRGC 34749-1	PacBio Sequel	44.4	111X	2.74	16.2

453 Table 3.

Variety Name	BioProject	BioSample	Genome size (bp)	# Contigs	Contig N50 (Mb)	# Gaps	Scaffold N50 (Mb)	BUSCO	Adjust BUSCO	Genome Accession	SRP	Supplementary Files (Bionano optical map)
CHAO MEO::IRGC 80273-1	PRJNA565484	SAMN12748601	376,856,903	55	11.02	43	30.35	97.60%	98.49%	VYIH00000000	SRP226088	SUPPF_0000003210
Azucena	PRJNA424001	SAMN08217222	379,627,553	28	22.94	16	30.95	97.80%	98.69%	PKQC000000000	SRP227255	SUPPF_0000003212
KETAN NANGKA::IRGC 19961-2	PRJNA564615	SAMN12718029	380,759,091	21	22.68	9	30.70	98.00%	98.89%	VYIC00000000	SRP226080	SUPPF_0000003204
ARC 10497::IRGC 12485-1	PRJNA565479	SAMN12748569	378,463,869	40	17.92	28	30.57	98.40%	99.30%	VYID00000000	SRP226093	SUPPF_0000003206
IR 64	PRJNA509165	SAMN10564385	386,698,898	104	7.35	92	31.22	95.70%	96.57%	RWKJ0000000	SRP227298	SUPPF_0000003213
PR 106::IRGC 53418-1	PRJNA563359	SAMN12672924	391,176,105	16	27.05	4	32.03	96.60%	97.48%	VYIB00000000	SRP226078	SUPPF_0000003202
LIMA::IRGC 81487-1	PRJNA564572	SAMN12715984	392,625,308	17	27.37	5	32.42	98.50%	99.40%	VXJH00000000	SRP226079	SUPPF_0000003203
KHAO YAI GUANG::IRGC 65972-1	PRJNA565481	SAMN12748590	393,737,720	19	21.82	7	32.08	98.60%	99.50%	VYIF00000000	SRP226086	SUPPF_0000003208
GOBOL SAIL (BALAM)::IRGC 26624-2	PRJNA564763	SAMN12721963	391,772,995	15	29.60	3	31.75	97.90%	98.79%	VXJI00000000	SRP226082	SUPPF_0000003205
LIU XU::IRGC 109232-1	PRJNA577228	SAMN13021815	392,033,263	17	30.91	5	32.30	98.40%	99.30%	WGGU0000000	SRP226085	SUPPF_0000003211
LARHA MUGAD::IRGC 52339-1	PRJNA565480	SAMN12748589	390,195,943	16	30.75	4	32.10	98.60%	99.50%	VYIE00000000	SRP226084	SUPPF_0000003207
NATEL BORO::IRGC 34749-1	PRJNA565483	SAMN12748600	383,720,936	16	27.83	4	31.31	98.10%	98.99%	VYIG00000000	SRP226087	SUPPF_0000003209

Variety Name	TOTAL	LTR-RT	LINEs	SINEs	DNA_TEs	Unclassified
NIPPONBARE	46.07	23.55	1.52	0.41	16.18	4.41
CHAO MEO::IRGC 80273-1	46.25	24.00	1.46	0.40	15.59	4.80
Azucena	47.07	24.48	1.47	0.40	15.82	4.89
KETAN NANGKA::IRGC 19961-2	46.99	24.87	1.47	0.40	15.72	4.53
ARC 10497::IRGC 12485-1	46.95	24.74	1.48	0.40	15.68	4.65
IR 64	47.87	26.82	1.42	0.40	14.97	4.26
PR 106::IRGC 53418-1	47.95	26.82	1.41	0.39	15.05	4.28
Minghui 63	47.97	26.61	1.44	0.4	15.3	4.22
Zhenshan 97	47.95	26.79	1.42	0.39	15.19	4.16
LIMA::IRGC 81487-1	48.04	26.87	1.40	0.39	15.01	4.37
KHAO YAI GUANG::IRGC 65972-1	48.27	27.27	1.40	0.39	14.87	4.34
GOBOL SAIL (BALAM)::IRGC 26624-2	48.15	26.99	1.40	0.39	14.99	4.38
LIU XU::IRGC 109232-1	46.92	27.06	1.26	0.32	14.31	3.97
LARHA MUGAD::IRGC 52339-1	48.05	26.74	1.41	0.39	15.09	4.42
NATEL BORO::IRGC 34749-1	47.33	25.75	1.42	0.40	15.12	4.64
N 22::IRGC 19379-1	47.79	25.95	1.44	0.39	15.20	4.81

454 Table 4.

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 539 rice. *Nature genetics* 50.2 (2018): 278.

540 Supplementary Information

- 541 Supplementary file1
- 542 Supplementary Table 1. Summary of Illumina genome survey sequences for 12 Oryza
- 543 *sativa* genomes.
- 544 Supplementary Table 2. Genome features of *de novo* assemblies for 12 *Oryza sativa*
- 545 accessions by Canu1.5, FALCON and MECAT2.
- 546 Supplementary Table 3. Genome features of 12 *Oryza sativa* accessions by GPM
- 547 editing.
- 548 Supplementary Table 4. Chromosome length (Mb) of 12 *Oryza sativa* genomes.
- 549 Supplementary Table 5. Bionano optical map statistics of 12 *Oryza sativa* genomes.
- 550 **Supplementary Table 6**. Summary of large structural variation (>50 bp) by comparison
- of each of 16 genomes to every other genome (including 12 genomes from this study and
- 552 4 previously reported: MH63, ZS97, N 22 and the IRGSP RefSeq).

553 Supplementary file2

- 554 Supplementary Figure 1. Admixture results for K=5 to 15. The samples are grouped
- according to the new classification. At K=9,12,13, the Q matrices converged to two
- 556 different modes, differing according to whether ind1A is split, or tropical japonica.
- 557 Supplementary Figure 2. Length distribution of PacBio long reads used for 12 Oryza
- 558 sativa genome assemblies.
- 559 Supplementary Figure 3. K-mer analysis of Illumina short sequences that were used for
- 560 genome size estimation with the GCE program.
- 561 Supplementary Figure 4. Bionano Access visualization view for 12 de novo assemblies
- 562 with Bionano optical maps and their underlying alignments.
- 563 Supplementary Figure 5. Summary of missing genes in the BUSCO gene space
- 564 evaluation of 12 de novo Oryza sativa assemblies, 4 public Oryza sativa PSRefSeqs and
- 565 3 high-quality Zea mays genomes.