Comparison of *Oryza sativa* and *Oryza brachyantha* Genomes Reveals Selection-Driven Gene Escape from the Centromeric Regions

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Centromeres are dynamic chromosomal regions, and the genetic and epigenetic environment of the centromere is often regarded as oppressive to protein-coding genes. Here, we used comparative genomic and phylogenomic approaches to study the evolution of centromeres and centromere-linked genes in the genus *Oryza*. We report a 12.4-Mb high-quality BAC-based pericentromeric assembly for *Oryza brachyantha* and 10 in *O. brachyantha*. Of these inversions, three resulted in centromere movement (*Chr1, Chr7*, and *Chr9*). Additionally, we identified a potential centromere-repositioning event, in which the ancestral centromere on chromosome 12 in *O. brachyantha* jumped about 400 kb away, possibly mediated by a duplicated transposition event (>28 kb). More strikingly, we observed an excess of syntenic gene loss at and near the centromeric regions (P < 2.2 × 10−16). Most (33/47) of the missing genes moved to other genomic regions; therefore such excess could be explained by the selective loss of the copy in or near centromeric regions after gene duplication. The pattern of gene loss immediately adjacent to centromeric regions suggests centromere chromatin dynamics (e.g., spreading or microrepositioning) may drive such gene loss.

INTRODUCTION

Centromeres in higher eukaryotes are typically composed of tandemly arranged repeats (Henikoff et al., 2001). The flanking pericentromeric regions often exhibit complex DNA structures such as the interspersed satellite DNA tracts and the large segmental duplications described in animal pericentromeres (Bailey and Eichler, 2006) or the Ty3/gypsy retrotransposons that colonize at plant pericentromeres (Neumann et al., 2011). Due to this complex and repetitive nature, the pericentromeric and centromeric regions are vastly underrepresented in whole genome assemblies, thus encumbering the pursuit of in-depth evolutionary and genomic analyses within such regions. Targeted efforts are necessary to confer the completeness and accuracy of the centromeric and pericentromeric assembly toward a reliable genomic analysis as shown in some important model genomes (Nagaki et al., 2004; She et al., 2004; Zhang et al., 2004; Hoskins et al., 2007; Wolfgruber et al., 2016).

Through evolution and speciation, the physical position of a centromere can change; a phenomenon most frequently described in conjunction with (and likely driven by) larger chromosomal rearrangements. However, there are reports of occurrences of novel centromere establishment, known as centromere repositioning, which occurs in the absence of any obvious causal chromosomal rearrangements. However, there are reports of occurrences of novel centromere establishment, known as centromere repositioning, which occurs in the absence of any obvious causal chromosomal rearrangements (Montefalcone et al., 1999). Centromere repositioning was used to explain the emergence of evolutionarily new centromeres (Ventura et al., 2001). Evolutionarily new centromeres are centromeres that form de novo at a new site and have been documented, mostly through cytological observations, in both animals (Rocchi et al., 2012) and plants (Han et al., 2009). However, the mechanism underlying their occurrence is not well understood. Several hypotheses have been proposed, including those driven solely through epigenetic changes (Ferreri et al., 2005; Ventura et al., 2007), while others point to latent centromeres (Ventura et al., 2004) and chromosomal rearrangements (Ventura et al., 2003). Important insights have been recently obtained using ChIP-seq (chromatin immunoprecipitation followed by sequencing) analyses. Centromeres from rice (*Oryza sativa*; Yan et al., 2008), potato (*Solanum*...
**Background:** Centromeres are necessary for faithful chromosome segregation in eukaryotic organisms. The genomic regions conferring centromere function/identity in most plants and animals are composed of large arrays of satellite repeats, surrounded by other highly repetitive DNA sequences. During evolution, centromere locations can change through chromosomal arrangements and some unclear epigenetic mechanisms such as centromere repositioning. Both centromeric chromatin and flanking pericentromeric heterochromatin are thought to be incompatible with gene transcription and thus represent an unfriendly environment for gene survival. However, centromeres may evolve from gene-containing regions and active genes have indeed been found within centromeric regions.

**Question:** How and to what extent have centromere locations changed and what is the evolutionary fate of genes within centromeric and pericentromeric regions? We study these questions in rice (*Oryza* spp) in which abundant high-quality genome sequences are available, facilitating comparative phylogenomic analysis.

**Findings:** We found that the physical positions of the centromeres in *Oryza sativa* and *Oryza brachyantha*, species that diverged approximately 15 million years ago, are generally conserved. Minor alterations in their centromere locations are due to local chromosomal arrangements (e.g., inversions) and/or centromere repositioning, which can be triggered by duplicated transposition. We observed an excess of syntenic gene loss in the centromeric and pericentromeric regions of the rice genome. Most of the lost genes were found to have moved to other genomic regions through segmental duplications, suggesting an evolutionary trend of centromere-linked genes escaping from centromeric regions. The excess gene loss is consistent with selective loss of the parental gene copy in dynamic centromeric region after gene duplications and supports the antagonistic relationship between the centromere environment and gene survival. We also observed abundant new genes at pericentromeric regions.

**Next steps:** Our study showed that selective forces conferred by centromere chromatin dynamics may drive gene relocation out of centromeric regions. It will be interesting to investigate how new genes evolve and fix at pericentromeric regions.

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**IN A NUTSHELL**

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interrogating available Oryza genomes (Stein et al., 2018) as well as closely related grass genomes (as outgroups), we were able to perform a comprehensive syntenic and phylogenomic analyses of the centromeric regions on all rice chromosomes. Our results provide a striking insight into centromere evolution, including conservation of centromere location, potential mechanisms underlying centromere repositioning, and the extent to which natural selection may have contributed to gene exclusion from centromeric regions.

RESULTS

Improvement of the O. brachyantha Pericentromeric/ Centromeric Sequence

In order to improve on the accuracy and completeness of the pericentromeric and centromeric sequence of O. brachyantha and therefore bolster interspecies genomic comparisons, we de novo sequenced 126 BAC clones (Supplemental Data Set 1) from such regions using a BAC-by-BAC strategy with 454 pyrosequencing (see more details in Methods). A total of 12.4 Mb of nonredundant BAC-based assembly for 10 centromeres was generated (Supplemental Table 1). We also confirmed the assembly of the other two centromeric regions by BAC end sequence alignments and manual inspection. In this genome assembly, the pericentromeric regions on all chromosomes except 6 and 10 extend into the CentO-F centromeric satellites (Supplemental Table 1), demonstrating that the sequence and structure of these pericentromeric regions are well covered. Furthermore, we compared our assembly to a separate build that was generated using optical mapping (BioNano Genomics Irys system) (Lam et al., 2012) to validate the contiguity and completeness of our BAC-based assembly. In comparison to the in silico Nt.BspQI recognition site maps, no obvious discrepancies were identified, demonstrating the high quality of the BAC-based assembly (Supplemental Figure 1).

In addition to contiguity, the sequence completeness is also significantly improved compared with the whole-genome shotgun assembly thereby allowing us to generate a high-quality annotation of transposable elements in the O. brachyantha pericentromeric and centromeric regions. Approximately 45.6% and 19.7% of the BAC assemblies are composed of retrotransposons and DNA transposons, respectively (Supplemental Table 2). This is slightly different from the centromeric and pericentromeric regions of O. sativa, in which 60.3% and 8.2% are made up of retroelements and DNA transposons, respectively (Supplemental Table 3). Six LTR-retrotransposon families that have intact elements were identified from the O. brachyantha pericentromeric and centromeric regions. Among them, centromeric retrotransposon FRetro3 (Gao et al., 2009) is the most abundant family having 45 intact elements and 252 solo-LTRs (Supplemental Data Set 2). Previous analysis of the Cen8 BAC sequence from O. brachyantha revealed that the amplification of FRetro3 only occurred within the last few million years (Gao et al., 2009). We found that nearly all FRetro3 elements were inserted into the O. brachyantha genome within the last two million years suggesting a more recent insertion than all other families identified (Figure 1). The overall ratio of solo-LTRs to intact elements in the O. brachyantha pericentromeric and centromeric regions is ~3.5:1, which is significantly higher than previously described in O. sativa (~0.9:1; Ma and Bennetzen, 2006), suggesting a considerable proportion of the O. brachyantha pericentromeric sequence has been removed due to unequal recombination events.

Comparison of local size variations in the centromeric regions between O. sativa and O. brachyantha revealed that nearly all centromeric regions in O. sativa have been expanded relative to the orthologous centromeric regions of O. brachyantha, and sizes increased from 0.05 in CEN3 to 1.07-fold in CEN4 (Supplemental Table 4). The gene content in the pericentromeric regions is generally low; for genes conserved between O. sativa and O. brachyantha, the density is ~2.6 per 100 kb in O. brachyantha, compared with ~1.86 in O. sativa (Supplemental Table 4).

ChiP-Seq with CenH3 in O. brachyantha

To determine which sequences/regions harbor the functional centromeres in O. brachyantha, we performed ChiP-seq analysis using antibodies targeting centromeric histones (CenH3).
We obtained a total of 295,118 high-quality 454 reads with an average length of 250 bp. The mapping strategy applied has been previously described in *O. sativa* (Yan et al., 2008). To avoid mapping bias due to an incomplete CentO-F assembly, we first filtered the reads containing CentO-F sequence. The subsequence mapping results revealed no obvious continuous genomic regions that were enriched for ChIP-seq reads (Supplemental Figure 2), indicating the functional centromeric regions in *O. brachyantha* were embedded exclusively in highly repetitive sequences, rather than in unique or low copy sequences (Yan et al., 2008; Wade et al., 2009). Surprisingly, we found some genomic regions, including a region from the short arm of chromosome 9 and the pericentromeric regions of chromosomes 6 and 11, which were significantly enriched in our read counts. Sequence analysis of these regions revealed they are 45S rRNA genes, suggesting that 45S rRNA clusters may contribute to the centromeric landscape in *O. brachyantha*. To test this hypothesis, we performed a classification analysis of the 454 reads in ChIP-seq data set and a control genomic data set also generated using 454 sequencing. The ChIP-seq data set contained 112,177 (~38%) and 11,684 (~4%) reads from CentO-F (a more detailed analysis of CentO-F is provided in Supplemental Text 1) and FRetro3, respectively, based on the RepeatMasker annotation (Supplemental Table 5). In contrast, we only observed ~1.6% and ~2.5% in the genomic data set, indicating Cento-F and Retro3 are indeed centromere-associated as shown in previous findings (Lee et al., 2005). As expected, we found that 10,863 reads (~3.7%) were derived from 45S rRNA genes in the ChIP-seq data set (Supplemental Table 5) compared with only 1.3% in the genomic data set, suggesting some functional centromeric regions may contain 45S rRNA genes in *O. brachyantha*. We found an ~600-kb 45S rRNA gene cluster immediately adjacent to the centromeric region of chromosome 9 (Supplemental Figure 3); thus, this cluster most likely contributed to the enrichment of 45S rRNA gene sequence in the ChIP-seq data set.

### Table 1. The Major Pericentric Inversions between Rice and O. brachyantha

<table>
<thead>
<tr>
<th>Name</th>
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<th>Specificb</th>
<th>Synteny Maps</th>
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</thead>
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<td>OB</td>
<td>Supplemental Figure 6A</td>
</tr>
<tr>
<td>PeriCen2</td>
<td>~1727</td>
<td>OB</td>
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<td>OS</td>
<td>Supplemental Figure 8</td>
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<tr>
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<td>~630</td>
<td>OS</td>
<td>Supplemental Figure 9</td>
</tr>
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<td>OB</td>
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<td>OB</td>
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</table>

aThe size of each pericentric inversion is determined by the length of the inverted segment in the species in which the inversion occurred.
bOB, *O. brachyantha*; OS, *O. sativa*.

### Centromere Synteny

We generated a comparative sequence map between *O. sativa* and *O. brachyantha* to infer centromere synteny. We used two genome assemblies for *O. sativa*: the *japonica* rice Nipponbare (MSU7) (Kawahara et al., 2013) and the *indica* rice Shuhui 498 (R498) (Du et al., 2017). Nipponbare was assembled using a traditional map-based, clone-by-clone approach, and R498 was assembled with the integration of single-molecule sequencing and mapping data set, a genetic map, and fosmid sequence tags. Approximately 2 Mb of homologous region flanking each centromere (Supplemental Table 6) was selected from a whole-genome alignment between Nipponbare and *O. brachyantha*, including ~5000 syntenic markers (Supplemental Table 7). The synteny map revealed extensive structural rearrangements in centromeric and pericentromeric regions between these two *Oryza* genomes (Figure 2; Supplemental Figure 4). Next, R498 was aligned to Nipponbare to ascertain whether significant discrepancies exist in the pericentromeric/centromeric regions between the genome assemblies of these two subspecies, which indeed verified the structural rearrangements identified between Nipponbare and *O. brachyantha*. Shifting of the centromere positions on chromosomes 1, 7, and 9 can each be resolved through a single inversion event. The differences in centromere position on chromosomes 3, 4, 5, 6, and 8 however appear to have arisen from a series of complex local arrangements. The centromere position is conserved on chromosomes 2, 10, and 11 between these two species. We found that only the centromere on chromosome 12 appears to have repositioned considerably, which likely resulted from a centromere repositioning event.

To infer the species of origin in which the inversions occurred, we performed an additional gene-based synteny analysis that included at least outgroup species from *Leersia perrieri* (Stein et al., 2018), a sister genus to *Oryza, Brachypodium distachyon, Setaria italica* (foxtail millet), and *Sorghum bicolor* (Supplemental Table 7).
Selection Drives Gene Escape from Centromeres

Figure 5). The homologous regions corresponding to the reference rice genome for the distantly grass species are provided in Supplemental Data Set 3. In total, we were able to trace the origin of 17 pericentromeric inversions, with sizes ranging from ∼50 to ∼1700 kb (Table 1, Figure 3; Supplemental Figures 6 to 13), 10 of which are specific to the *O. brachyantha* lineage and seven to *O. sativa*. A pericentric inversion specific to *O. sativa* has previously been reported to cause the physical shift of Cen8 (Ma et al., 2007), which is inconsistent with our result. We found both *L. perrieri* and *B. distachyon* share a relatively similar gene order to *O. sativa*; therefore, the inversion is presumably specific to *O. brachyantha* (Supplemental Figure 6) and appears to have no direct effect on the centromere position. We caution that synteny analysis within complex genomic regions such as centromeres can be ambiguous if pursued with only a few available syntenic markers.

Our syntenic data demonstrates that the centromere movement involving chromosome 1 in *O. brachyantha* (Supplemental Figure 6) and chromosome 7 in *O. sativa* (Figure 3A) are unique to their respective species and each was caused by a single pericentric inversion. Our data also reveal that the centromere movement on chromosome 9 is specific to *O. brachyantha* (Figure 3B). We detected a hemicentric inversion (one break in the active centromere and a second break on the chromosome arm) (Lamb et al., 2007) within the *O. brachyantha* Cen9 region. However, the new centromere is not located exactly at the distal breakpoint region; rather, it moved across a genomic segment of ∼120 kb, distally. Therefore, the centromere movement on *O. brachyantha* chromosome 9 might have involved a micro-repositioning event, in addition to the inversion. We also observed hemicentric inversions in the Cen5 and Cen10 regions of *O. brachyantha* (Supplemental Figures 10 and 12).

We were unable to resolve the discrepant centromere synteny observed in Chr3, Chr5, and Chr6 due to extreme syntenic decay within these centromeric regions. Although extensive chromosomal rearrangements have occurred within the centromeric regions between *O. brachyantha* and *O. sativa*, the centromeric synteny (approximate centromere position) is preserved with the exception of a few “subtle” changes (<500 kb). Our results suggest that the shift in centromere position between *O. sativa* and
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O. brachyantha can be explained by small local inversions and repositioning.

Evolution of Cen12

Centromere movement on chromosome 12 can either be explained via two successive inversions or as the result of a repositioning event. To elucidate the evolutionary origin of this event, we performed a gene-based synteny analysis of Cen12 regions, using L. perrieri as the outgroup (Figure 4A). We verified the sequence contigs from L. perrieri using BAC-end sequence alignments (only uniquely mapped sequences were used) (Supplemental Figure 14). The ancestral centromere position of chromosome 12 can be determined by the outgroup species, L. perrieri, in which the centromere position was shown to be syntenic with O. sativa. This is supported by the fact that the corresponding region of ancestral centromere position in O. brachyantha still retains a small cluster of residual centromeric satellite repeats (Figure 4A). Therefore, we conclude that the centromere of O. brachyantha has likely changed from its ancestral position. The new centromere location separated by ∼400 kb from the ancestral position. An inversion involving only one gene might have caused the centromere movement between the centromere of O. sativa and the inactive centromere of O. brachyantha (Figure 4A). However, due to the lack of syntenic information from L. perrieri, we were unable to assign this inversion and its associated centromere movement to a specific lineage.

To investigate whether specific genomic features are associated with this centromere movement, we performed a detailed sequence analysis of the ∼1.2 Mb Cen12 region in O. brachyantha. Interestingly, we identified an inverted segmental duplication of ∼28 kb in the Cen12 region (Figure 4A). The duplicated segments were located precisely at the ancestral and new centromeric positions. The paralogous segments showed a high degree (∼94.5%) of sequence similarity (Figure 4D), suggesting that the duplication occurred recently. The calculation of nucleotide substitutions per site between these two paralogous sequences allow us to estimate an approximate divergent time of ∼1.54 Ma, using the formula: $T = \frac{K}{2r}$, where $K$ is the average number of substitutions per aligned site and $r$ is the average substitution rate of $1.3 \times 10^{-8}$ (Ma and Bennetzen, 2004). It should be
Figure 4. Evolutionarily New Centromere on O. brachyantha Chromosome 12 Probably Resulted from a Duplicated Transposition.

(A) Gene synteny map revealed centromere on O. brachyantha chromosome 12 has repositioned. Double slashes indicate the positions of centromeric gaps, representing the potential centromere positions in each species. Orthologous genes (black rectangles) are connected by dashed lines. The predicted ancestral centromere retains a small cluster of residual centromeric satellite repeats (shaded in green). The dashed line in green indicates an inversion containing only one gene has occurred between O. sativa and O. brachyantha.

(B) Annotation map of the 1.2-Mb region of O. brachyantha Cen12. Gene models, transposable elements (see detailed figure legend in the black box below), CentO-F clusters (red squares, with directions indicated by green arrows above), and a segmental duplication event (gray polygon) are depicted in the map. The size of duplicated segment is approximate 28 kb. The paralogous segments are exactly adjacent to the CentO-F sequence in the ancestral centromere position. Two genes in the dashed orange box were inserted into the ancestral centromeric position in O. brachyantha.
noted that since genomic regions near centromeres always have a higher mutation rate (Ossowski et al., 2010), the divergent time may still be overestimated. This duplicated segment is immediately adjacent to the CentO-F cluster, indicating at least a portion, if not all, of the ancestral CentO-F sequences are involved in this segmental duplication, consistent with the inverted orientation of CentO-F clusters between the new and old centromeric loci. Since segmental duplications have been proposed to play a role in the new centromere seeding (Cardone et al., 2007) and local segmental duplications facilitate centromere expansion in rice (Ma and Jackson, 2006), we suggest that Cent12 movement in O. brachyantha was more likely as the result of centromere repositioning triggered by segmental duplications.

Another notable feature of the O. brachyantha Cent12 region is that the genomic region (~66 kb) immediately adjacent to CentO-F cluster of the new centromeric region is composed almost entirely of nested LTR-retrotransposons (Figure 4C). Therefore, LTR-retrotransposon accumulation may have contributed to the new centromere expansion.

Sequence analysis of the duplicated segments revealed a mosaic composition of genomic segments that originated from three chromosomes (Figure 4D; Supplemental Table 8). The structural and evolutionary patterns observed in pericentromeric segmental duplications is similar to those reported in humans (Horvath et al., 2000), suggesting an example of convergent evolution among pericentromeric regions between distant eukaryotic lineages.

An Excess of Syntenic Gene Losses near the O. sativa Centromeres

Gene loss is one of the major reasons for synteny decay in plant genomes. To characterize the rate and distribution of syntenic gene loss in O. sativa, we attempted to identify single or low-copy genes (identified as gene copies equal to or less than 3 based on OrthoMCL analysis) that are present in O. brachyantha and have syntenic orthologs in at least one outgroup species (L. perrieri and/or B. distachyon), yet are absent in the syntenic regions of O. sativa. Using intensive manual inspection, we identified 206 genes that had been lost from their original position in the O. sativa genome after its divergence from O. brachyantha, which accounted for ~1% of the total syntenic gene pairs (206/20,467). We further assigned each gene loss event to the Oryza phylogenetic tree by including four additional Oryza genomes, i.e., O. glaberrima, O. glumeaapatula, O. meridionalis, and O. punctata, which diverged from O. sativa 0.8 to 6.8 Mya (Stein et al., 2018) (Figure 5A). We found that the relative syntenic gene loss along the branches leading to O. sativa are proportional to the branch lengths (Figure 5B) with an exception of the branch between nodes II and III, suggesting that gene loss occurred at a fairly constant rate during Oryza genome evolution.

We analyzed the distribution of these 206 genes along the 12 chromosomes in O. sativa. We divided each chromosome arm into nine bins from the telomere to the centromere, each bin with an identical gene number (Figure 5C). The gene loss events are enriched at the bins closest to centromeres and telomeres, a phenomenon that also been described in Drosophila melanogaster (Han and Hahn, 2012). Approximately a quarter of the gene loss events (47/206) fell into the regions immediately flanking each centromere with 10 syntenic genes on each side (Supplemental Figure 15 and Supplemental Data Set 4), which are generally within previously defined crossing-over suppressed pericentromeric regions (Yan et al., 2008; Tian et al., 2009). Therefore, centromeric or pericentromeric regions have experienced a significant excess of syntenic gene loss compared with other genomic regions. This is especially apparent in the centromeric regions of chromosome 6 and 9 other chromosomes with the exception of chromosomes 10 and 12 (Binomial test, P < 2.2 × 10−16; Table 2).

Most Genes Absent from Syntenic Regions near Centromeres Have Duplicated to Nonsyntenic Positions

Since the genes that were identified as missing at the pericentromeric and centromeric regions are of single or low-copy number and also evolutionarily conserved between distantly related grass species, they should be essential for Oryza genomes. Indeed, the functional annotation of these genes revealed that some of them encode important proteins, such as FAD binding and arabinol-lactone oxidase domains, AMP binding domain, dolichol phosphate-mannose biosynthesis regulatory, AMP binding domain containing, glycosyltransferase, and so on (Supplemental Data Set 4). Thus, the loss of such proteins at the genomic level seems unlikely since it would otherwise affect gene dosage balance and/or result in lethality. Two circumstances provide reasonable explanations for the missing syntenic genes: These genes existed in two or more copies in the ancestral species when it diverged into O. sativa and O. brachyantha or, alternatively, the genes have simply moved to other genomic positions after the divergence of O. sativa and O. brachyantha. As we expect, among the 47 missing genes in syntenic positions near centromeres in O. sativa, 41 have homologs elsewhere in the genome (Supplemental Data Set 4). To investigate which circumstance lead to the gene loss, we sought to...
examine whether those nonsyntenic homologs (41 cases) arrived before or after the divergence of *O. sativa* and *O. brachyantha*. To achieve this goal, we performed synteny analyses that included five outgroup species, i.e., *L. perrieri*, *B. distachyon*, foxtail millet, sorghum, and maize. If a gene was present in a syntenic region with any of the outgroup genomes, it was regarded as an ancestral gene, whereas if it was absent in the syntenic regions from all outgroup species, it would be regarded as a derived gene (Supplemental Figure 16). Using these criteria, we are able to determine that at least 33 genes were derived loci in *O. sativa*, suggesting that genes moved here from centromeric regions after the split of the *O. sativa* and *O. brachyantha* lineages. Of these 33 genes, 32 are subject to purifying selection as shown that the nonsynonymous to synonymous nucleotide substitution rates are significantly less than one by comparing them with homologous genes in the centromeric regions in *O. brachyantha*.

**Figure 5.** Syntenic Gene Loss/Movements during Rice Genome Evolution.

(A) Placement of syntenic gene loss events along the branches leading to *O. sativa* in the *Oryza* species tree. Nodes (red circles) are labeled with Roman numerals. The length of branches (units, million years) is drawn to scale. The number of syntenic gene loss events is shown at each node. The phylogenetic inference and species divergence times are resolved and described in a recent report (Stein et al., 2018).

(B) Correlation of gene loss event and divergence time. The number of gene loss events was plotted against the divergence time of each node (see [A]). A strong correlation is found ($R^2 = 0.995$), suggesting gene loss events occurred at a constant rate during the *Oryza* genome evolution.

(C) Syntenic gene loss events are enriched at centromeric or telomeric regions. Each chromosomal arm was divided into 9 bins, each with the same gene number. Gene loss events were calculated for each bin. An apparent enrichment of gene loss events in the bin closest to the centromere was observed.

(D) An example of pseudogenization event in the *O. sativa* Cen8 region. The purple inverted triangle indicates an insertion event, while the red line indicates a reading-frame shift mutation (2-bp deletion).

(E) LOC_OS08g41460 (MSU7 Chr8: 26,174,014–26,185,145) is highly homologous to Ob08g19210 and the pseudogene sequence in Cen8 region. Analysis of this locus reveals it was derived before the insertion event occurred between the divergence of *O. sativa* and *O. meridionalis* but after its divergence with *O. punctata*, implying this gene was a duplicated copy of the donor sequence. The donor gene at Cen8 became a pseudogene.

(F) Breakpoint analysis of the genomic segment containing locus LOC_OS08g41460 suggests the insertion was associated with the repair of double-strand breaks through nonhomologous end joining. A target site duplication of 6 bp was detected exactly at the insertion point.
(Supplemental Table 9). In addition, 31 genes show expression in seedlings (Zhang et al., 2012; Supplemental Table 9), 27 of which have an FPKM (fragments per kilobase of transcript per million mapped reads) value larger than one. These results indicate the set of relocated genes described retain function after settling into their new locus. Notably, we only found two duplication events that involved more than one gene, each with three and two genes, respectively, indicating that most of gene movement occurred independently (Supplemental Data Set 4).

Most of the missing genes have moved to other genomic regions, thus mitigating the effect of gene dosage imbalance; however, mechanistic action of these movements is poorly understood. In our previous study, we reported that the breakpoint sequence analysis suggested nonhomologous end joining through the repair of double-strand breaks is the major mechanism account for creating the non-collinear genes in the rice genome (Chen et al., 2013). In this study, most of the derived genes (32/33) moved from centromeric regions prior to the divergence of the AA genome species (Supplemental Data Set 4), accounting for ~2.5 million years of divergence. During such an evolutionary time period, sequence features necessary for a mechanistic characterization are sparse, especially within the rapidly evolved centromeric regions, resulting in only six cases that we were able identify degenerated sequences of parental genes in the ancestral loci and propose a mechanism. In the cases within the rice Cent8 region, two genes (homologs of Os08g1460, located on the long arm of chromosome 8, which may be the derived copy (acceptor). The derived copy appears to have originated from an insertion event (Figure 5E) by the double-strand break repair through nonhomologous ending joining (Figure 5F).

If gene duplication is indeed the major mechanism responsible for genes moving out of centromeric regions, we expect there should be considerable and recent gene duplications in centromeric gene regions. An example of this can be observed in the rice Cent4 region, in which a segment from the core region of this centromere was duplicated to the long arm of chromosome 4. This event was demonstrated to have occurred only after the divergence of O. sativa ssp japonica and O. sativa ssp indica (Supplemental Figure 17), suggesting it is a very recent event. The segment contains four annotated genes, including Os04g17650, Os04g17660, Os04g17680, and Os04g17700. In the derived loci, three genes were retained (Os04g24410, Os04g24430, and one unannotated gene model). Thus, gene movements in centromeric regions are still ongoing.

### Gene Gain at the Genomic Regions near the O. sativa Centromeres

To investigate the extent of gene gain near centromeres in O. sativa, we focused on 240 genes, which are the most centromere-proximal (10 on each side of each centromere) and with high confidence, i.e., supported by full-length cDNAs, expression evidence (Supplemental Data Set 5), or evolutionary conservation. Of these 240 genes, 146 are shared with O. brachyantha and 14 were lost from the syntenic positions in O. brachyantha, leaving the remaining 80 candidates of gene gain. Since most of them (64/80) have no homologs in O. brachyantha or any other outgroup species, most likely, they have originated after the divergence of O. sativa and O. brachyantha. By introducing a number of intermediate species (Figure 5A), we found 59/64 are specific to the AA lineage. These results are consistent with the recent report (Stein et al., 2018) that the AA-type genomes experienced a higher rate of new gene emergence and new genes are more likely to be found near centromeres over low-recombination regions.

Despite of a striking number of gene gains near the O. sativa centromeres, most are likely new genes and not present before the divergence of O. sativa and O. brachyantha. For those genes that existed before the divergence of O. sativa and O. brachyantha, only 10 can be unambiguously traced from other genomic regions of origin. Notably, during the evolutionary period from the O. sativa-O. brachyantha split to the O. sativa-O. punctata split, only four genes moved in, compared with 24 genes moved out in the same evolutionary duration. This result indicates that gene translocations to O. sativa centromeres are less frequently than moving out, at least during the evolutionary period before it divergent from O. punctata.

### Reciprocal Syntenic Gene Loss between O. sativa and O. brachyantha Centromeric Regions Provides Evidence for Antagonism between Gene and Centromere

We identified 45 syntenic gene losses (Supplemental Data Set 6) in the pericentromeric and centromeric regions (genome-wide
DISCUSSION

Our study presents a sequence-level comparative genomic analysis of pericentromeric/centromeric regions in a relatively complex eukaryotic genome. The high-quality data generated here bolster our ability to accurately identify structural variations and hence permit reliably tracking the evolutionary history within complex pericentromeric and centromeric regions, which in turn allow us to relate each centromere movement to specific chromosomal rearrangements. We also showed that gene loss near centromeres may have accompanied the evolution of centromeres.

Conservation of Centromere Location

Although O. sativa and O. brachyantha separated each other for more than 15 million years and left <10% sequence conserved at the flanking regions of centromeres, our comparative analysis revealed that centromere syntenry between these two Oryza genomes is still well maintained with the exception of a few subtle changes. This conservation can even extend to more distantly related grass species, such as B. distachyon, sorghum, maize, and wheat (Triticum aestivum; Qi et al., 2009, 2010; Wang and Bennetzen, 2012). Therefore, from a genomic perspective, the location of the plant centromeres as observed here is unlikely to change substantially, supporting the view that chromosomal location may be an evolutionarily conserved primary determinant for functional centromere identity (Thakur and Sanyal, 2013). This viewpoint could be further supported by recent observations that centromere positions between Z. mays and some distantly related species, e.g., Zea diploperennis, Zea luxurians, and Tripsacum dactyloides are stable within a constraint genomic region although extensive variations existed in CentC content (Gent et al., 2017).

Genomic Variations and Centromere Location

We documented many pericentric inversions in O. sativa and O. brachyantha, and some of the events that gave rise to centromere movements can be explained via evolutionary analysis. It should be noted that the overall inversion number can’t be accurately measured due to the remarkable divergence of sequences and therefore we anticipate the exact number to be larger than what is reported here. Several hemi-centric inversions were found in centromeric regions, suggesting such a phenomenon is common in plant centromeric regions (Lamb et al., 2007; Schneider et al., 2016; Wolfgruber et al., 2016). The frequency of detected hemi-centric inversions is indeed puzzling given the propensity of these types of inversions to disrupt centromere integrity.

Centromeric chromatin (CenH3-containing nucleosomes) can spread over the flanking noncentromeric regions (Lam et al., 2006; Müller and Almouzni, 2017). In contrast to the dynamic genomic architecture of centromeres, the physical size of a centromere may be a critical element in maintaining a robust epigenetic inheritance (Bodor et al., 2014). For example, in grass species, centromere sizes are positively correlated with genome sizes (Zhang and Dawe, 2012) and are found to be uniform within a given species (Wang et al., 2014b). Centromere expansion may be the key factor in determining whether a genomic variation is tolerated within a centromeric region. Hemi-centric inversions can split centromeric chromatin and this centromeric chromatin can then spread into the flanking regions of the breakpoint in order to restore the optimal size of the centromere. The abundance of repetitive sequences around centromeres might be beneficial, as they can provide a mechanism to buffer genomic sequence from variation and exchanges within centromeric regions, thereby stabilizing the region (Fukagawa and Earnshaw, 2014). While some regional centromeres may cover hundreds of kilobases to several megabases along the chromosome (Wolfgruber et al., 2009), any degree of genomic variation within centromeres, both small (e.g., single-nucleotide polymorphisms and indels) and large (e.g., copy number variations and inversions), may have the potential to reshape centromeric chromatin.

Genes Escaping from Genomic Regions near/within Functional Centromeres during Evolution

Genomic positional bias in gene loss patterns have been observed in several evolutionary events, such as during the diploidization that follows whole-genome duplication (Langham et al., 2004; Thomas et al., 2006) and during the evolution of sexual chromosomes from autosomes (Cortez et al., 2014; Bergero et al., 2015). In this study, we observed an excess of syntenic gene loss at the centromeric regions of Oryza genomes, and this bias is generally resulted from selective loss of parental gene copies in the centromeric regions after gene duplications. Two major explanations are proposed to address this pattern, i.e., why duplicated genes in centromeric and pericentromeric regions are more likely to be pseudogenized or lost. First, meiotic
recombination is known to be severely repressed or even absent near centromeres; thus, natural selection against deleterious gene mutations would be reduced, a scenario similar to the degeneration of Y chromosome (Charlesworth and Charlesworth, 2000). Our results reveal that most of the relocated genes currently reside within recombination-repressed regions. In addition, deleterious variants are indeed found enriched in regions of low recombination in the rice genome (Liu et al., 2017). Second and more likely, our result revealed genes immediately adjacent to functional centromeric regions are more likely to be lost, which can be explained by the antagonistic relationship between centromeric chromatin (CenH3) and active genes (Wolfrubger et al., 2016). The encapsulation of CenH3 across an active gene may result in the silencing of that gene and is thereby deleterious, especially when the gene is essential. Therefore, genes located close to the centromere may act as a barrier to prevent the spread of centromeric chromatin (CenH3) (Lomiento et al., 2008; Alonso et al., 2010), analogous to the boundary elements for maintaining a local balance of heterochromatin and euchromatin (Wang et al., 2014a). However, if the gene has paralogs (e.g., as a result of gene duplication) in other genomic regions, the barrier function will become weakened or completely lost because redundant genes can compensate the function of this gene. The spread of CenH3 encompassing this gene will have greatly reduced effect on genome function. Once this gene is silenced by CenH3 encapsulation, it would likely accumulate mutations due to the relaxation of functional constraints and eventually become pseudogenized or lost. In addition, gene loss at centromeric regions may be beneficial for centromere establishment and maintenance. Genomic variations are frequently found within centromeric regions; therefore, the expansion of CenH3 may provide a proactive mechanism to maintain optimal centromere size and organization. In this regard, selective loss of a duplicated gene within centromeric regions could provide a sequence resource for use during CenH3 expansion. This would be especially crucial to new centromeres because they are relatively small and successful expansion is thought to play a key role on their final survival and maturity.

Selective forces contributing to biased gene movement have been documented in animal sex chromosome evolution. For example, genes moving out of the mammalian X chromosome are thought to be driven by natural selection to attain male germline function (Emerson et al., 2004). Such forces have also been reported in the X chromosome in Drosophila (Betran et al., 2002; Vibranovski et al., 2009) and the Z chromosome in silkworm (Wang et al., 2012). In our study, we reveal a potential novel selective force where CenH3 chromatin dynamic spreading or repositioning can drive gene movement or gene relocation in the *Oryza* genomes.

The observation of abundant new genes at pericentromeric regions appears paradoxical to our conclusion. Several assumptions may help to explain this paradox. First, although new genes can generate at a high rate within the pericentromeric regions as driving by the rapid evolutionary turnover in such regions (She et al., 2004), most would be rapidly lost and a small set will be retained by selection (Carvunis et al., 2012; Neme and Tautz, 2013). New genes have relatively shorter exons and lower expressions, even with relaxed selective constraints comparing to old genes (Stein et al., 2018), suggesting their “dispensable” role to the genome. Our result that most new genes are specific to AA lineage (∼2.5 million years), few were found to be emerged within the duration (∼8.2 million years) between the *O. sativa*- *O. punctata* split and the *O. sativa*- *O. brachyantha* split, supports the assertion that only a small set of new genes can survive over time if the rate of new gene emergence is constant during evolution. Second, centromere maturity level can affect the rate of new gene creation and/or survival. While centromeres have not been completely evolved to the repeat-based centromeres, their immediate pericentromeric regions are more resistant to accept new genes. However, once centromeres are gradually becoming mature by acquiring sufficient satellite repeats within their core, the expanded flanking pericentromeric regions would become less sensitive to new gene creation because the new created genes are already far away from the centromeric chromatin. Therefore, the distinct extend of new genes observed before and after *O. sativa* diverged from *O. punctata* may reflect this effect.

**METHODS**

**BAC Selection, Sequencing, Assembling, and Verification**

The BAC library, BAC-end sequences, and fingerprinted contig data of *Oryza brachyantha* were obtained from the Arizona Genomics Institute (http://www.omap.org/resources.html) (Ammiraju et al., 2006; Kim et al., 2008). BAC clones selected for sequencing were identified by alignments of BAC-end sequences onto pseudomolecules representing the pericentromeric regions of *O. brachyantha*. The pseudomolecules were constructed by manually ordering and orienting the raw scaffolds that were used to generated the whole genome assembly (Chen et al., 2013). Tiling-path BAC physical maps were also constructed for each centromeric region (covering at least 2 Mb on each side).

We sequenced a total of 126 BAC clones using the Roche 454 platform. These BAC clones represent 21 contigs, including 18 pericentromeric regions, two regions from chromosome 2 that were sequenced to verify the inversion breakpoints, and one region from the telomere on the short arm of chromosome 9 (Supplemental Data Set 1). In addition, we sequenced clone 45C05 to extend the centromeric region of chromosome 8. Several clones for the *O. brachyantha* centromere 8 have already been sequenced using the standard Sanger methods (Gao et al., 2009). We sequenced 56 clones using a pooled BAC clone strategy (each pool contained two to three clones), and independently sequenced 71 clones due to their repetitive nature (Supplemental Data Set 1).

All BAC sequences were assembled with Newbler (version 2.6) using the default parameters, followed by intensive manual correction. We determined overlaps between neighboring clones using BLASTN and constructed the final pseudomolecules with careful inspection and verification of each overlap.

We compared our BAC-based assemblies to an optical physical genome map to validate accuracy and completeness. We produced the genome map using the BioNano Genomics Irys system (Lam et al., 2012). In total, we collected 39 GB of data (>100 kb) representing ~130x genome coverage with a N50 length of 154 kb (Supplemental Table 10). The assembly has 244 maps with a N50 of 1.33 Mb that spans 255 Mb.

**ChIP-Seq Analysis**

ChIP experiments performed as previously described (Yan et al., 2008). ChIP DNA was subjected to 454 sequencing. Reads contain CentO-F...
sequences were first filtered using BLASTN and RepeatMasker (www.
repeatmasker.org). The rest reads were mapped onto the improved O. brachyantha genome assembly using LASTZ (Schwartz et al., 2003) with coverage of 90% and identity of 95% (Supplemental Methods). We plotted the number of CenH3 ChIP-seq reads along individual chromosomes in widow size of 20 kb, spaced every 10 kb.

Identification of LTR-Retrotransposon and Timing of Its Integration

LTR-retrotransposons were identified using LTR_STRUC (McCarthy and McDonald, 2003) with default settings and RepeatMasker with a custom TE library for O. brachyantha (Supplemental Methods). The integration times of full LTR-retrotransposon elements were estimated as previously described (Yang et al., 2012).

Sequence-Based Synteny Map

Genome sequences of Oryza sativa ssp japonica (IRGSP1.0), O. sativa ssp indica R498, and O. brachyantha were soft masked using RepeatMasker (www.repeatmasker.org) with independent custom TE libraries. Chromosome one-to-one alignments were performed with LASTZ, with parameters set as: K = 2200, L = 6000, E = 30, H = 0, O = 400, and T = 1. The resultant alignment blocks were further parsed with the Chain/Net package (Kent et al., 2003) as described previously (Chen et al., 2013). A custom Perl script (https://github.com/yiliao1022) was used to identify syntenic blocks, requiring neighboring blocks to be no more than five blocks apart. Manual inspection was conducted if necessary. Syntenic blocks identified for 2 Mb on each side of centromeres were extracted and used for constructing the synteny map.

Gene Synteny Map

We used syntenic and orthologous genes as markers to construct the synteny map between O. sativa and O. brachyantha at each centromeric region, using appropriate species as outgroups, such as Leersia perrieri, Brachypodium distachyon, Setaria italica, Sorghum bicolor, and/or Zea mays (Supplemental Methods).

Centromere Satellite Repeats for L. perrieri

Approximate 2-Mb sequences from the pseudomolecules of L. perrieri (Stein et al., 2018) corresponding to the homologous regions of rice centromeres were searched by Tandem Repeat Finder (Benson, 1999). The most abundant tandem repeat sequences found were the satellite repeats with a unit size of 180 bp, which were exclusively mapped at the homologous regions of rice centromere.

Sequence Annotation for O. brachyantha Cen12 Region

We annotated genes, transposable elements, and inter/intrasegmental duplications in the O. brachyantha Cen12 region (~1.2 Mb). Gene models identified using the MAKER v2.31 annotation engine (http://www. yandell-lab.org/software/maker.html), incorporating homology, expression evidence, and ab initio gene prediction methods. TE annotation is described as the above. Segmental duplications refer to paralogous sequences with coverage >1 kb and identity >90%. More details are provided in Supplemental Methods.

Syntenic Gene Losses

In this study, syntenic gene loss refers to a gene that has been lost in a syntenic position between O. sativa and O. brachyantha. The gene may be either completely lost from the genome or moved to other genomic locations. To identify syntenic gene losses in O. sativa after its divergence from O. brachyantha, we searched genes that are syntenically conserved between O. brachyantha and at least one outgroup species, L. perrieri and/or B. distachyon, but absent at the corresponding syntenic region in O. sativa. To filter false positives that may result from a missing annotation or sequence gap, we conducted a three-step process as previously described (Schnable et al., 2012). To place gene loss events on the branches of the Oryza phylogenetic tree leading to O. sativa, we also assessed these events in four internal branch species, including O. glaberrima, O. glumaeapativa, O. meridionalis, and O. punctata, which represent an evolutionary gradient.

Accession Numbers

The CenH3 ChIP-seq 454 reads have been deposited in GenBank under accession number SRX4224850. The monomer sequence of centromeric satellite repeat of L. perrieri has been deposited in EMBL under accession number LS9738733.1. The assembly of BAC clones are available at https://de.cyverse.org/dl/d/A07342B7-6E3D-485D-8605-867C93509F74/Obrac_Centromere_BAC_seq.fasta.

Supplemental Data

Supplemental Figure 1. Alignment of the in silico reference sequence motif map for the pericentromeric regions (upper panel) and the map of the same regions produced by genome mapping (lower panel).

Supplemental Figure 2. Plot of CenH3 ChIP-seq reads along the O. brachyantha chromosomes.

Supplemental Figure 3. BioNano genome maps aligned to chromosome 9.

Supplemental Figure 4. Comparative sequence analysis of the centromeres and flanking regions between O. sativa and O. brachyantha.

Supplemental Figure 5. Phylogenetic tree and divergence times of some grass species

Supplemental Figure 6. Synteny map of orthologous genes at Cen1 and Cen8 regions.

Supplemental Figure 7. Gene synteny at the centromeric regions of chromosome 2 between O. sativa and O. brachyantha, and the homologous region (scaffold_1: 16.6–21.6 Mb) from S. italica (outgroup).

Supplemental Figure 8. Gene synteny at the centromeric regions of chromosome 3 in rice (O. sativa) and O. brachyantha, and the homologous regions from S. italica (outgroup 1) and S. bicolor (outgroup 2).

Supplemental Figure 9. Gene synteny at the centromeric region of chromosome 4 in rice (O. sativa) and O. brachyantha, and the homologous regions from B. distachyon (Bd4: 51.80–68.45 Mb) and S. bicolor (Chromosome 6: 6.68–10.11 Mb).

Supplemental Figure 10. Gene synteny at the centromeric region of chromosome 5 in rice (O. sativa), O. brachyantha, and L. perrieri, and the homologous region from S. italica (scaffold3: 28.7–34.4 Mb).

Supplemental Figure 11. Gene synteny at the centromeric region of chromosome 6 in rice (O. sativa) and O. brachyantha, as well as the homologous region from S. italica (scaffold 4: 18.50–22.87 Mb).

Supplemental Figure 12. Gene synteny at the centromeric region of chromosome 10 in rice (O. sativa), O. brachyantha, and L. perrieri (outgroup).

Supplemental Figure 13. Gene synteny at the centromeric region of chromosome 11 in rice (O. sativa), O. brachyantha, and L. perrieri, and the homologous region from B. distachyon (Bd4: 22.30–23.22 Mb).
Supplemental Figure 14. Two contigs from the L. pennieri centromeric regions of chromosome 12 are verified by BAC end sequence alignments.

Supplemental Figure 15. Our targeted regions (10 syntenic genes on each side of the centromere) generally overlap with crossing-over suppressed pericentromeric regions in rice.

Supplemental Figure 16. Examples illustrating the identification of an ancestral or derived gene locus in O. sativa.

Supplemental Figure 17. A segment containing multiple genes from rice Cen4 has duplicated to the long arm of chromosome 4.

Supplemental Figure 18. Chromosome-specific distribution of CentoF.

Supplemental Table 1. Statistics of BAC-based assembly.

Supplemental Table 2. O. brachyantha pericentromeric region repeat counts by repeat classes.

Supplemental Table 3. O. sativa pericentromeric/centromeric region* repeat counts by repeat classes.

Supplemental Table 4. Comparison of the conserved gene content and local size variation at the centromeric/pericentromeric regions between O. sativa and O. brachyantha.

Supplemental Table 5. Comparison of 454 reads between CenH3 ChIP data set and whole-genome shotgun data set.

Supplemental Table 6. Targeted regions for comparative mapping.

Supplemental Table 7. Number of syntenic blocks detected using LASTZ alignments between O. sativa and O. brachyantha.

Supplemental Table 8. Segmental duplications within the O. brachyantha Cen12 region.

Supplemental Table 9. Nonsynonymous to synonymous substitution rates (ka/ks) between O. sativa centromeric moved genes and their homologs in the centromeric regions of O. brachyantha, and the expression level of the moved genes in seedling.

Supplemental Table 10. Summary of Oryza brachyantha raw BioNano molecule data.

Supplemental Table 11. Average sequence similarity (%) of CentO-F monomers within and between O. brachyantha centromeres.

Supplemental Data Set 1. List of BAC clones sequenced for the O. brachyantha pericentromeric and/or centromeric regions.


Supplemental Data Set 3. Major homologous/syntenic chromosomal segments of other grass species compared with O. sativa.

Supplemental Data Set 4. Genes identified that moved or were lost from the rice genome after its split with O. brachyantha.

Supplemental Data Set 5. Investigating of the most 240 centromere-proximal genes in O. sativa.

Supplemental Data Set 6. Genes identified that moved or were lost from the O. brachyantha pericentromeric regions after it split with O. sativa.

Supplemental Data Set 7. Gene synteny at 12 centromeric regions.

Supplemental Methods.

Supplemental Text 1. Centromeric satellite repeat sequence (CentO-F) in Oryza brachyantha.

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AUTHOR CONTRIBUTIONS

Y.L. and M.C conceived and carried out the experiments, analyzed the data, and wrote the article. X.Z., B.L., T.L., J.C., Z.B., M.W., and J.S. carried out the experiments and analyzed the data. J.G.W. and J.J. provided CenH3 ChIP-seq data and edited the article. R.A.W. provided unpublished Oryza genome assemblies and edited the article. All authors read and approved the final manuscript.

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Comparison of *Oryza sativa* and *Oryza brachyantha* Genomes Reveals Selection-Driven Gene Escape from the Centromeric Regions

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