

RESEARCH ARTICLES

Conservation and Purifying Selection of Transcribed Genes Located in a Rice Centromere^W

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Recombination is strongly suppressed in centromeric regions. In chromosomal regions with suppressed recombination, deleterious mutations can easily accumulate and cause degeneration of genes and genomes. Surprisingly, the centromere of chromosome8 (*Cen8*) of rice (*Oryza sativa*) contains several transcribed genes. However, it remains unclear as to what selective forces drive the evolution and existence of transcribed genes in *Cen8*. Sequencing of orthologous *Cen8* regions from two additional *Oryza* species, *Oryza glaberrima* and *Oryza brachyantha*, which diverged from *O. sativa* 1 and 10 million years ago, respectively, revealed a set of seven transcribed *Cen8* genes conserved across all three species. Chromatin immunoprecipitation analysis with the centromere-specific histone CENH3 confirmed that the sequenced orthologous regions are part of the functional centromere. All seven *Cen8* genes have undergone purifying selection, representing a striking phenomenon of active gene survival within a recombination-free zone over a long evolutionary time. The coding sequences of the *Cen8* genes showed sequence divergence and mutation rates that were significantly reduced from those of genes located on the chromosome arms. This suggests that *Oryza* has a mechanism to maintain the fidelity and functionality of *Cen8* genes, even when embedded in a sea of repetitive sequences and transposable elements.

INTRODUCTION

Recombination via chromosomal crossing-over plays a significant role in gene and genome evolution (Gaut et al., 2007; Li et al., 2007). In chromosomal regions with suppressed or reduced recombination from crossing-over, deleterious mutations can easily accumulate due to inefficient natural selection caused by Hill-Robertson Interference (Haddrell et al., 2007; Comeron et al., 2008; Betancourt et al., 2009; Charlesworth et al., 2009). Such interference is thought to be a major factor leading to genetic degeneration of genes and genomes. Suppression or reduction of recombination resulting from structural rearrangements along chromosomes can also prevent gene flow and hinder the introgression of alleles, thereby contributing to speciation and/or persistence as demonstrated by many empirical studies (Noor et al., 2001; Rieseberg, 2001; Ortiz-Barrientos et al., 2002; Navarro and Barton, 2003; Butlin, 2005; Stump et al., 2005).

Recent studies have demonstrated the presence of active genes in recombination-suppressed chromosomal domains of mammals (Mudge and Jackson, 2005), *Drosophila mela-*

nogaster (Hoskins et al., 2002), and plants (Haupt et al., 2001; Yan et al., 2005, 2008), which raises questions as to how the fidelity and function of such genes are maintained in an environment presumed to be void of recombination. Maintenance of structure and function of genes in the human Y chromosome has been shown to occur by intrachromatid gene conversion mediated via homologous recombination between opposing arms of large palindromic sequences (Lange et al., 2009). In *Drosophila*, regional and ancient recombination events in heterochromatin have been postulated to maintain heterochromatic genes (Schulze et al., 2006). Polymorphism and divergence data from chromosome 4 of *Drosophila*, which was initially believed to be recombination suppressed, revealed the presence of extremely low levels of recombination; this reduced level was sufficient to maintain normal gene density and gene functionality (Arguello et al., 2010).

Centromeres are defined by the presence of a centromere-specific histone variant CENH3 (CID in *Drosophila*, CENP-A in humans) (Allshire and Karpen, 2008). Unlike other chromosomal domains, such as pericentromeric heterochromatin, in which self- or low-frequency recombination events have been observed, centromeres are thought to be completely devoid of crossover recombination (Beadle, 1932; Lambie and Roeder, 1986; Jackson et al., 1996; Anderson et al., 2003; Shi et al., 2010). Interestingly, sequence and transcriptome analysis of the centromere of rice (*Oryza sativa*) chromosome8 (*Cen8*) revealed the presence of 16 transcribed genes (Nagaki et al., 2004; Yan et al., 2005); the first set of genes found in the functional domain of a eukaryotic centromere. An intriguing question emerged from this discovery: What selective forces are driving the evolution and existence of transcribed genes in genomic regions devoid of crossover recombination?

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To address this question, we sequenced *Cen8* in two additional *Oryza* species, *Oryza glaberrima* and *Oryza brachyantha*, which have diverged from rice for ~ 0.5 to 1 and ~ 10 to 15 million years (MY), respectively (Ammiraju et al., 2008). The centromeres of *O. brachyantha* chromosomes were previously demonstrated to contain completely different sets of repetitive DNA sequences compared with cultivated rice (Lee et al., 2005; Gao et al., 2009). Here, we demonstrate the persistence of orthologous transcribed *Cen8* genes in these three *Oryza* species. The conserved *Cen8* genes showed strong functional constraints in both *O. glaberrima* and *O. brachyantha*, representing a striking phenomenon of active gene survival in a recombination-free zone over a 10 to 15 MY evolutionary time span.

RESULTS

Identification of Active Genes in *Cen8* of *O. glaberrima* and *O. brachyantha*

We sequenced and assembled *Cen8* sequences of *O. glaberrima* (1.3 Mb) and *O. brachyantha* (1.1 Mb) corresponding to ~ 1 Mb of the *O. sativa* ssp *japonica* var *Nipponbare* (referred to hereafter simply as *O. sativa*) *Cen8* (Nip-*Cen8*), including the ~ 750 -kb CENH3 binding domain (Yan et al., 2008). The crossover-suppressed domain in *Cen8* is 2312 kb and encompasses this 750-kb CENH3 binding domain (Yan et al., 2005). The pseudomolecules of both species included centromere-specific satellite repeats of unknown size.

The 1-Mb sequence of *O. sativa* contains 17 transcribed genes based on expression evidence and gene ontology (Yan et al., 2005) (see Supplemental Table 1 online; *Cen8.t00802* was not described in Yan et al., 2005). To determine the extent of gene conservation in orthologous *Cen8* regions of *O. glaberrima* and *O. brachyantha*, we manually annotated these regions and identified 15 of the 17 (88%) orthologous genes in *O. glaberrima* (13 with intact open reading frames and two pseudogenes [*Cen8.t00969* and *Cen8.t01075*]) and 7 of the 17 (41.2%) orthologous genes in *O. brachyantha* (Figure 1, Table 1). Two orthologous genes in *O. glaberrima* (*Cen8.t00969* and *Cen8.t01075*) appeared to be pseudogenes as each had single base pair deletions that resulted in frame-shift mutations in their protein coding sequences (Figure 1). The order and orientation of all orthologous *Cen8* genes were conserved across the three *Oryza* species (Figure 1).

Nine *Cen8* genes conserved between *O. sativa* and *O. glaberrima* and seven conserved *O. brachyantha* *Cen8* genes were tested for transcriptional activity using cDNA synthesized from both leaf and root tissues. Genomic DNA and an RT negative control (produced using cDNA synthesis reaction without addition of reverse transcriptase) were also included in the analysis as controls. Each of the nine primer sets in *O. glaberrima/O. sativa* and the seven sets in *O. brachyantha* faithfully amplified products of predicted size from both tissues, except *Cen8.t00793* in *O. brachyantha*, which produced a much larger genomic product (Figure 2; for primers, see Supplemental Table 2 online). Furthermore, transcripts were also detected for all seven *O. brachyantha* *Cen8* genes based on Illumina-based transcriptome profiling data obtained from leaf and root tissues (see Supple-

mental Table 3 online). Overall, by combining RT-PCR results, transcriptome analysis, and an EST search analysis, our data showed that 12 of the 15 orthologous *O. glaberrima* *Cen8* genes and all seven *O. brachyantha* *Cen8* genes were transcribed (see Supplemental Table 4 online).

Confirmation of the Centromeric Position of the Orthologous *Cen8* Sequences from *O. glaberrima* and *O. brachyantha*

We conducted chromatin immunoprecipitation (ChIP) experiments to confirm the centromeric location of the *Cen8* sequences produced from *O. glaberrima* and *O. brachyantha*. PCR primers were designed from regions near the seven conserved *Cen8* genes in both species (see Supplemental Table 5 online). ChIP was performed using an antibody against rice CENH3 (Nagaki et al., 2004). Here, we surveyed the *Cen8* landscape in *O. glaberrima* and *O. brachyantha* for evidence of CENH3 binding. Eight *O. glaberrima* and four *O. brachyantha* primers were designed to interrogate this region spanning ~ 1 Mb of *Cen8* sequence. In *O. glaberrima*, 13 of the 15 conserved *Cen8* genes are localized within a region where eight primer sets showed significant CENH3 enrichment using ChIP-PCR (Figure 1). The remaining two genes, *Cen8.t01152* and *Cen8.t01153*, are 100 kb away from one of the ChIP-PCR markers (Figure 1). In *O. brachyantha*, significant CENH3 enrichment of the centromere specific CentO-F satellite array was confirmed (Figure 1). DNA sequences outside of the seven *Cen8* genes were not enriched by ChIP-PCR. However, active genes are associated with H3 (rather than CENH3) nucleosomes (Yan et al., 2008); thus, the *Cen8* genes cannot be used for testing CENH3 enrichment. Since the *O. brachyantha* *Cen8* genes are only 500 kb away from the CentO-F satellite, we predict that these genes are either within the CENH3 domain or immediately outside of the CENH3 domain, depending on the size of the CentO-F satellite array in this centromere. Based on the fact that the crossover suppressed domains are severalfold larger than the CENH3 domains in all rice centromeres (Yan et al., 2008), we presumed that all *O. brachyantha* *Cen8* genes are located within the crossover-suppressed domain in this centromere.

Purifying Selection of Conserved Orthologous *Cen8* Genes in Three *Oryza* Species

Annotation and expression data revealed the presence of seven conserved and transcribed orthologous genes between two independently domesticated rice species and their distant wild relative *O. brachyantha*. To determine if any of these genes were under functional constraints, the ratios of nonsynonymous substitution rate (K_a) and synonymous substitution rate (K_s) for all seven *Cen8* orthologs were calculated (Table 1). The results showed that all K_a/K_s (ω) ratios between either *O. sativa* and *O. brachyantha* (OS versus OB) or *O. glaberrima* and *O. brachyantha* (OG versus OB) were <0.5 (average 0.1811 for OS versus OB, and 0.1625 for OG versus OB) (Table 1), suggesting that all seven genes are under strong functional constraint with purifying selection. Likelihood ratio tests (LRTs) for all seven K_a/K_s values deviated significantly from neutrality ($\omega = 1$).

K_a/K_s ratios for the seven genes between *O. sativa* and *O. glaberrima* as well as the six remaining genes shared by these

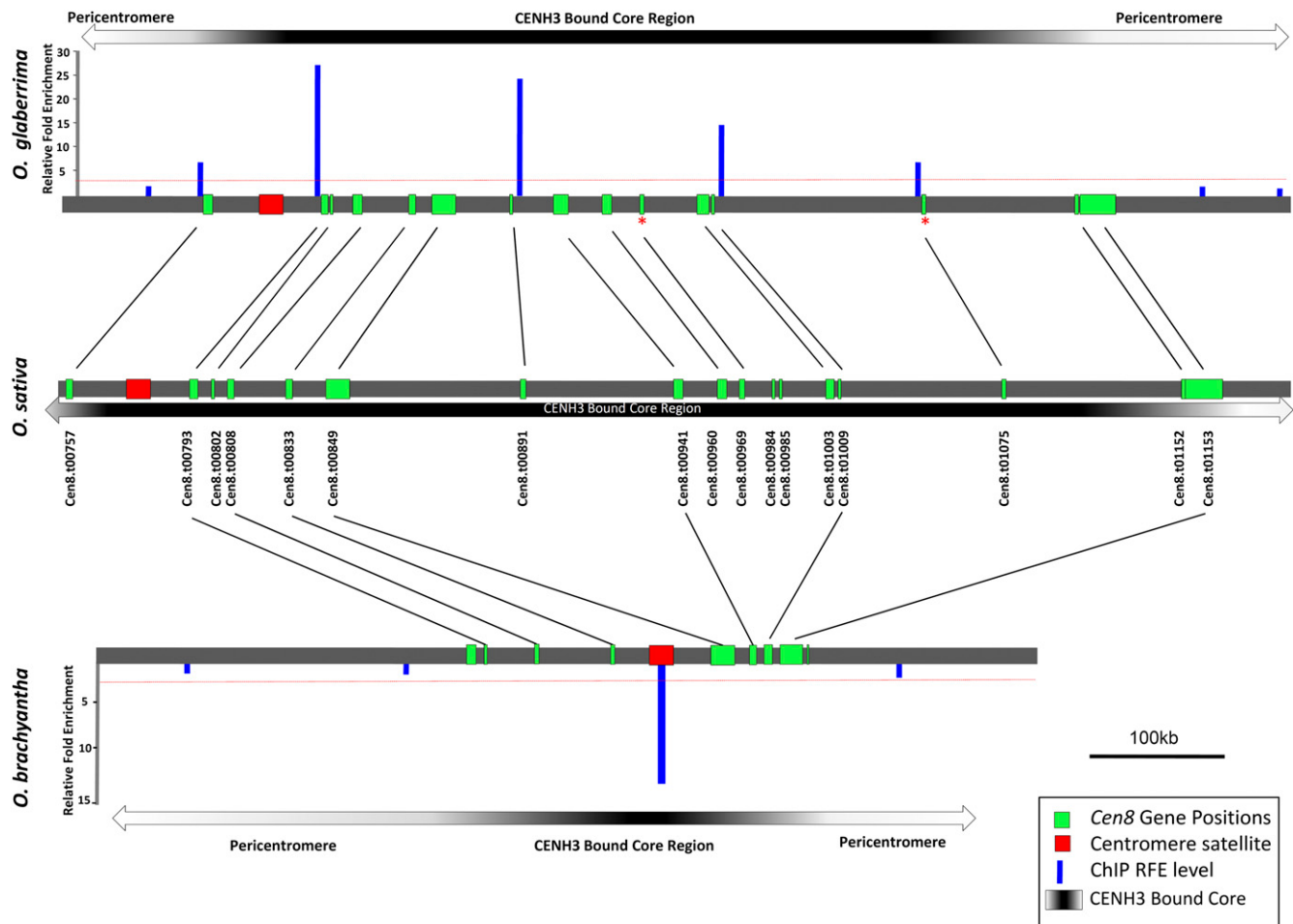


Figure 1. ChIP Analysis and Orthologous *Cen8* Gene Alignment in *O. sativa*, *O. glaberrima*, and *O. brachyantha*.

O. sativa Refseq and gene annotation from the Rice Genome Annotation database were used as references, and *O. glaberrima* and *O. brachyantha* orthologous genes are matched by straight lines. Two *O. glaberrima* orthologous pseudogenes are marked with a red asterisk and contain single frame-shift point mutations. The size of the centromere satellite (CentO in *O. sativa* and *O. glaberrima* and CentO-F in *O. brachyantha*) was not drawn to scale as the full sequence could not be determined. The CENH3 binding functional domains are shown as gray shaded bars. The black to white gradient illustrates the decrease in CENH3 binding from the satellite core of the centromere to more sparsely bound regions approaching the pericentromere. The red dashed lines indicate the ChIP significance level at a P value of 0.01.

two species indicated that 10 genes were under purifying selection. The remaining three genes (*Cen8.t00802*, *Cen8.t00808*, and *Cen8.t00960*) had K_a/K_s ratios > 1 (Table 1). However, LRT tests for all 13 comparisons between *O. sativa* and *O. glaberrima* did not deviate significantly from 1, suggesting that the short divergence time between these two species limits the statistical power of this analysis.

Low Molecular Evolution Rates of *Cen8* Coding Sequences versus Noncentromeric Genes

To investigate the pattern of molecular evolution of *Cen8* genes versus genes located in recombining regions, we analyzed orthologous gene sets derived from sequenced short arms of chromosome 3 (Chr3S) of *O. sativa*, *O. glaberrima*, and *O. brachyantha*. Comparisons between *O. sativa* and *O. glaberrima* were made

using 1515 orthologous gene pairs (see Supplemental Figure 1 online). These Chr3S genes were scattered along the entire chromosome arm excluding the recombination free centromeric region. Both the mean K_s and K_a values for all 1515 pairs were significantly higher than that of the 13 pairs of *Cen8* genes ($P = 7.7E-11$ for K_s and $P = 0.00083$ for K_a , one-sided t test; Table 2), and tests of mutation rate in the coding region (/bp/MY) yielded similar results ($P = 1.35E-5$; Table 2). For *O. brachyantha*, we selected a total of 268 and 230 Chr3S genes that are orthologous to *O. sativa* and *O. glaberrima* genes, respectively (see Supplemental Figures 2 and 3 online). The average K_s of orthologous genes pairs between *O. sativa* and *O. brachyantha* and between *O. glaberrima* and *O. brachyantha* (Supplemental Figures 2 and 3 online) was significantly higher ($P = 0.00182$, $3.62E-5$, respectively, one-sided t test) than that for the seven pairs of *Cen8* genes (Table 2). However, the K_a of *Cen8* genes and Chr3S genes were not

Table 1. Sequence Divergence of Orthologous *Cen8* Active Genes

OS_CDS	TIGR Gene ID	Comparison	Identity	Length	Ka	Ks	Ka/Ks	LRT P Value
<i>Cen8.t00757</i>	Os08g21660	OG-OS	99.9	978	0.0000	0.0034	0.0010	0.1261
<i>Cen8.t00793</i>	Os08g21700	OG-OS	99.7	576	0	0.0106	0.0000	0.0365
		OB-OS	89.9	555	0.0827	0.1908	0.4332	0.0044**
		OG-OB	89.5	555	0.0830	0.2062	0.4026	0.0016**
<i>Cen8.t00802</i>	Os08g21720	OG-OS	98.6	285	0.0207	0.0000	∞	0.0928
<i>Cen8.t00808</i>	Os08g21760	OG-OS	99.8	483	0.0026	0.0000	∞	0.4997
		OB-OS	92.8	483	0.0203	0.2949	0.0689	0.0000***
		OG-OB	93.5	483	0.0116	0.2818	0.0412	0.0000***
<i>Cen8.t00833</i>	Os08g21840	OG-OS	99.5	885	0.0035	0.0069	0.5010	0.4949
		OB-OS	90.5	873	0.0464	0.2621	0.1770	0.0000***
		OG-OB	92.1	873	0.0286	0.2420	0.1182	0.0000***
<i>Cen8.t00849</i>	NA	OG-OS	98.4	567	0.0142	0.0223	0.6384	0.5475
		OB-OS	93.3	570	0.0222	0.2223	0.0997	0.0000***
		OG-OB	93.5	567	0.0198	0.2255	0.0878	0.0000***
<i>Cen8.t00891</i>	Os08g22060	OG-OS	100	102	0.0000	0.0000	∞	0.9980
<i>Cen8.t00941</i>	Os08g22149	OG-OS	99.7	705	0.0020	0.0050	0.4015	0.5288
		OB-OS	91.0	696	0.0538	0.2215	0.2431	0.0000***
		OG-OB	90.7	696	0.0560	0.2170	0.2442	0.0000***
<i>Cen8.t00960</i>	Os08g22200	OG-OS	99.7	882	0.0049	0.0000	∞	0.1515
<i>Cen8.t01003</i>	Os08g22354	OG-OS	99.8	1980	0.0014	0.0037	0.3891	0.3603
		OB-OS	93.5	1968	0.0233	0.2235	0.1043	0.0000***
		OG-OB	93.6	1968	0.0234	0.2170	0.1077	0.0000***
<i>Cen8.t01009</i>	NA	OG-OS	99.0	294	0.0096	0.0121	0.7918	0.7152
<i>Cen8.t01152</i>	Os08g22852	OG-OS	99.8	564	0.0000	0.0046	0.0000	0.1639
<i>Cen8.t01153</i>	Os08g22864	OG-OS	99.7	3174	0.0020	0.0048	0.4106	0.2465
		OB-OS	95.0	3108	0.0211	0.1487	0.1416	0.0000***
		OG-OB	95.0	3108	0.0206	0.1511	0.1361	0.0000***

NA, not available; OB, *O. brachyantha*; OG, *O. glaberrima*; OS, *O. sativa*; TIGR, The Institute for Genomic Research. ∞, infinite when there is no synonymous substitution and therefore the denominator (Ks) as 0; **, significant level as $P < 0.01$; ***, significant level as $P < 0.001$.

statistically different (Table 2). The mutation rate across the entire Chr3S of *O. brachyantha* revealed nearly significant higher mutation rates than those found for all conserved active *Cen8* genes across the three *Oryza* species ($P = 0.169$ and 0.059) (Table 2).

Functional Constraints of *Cen8* Genes Suggested by *O. glaberrima* and *O. brachyantha* *Cen8* Polymorphism Patterns

To obtain supporting evidence for purifying selection of the conserved *Cen8* genes, we performed a population genetic analysis using two data sets. First, we obtained *O. glaberrima* *Cen8* region polymorphisms using whole-genome single nucleotide polymorphism (SNP) data from eight accessions of *O. glaberrima*. In a 651,584-bp region of *O. glaberrima* *Cen8*, we found a total of 388 SNPs giving a polymorphism rate of 0.595/kb, which is equal to half of the average polymorphism rate across the whole genome (1.118/kb). Of the 388 SNPs in *Cen8*, 358 SNPs were located in intergenic regions, and only 30 were present in *Cen8* genes. Of the 30 SNPs, only five were located in exons, including one nonsynonymous substitution and four synonymous substitutions (see Supplemental Table 6 online). Statistical tests indicated that both an excess of SNPs in *Cen8* genes and synonymous substitutions in coding regions significantly devi-

ated from neutral expectations (see Supplemental Tables 7 and 8 online).

Second, we amplified and sequenced a complete orthologous *Cen8* gene (*OB_t00833* [3028 bp, excluding gaps]) and part of the *OB_t01153* (1022 bp) gene, from 15 *O. brachyantha* accessions. Sequence analysis from pooling two sequence regions revealed the presence of 35 polymorphic sites (31 in *OB_t00833* and four in *OB_t01153*), 30 of which were in noncoding sequences (27 in *OB_t00833* and three in *OB_t01153*) and five of which were in coding sequences (four in *OB_t00833* and one in *OB_t01153*): four as synonymous substitutions and one as a replacement mutation (see Supplemental Figure 4 online). Having both an excess of polymorphic sites in noncoding regions and an excess of synonymous substitutions in coding sequences significantly deviated from neutral expectations (χ^2 test, $P = 0.048$ and 0.029 , respectively; see Supplemental Tables 7 and 8 online), thus supporting our observation that the conserved *OB_t00833* and *OB_t01153* coding sequences are under strong functional constraints.

We conducted additional population genetic analyses using the polymorphisms identified from the eight *O. glaberrima* accessions and 15 *O. brachyantha* accessions to infer the evolutionary pattern of *Cen8* genes in *O. glaberrima* and *O. brachyantha*. Each of the three recombination and gene conversion tests we performed failed to reject the null hypothesis

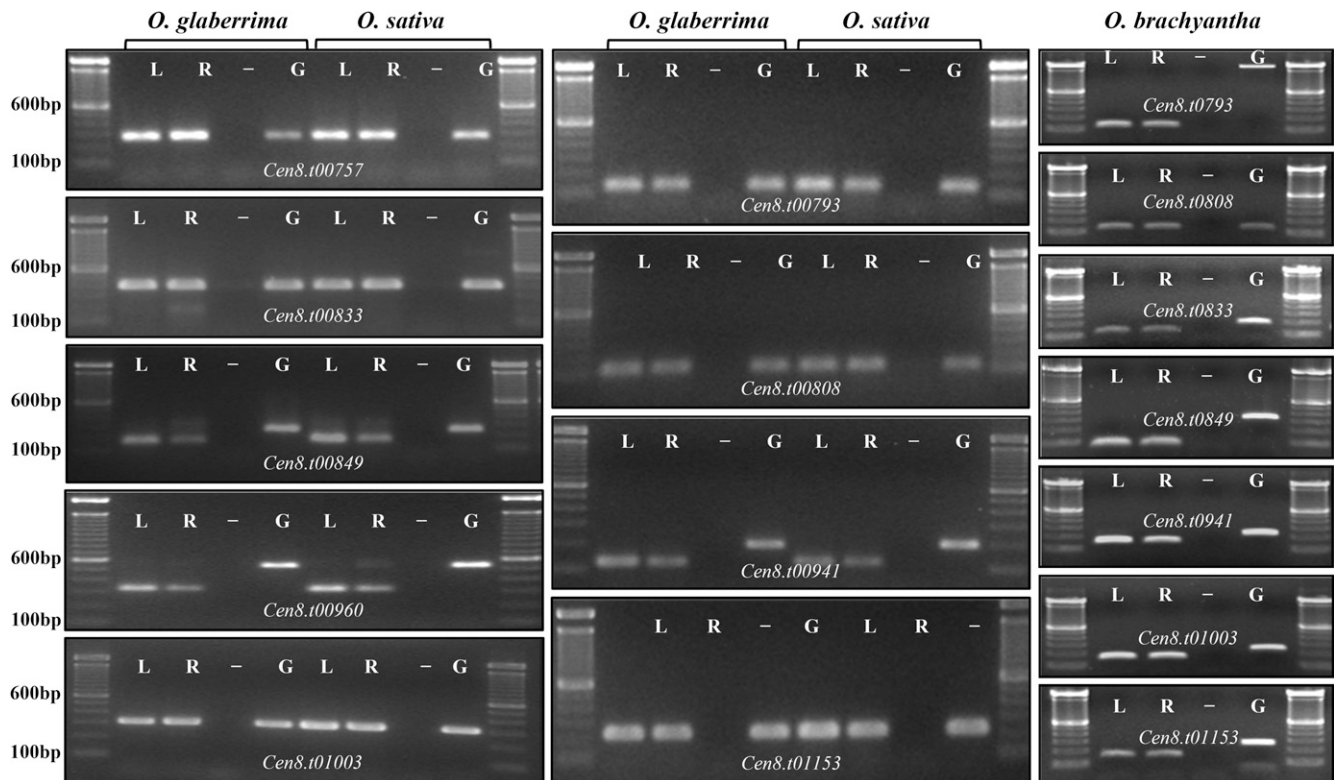


Figure 2. RT-PCR Results of *Cen8* Genes from *O. sativa*, *O. glaberrima*, and *O. brachyantha*.

L, leaf; R, root; (-), negative control; G, genomic DNA.

of no recombination (see Supplemental Tables 9 to 11 online), indicating that no recombination or conversion within these centromeric genes could be detected. A negative Tajima's D value in *O. glaberrima* significantly deviated from neutrality, indicating *O. glaberrima* *Cen8* is likely undergoing purifying selection (Table 3). Tajima's D values inferred from either silent polymorphic sites or total polymorphic sites in *O. brachyantha* were positive but did not deviate from neutral expectations ($P = 0.84$) (Table 3).

Pfam Analysis of Seven Conserved *Cen8* Genes

The presence of seven highly conserved genes within a functional centromere, spanning the *Oryza* phylogeny, may indicate that these genes are biologically essential. Pfam analysis (<http://pfam.sanger.ac.uk/>) of these genes revealed that five of the seven (except *Cen8.t00793* and *Cen8.t00849*) belong to gene families assigned to physiological and/or cellular functions (see Supplemental Table 12 online). In particular, *Cen8.t01003* (Os08g22354) was found to contain nine conserved domains: four RNA/DNA binding sites, four RRM dimerization sites, and poly-adenylate binding protein, providing scaffolds to which proteins can bind and mediate processes such as export, translation, transcript turnover, and regulation of development at the transcriptional level (Bandziulis et al., 1989; Birney et al., 1993; Mangus et al., 2003).

DISCUSSION

Centromeres in higher eukaryotes are embedded within highly heterochromatic pericentromeric chromatin. In most plant and animal species described, centromeres contain satellite repeats and transposable elements (Henikoff et al., 2001; Jiang et al., 2003) and are nonrecombinogenic. These attributes have hindered centromere research, especially at the sequence level. In fact, virtually all whole-genome shotgun assemblies have completely ignored centromeres as they are difficult to recognize and assemble. The genus *Oryza* provides a unique model for centromere research for three reasons. First, the centromeres of several rice chromosomes have been fully or partially sequenced (Zhang et al., 2004; Yan et al., 2008; Wu et al., 2009). Second, transcribed genes located within CENH3-associated chromatin domains in rice provide a platform to study the evolution of genes located in recombination suppressed chromosomal domains. Third, a set of BAC-based physical maps representing 13 *Oryza* species and all 10 *Oryza* genome types (six diploids and four polyploids) has been developed, thereby providing unprecedented access to virtually any region of the collective *Oryza* genome for interrogation, including centromeres (Ammiraju et al., 2006; Kim et al., 2008).

We sequenced and compared the *Cen8* regions of *O. glaberrima* and *O. brachyantha* that span the CENH3 binding domains and contain centromere-specific satellite repeats. We demonstrated that *O. glaberrima* shared 12 active *Cen8* genes with

Table 2. Statistical Tests of Synonymous (Ks) and Nonsynonymous (Ka) Substitution Rates and Mutation Rates between *Cen8* Genes and Chr3S Noncentromeric Genes

Test	Gene	OS versus OG	OS versus OB	OG versus OB
Ks	Chr3S genes	0.0287 (1515)	0.3000 (268)	0.3245 (238)
	<i>Cen8</i> genes	0.0056 (13)	0.2234 (7)	0.2219 (7)
Welch <i>t</i> test P value (one tail)		7.716E-11***	0.00182**	3.62E-5***
Ka	Chr3S genes	0.0113 (1515)	0.0436 (268)	0.0505 (238)
	<i>Cen8</i> genes	0.0043 (13)	0.0385 (7)	0.0347 (7)
Welch <i>t</i> test P value (one tail)		0.000828***	0.3000	0.0796
Mutation rate (/bp/MY)	Chr3S gene	0.0066 (1515)	0.00466 (268)	0.00486 (238)
	<i>Cen8</i> genes	0.00250 (13)	0.00393 (7)	0.00381 (7)
Welch <i>t</i> test P value (one tail)		1.3553E-5***	0.169	0.0594

OB, *O. brachyantha*; OG, *O. glaberrima*; OS, *O. sativa*. **, significant level as $P < 0.01$; ***, significant level as $P < 0.001$. The numbers in parentheses indicate the number of genes used for each test.

O. sativa-Cen8, whereas only six orthologous *Cen8* genes were shared between the two subspecies *O. sativa ssp japonica* and *O. sativa ssp indica* (Wu et al., 2009). A significant finding was that seven active genes were conserved in *O. brachyantha*, which diverged roughly 10 to 15 MY ago from *O. sativa* and *O. glaberrima*. This observation raises important questions regarding gene loss, gene gain, and gene mutation in the three *Oryza* centromeres. It is unknown if the ancestral state of *Cen8* more closely resembled *O. brachyantha Cen8* or *O. sativa Cen8*. If the structure of *O. sativa Cen8* reflects the ancestral state, then both *O. glaberrima* and *O. brachyantha* have undergone gene loss (2 to 10 genes, respectively) and pseudogenization in *Cen8*. Alternatively, if *O. brachyantha Cen8* is a closer to reflection of the ancestral state, then *O. glaberrima Cen8* has acquired additional expressed genes, and *O. sativa Cen8* is still acquiring genes. The latter scenario is highly unlikely because centromeres are thought to have evolved from noncentromeric regions via neocentromere formation and accumulation of repetitive DNA (Nagaki et al., 2004). In addition, an analysis of sequences flanking the *O. glaberrima Cen8* genes did not reveal the presence of *helitron* or MULE sequences, which could be used to explain such gene acquisition.

The most surprising discovery was that all seven genes were not only transcribed but appear to be under strong purifying selection based on two lines of evidence (Ka/Ks ratios and a population genetics analysis of centromeric genes). Such syntenic conservation and purifying selection implies that the fidelity of centromeric genes can be preserved without crossover recombination, even when embedded in a sea of highly dynamic and constantly evolving transposable elements and tandem satellite repeats.

It is generally understood that low recombination rates will reduce sequence diversity due to rapid elimination or fixation of mutations (Nachman, 2002). We tested the synonymous substitution rate, which is assumed to be neutral, between three *Oryza* species. The significantly smaller Ks rates for centromeric genes compared with noncentromeric genes are consistent with the observation of low intraspecific polymorphisms detected in centromeric regions from various organisms, including yeast, maize (*Zea mays*), *Drosophila*, and rice (Aguade et al., 1989; Begun and Aquadro, 1992; Gerton et al., 2000; Gore et al., 2009; Schacherer et al., 2009). It is known that recombination can

facilitate chromosomal rearrangements, gene copy number changes, and even the generation of single-nucleotide mutations (Lercher and Hurst, 2002; Hellmann et al., 2003; Jelesko et al., 2004; Schuermann et al., 2005). Therefore, given the lack of crossover recombination in centromeres, a reduced mutation rate could also contribute to low sequence divergence rates of centromeric genes.

Speciation and species differentiation may be enhanced by the suppression of recombination. Both modeling and recent empirical studies suggest that recombination can reduce speciation events (Noor et al., 2001; Ortiz-Barrientos et al., 2002), while suppression of recombination can allow species to diverge by preventing gene flow between individuals. For example, in *Drosophila*, the hybrid incompatibility genes *Lhr*, *Zhr*, and *OdsH* are associated with speciation, and all map to recombinationally suppressed pericentric and heterochromatic regions that showed reduced or undetectable levels of recombination (Sawamura et al., 1993; Brideau et al., 2006; Bayes and Malik, 2009). In plants, several sets of genes involved in speciation and reproductive isolation have been localized to highly heterochromatic regions where recombination is suppressed. For example, the "A" locus for gametophytic apomixis, a phenomenon that results in asexual reproduction, was identified in a region completely devoid of recombination (Ozias-Akins and van Dijk, 2007). Moreover, self-incompatibility genes (S-locus) were found to be recombinationally suppressed due to their subcentromeric location in *Petunia* (Coleman and Kao, 1992; Entani et al., 1999).

Table 3. Average Sequence Diversity of *O. brachyantha Cen8* Genes and *O. glaberrima Cen8* Region

Statistic	OB_t00833 and OB_t01153	OG_Cen8
Sample size	16	8
Length (bp)	4,427	651,484
Segregating sites (S)	35	388
θ_w per kb	3.17	0.23
θ_π per kb	2.66	0.15
Tajima's D	0.82494, P = 0.839	-1.84077, P = 0.00001**

OB, *O. brachyantha*; OG, *O. glaberrima*. **, significant level as $P < 0.01$.

and *Antirrhinum* (Ma et al., 2003; Yang et al., 2007), the presence of repetitive DNA in *Nicotiana* (Matton et al., 1995), and conserved linkage in *Prunus* (Ikeda et al., 2005). It is believed that S-locus genes experienced strong balancing selection that resulted in high local population polymorphisms but low population differentiation (Ruggiero et al., 2008). The low levels of sequence divergence found between *Cen8* genes in our three species comparisons, along with the population genetic data analysis from the *O. brachyantha* accessions, suggests that the conserved genes found within these functional centromeres have undergone molecular evolutionary events similar to those observed in S-locus genes. Functional assays of the conserved *Cen8* genes may provide new evidence that centromeres serve as islands of speciation (Noor and Bennett, 2009).

METHODS

Sequencing of BAC Tiles from *Cen8* Centromeric Regions of *Oryza glaberrima* and *Oryza brachyantha*

We used the rice (*Oryza sativa*) *Cen8* sequence (~1 Mb DNA), which includes the 750-kb CENH3 binding domain (Yan et al., 2008), as a reference sequence to identify *Cen8* genomic regions of *O. glaberrima* and *O. brachyantha*. Minimum tiling paths of overlapping BAC clones spanning *Cen8* in the two species were developed, including 12 (*O. glaberrima*) and 9 (*O. brachyantha*) BACs. Each BAC was shotgun Sanger sequenced and finished using previously described methods (Project, 2005). The *O. glaberrima* and *O. brachyantha* BAC sequences were assembled into 1.3- and 1.0-Mb pseudomolecule sequences, respectively. Each individual BAC ID and GenBank accession number is listed in Supplemental Table 13 online.

ChIP Analysis to Determine Functional Centromeres

Nuclei were isolated from young leaf tissue of *O. glaberrima* and *O. brachyantha*, and ChIP was performed using antibodies against the centromere histone H3 (CENH3) of rice (Nagaki et al., 2004; Lee et al., 2005). As a negative control, a mock ChIP experiment was also performed, in tandem, by replacing the anti-CENH3 antibody with normal rabbit serum. Centromere sequences bound to CENH3 were identified with quantitative PCR using primers that spanned and flanked the proposed set of centromeric genes. Primers were designed to amplify products between 119 and 325 bp. Quantitative real-time PCR was conducted to determine the enrichment of centromere sequences within the ChIP samples compared with the mock. The quantitative real-time PCR was performed in triplicate using a DyNAmo HS SYBR Green qPCR kit (Finnzymes) using the following cycling parameters: 94°C for 15 min, 45 cycles of 95°C for 10 s, 60°C for 30 s, and 72°C for 30 s. The relative fold enrichment was calculated for each primer pair using a noncentromere primer set (NonCenControl 1) as a reference. For each primer pair, ΔC_t cycle threshold (C_t) for mock was calculated as $\Delta C_t(\text{mock}) = C_t(\text{centromere primer}) - C_t(\text{NonCenControl 1})$, and ΔC_t for ChIP was calculated as $\Delta C_t(\text{ChIP}) = C_t(\text{centromere primer}) - C_t(\text{noncentromere primer})$. Lastly, the relative fold enrichment ($2^{-\Delta\Delta C_t}$) was calculated, where $\Delta\Delta C_t = \Delta C_t(\text{ChIP}) - \Delta C_t(\text{mock})$. An enrichment cutoff line was placed based on P values assigned using a one-tailed Student's *t* test at a significance level of $\alpha = 0.01$.

Identification and Expression Analysis of Orthologous *Cen8* Genes in *O. sativa*, *O. glaberrima*, and *O. brachyantha*

Coding sequences (CDSs) of all annotated genes in the *O. sativa* *Cen8* region were used as queries to search for orthologous genes in the

O. glaberrima and *O. brachyantha* *Cen8* pseudomolecules using MEGA-BLAST (parameters: *O. glaberrima* e-value <1e-3, >95% sequence identity over the entire CDS; *O. brachyantha* e-value <1e-1, >90% sequence identity). Active genes were classified as those having RT-PCR and/or ESTs expression evidence. Gene ontology annotations were described in the Rice Genome Annotation database (<http://rice.plantbiology.msu.edu/cgi-bin/gbrowse/rice/>). Expression profiles of all *Cen8* genes in *O. sativa* were obtained from RT-PCR experiments (Yan et al., 2005) and/or the most recent whole-genome UniGene rice EST/mRNA data set (<http://www.ncbi.nlm.nih.gov/UniGene/UGOrg.cgi?TAXID=4530>). Evidence for expression of *O. glaberrima* *Cen8* genes was obtained using RT-PCR from leaf and/or root tissue. Evidence for expression of *O. brachyantha* *Cen8* genes was obtained by RT-PCR and analysis of an Illumina sequence-based genome-wide transcriptome data set derived from root and shoot cDNA, which were kindly provided by M. Chen at the Chinese Academy of Sciences, Beijing, China. Gene expression levels were expressed as reads (number of Illumina reads that mapped onto CDS without mismatch) and RPKM (number of reads per kilobase per million reads).

A total of nine non-transposable element related *O. glaberrima* *Cen8* genes and seven *O. brachyantha* *Cen8* genes found within *Cen8* in *O. sativa* were selected for transcriptional analysis via RT-PCR. Primer pairs were designed from conserved regions within the gene exons such that the same primers could be used to perform RT-PCR in both *O. sativa* and *O. glaberrima*. However, since the priming sites were not conserved between *O. glaberrima* and *O. brachyantha*, the primers used for *O. brachyantha* were designed de novo. Total mRNA was isolated from leaf and root tissues sampled from *O. sativa*, *O. glaberrima*, and *O. brachyantha* plants grown in the Biotron facilities at the University of Wisconsin-Madison under normal rice growth conditions. First-strand cDNA synthesis was completed using the SuperScript III first-strand synthesis system for RT-PCR (Invitrogen). One mRNA sample from each species was taken through an amended cDNA synthesis procedure in which the addition of the reverse transcriptase enzyme was omitted. These samples represent the negative controls used in subsequent PCR to confirm the absence of contaminating genomic DNA. PCR from the synthesized cDNAs was performed under the following cycling conditions: 95°C for 5 min followed by 33 cycles of 95°C for 30 s, 60°C for 30 s and 72°C for 30 s, ending with a final 4-min extension at 72°C.

Sequence Divergence Analysis and Estimation of Mutation Rate

Genomic, CDS, and protein sequences of orthologous *Cen8* genes from the three *Oryza* species were compared to reveal the structure, origin, and evolution of the orthologous genes. Ka/Ks ratios (ω) using the maximum likelihood algorithm were computed using PAML (Yang, 2007). The significance of ω that deviated from neutrality ($\omega = 1.0$) was tested using LRT. Protein sequences of homologous gene pairs were aligned using MUSCLE (Edgar, 2004), and codon-based DNA sequences were aligned using the aligned protein sequences as guides with the Pal2nal script (Suyama et al., 2006). Codeml with fixed ($\omega = 1$) and free omega ($\omega = \text{estimated}$) models was used to test whether any homologous gene pairs were under selective constraint (Yang, 2007). We further calculated the mutation rate in coding regions (/base pair/MY) using the total number of substitutions (synonymous and nonsynonymous) divided by the product of CDS length (bp) and double species divergent time (Gillespie, 2004), 1 MY for *O. sativa* and *O. glaberrima* and 10 MY for *O. sativa* and *O. brachyantha*, and *O. glaberrima* and *O. brachyantha* (Ammiraju et al., 2008, 2010).

Population Polymorphism Analysis of *O. glaberrima* and *O. brachyantha* *Cen8* Genes

SNP data of *O. glaberrima* *Cen8* genes were generated from eight accessions (see Supplemental Table 14 online) using Illumina Solexa

resequencing (R.A. Wing, Y. Yu, and C. Fan, unpublished data). By mapping the resequencing reads to the *O. glaberrima* *Cen8* pseudomolecule, we obtained sequence polymorphisms for *O. glaberrima* *Cen8* genes and intergenic regions. The number of SNPs and the rate of SNPs per kilobase was further calculated and compared. For *O. brachyantha*, one *O. brachyantha* *Cen8* gene, *OB_t00833* (3.9 kb) and one gene fragment *OB_t01153* (partial sequence is 1022 bp), were PCR amplified and sequenced from 15 *O. brachyantha* accessions collected from several African countries (see Supplemental Table 15 online). *OB_t00833* is located immediately adjacent to the centromeric satellite domain (CentO in *O. sativa* and *O. glaberrima*, and CentO-F in *O. brachyantha*; Figure 1), and *OB_t01153* resides in the left boundary of the satellite domain, where crossover recombination is presumed to be totally suppressed. Sequence alignment of population polymorphism data allowed us to calculate several population genetic parameters, including polymorphism frequency spectra, and Tajima's D (Tajima, 1989) as implemented in DnaSPv5 (Librado and Rozas, 2009) using nucleotide diversity (θ_{π}) and Watterson's sequence variation (θ_w) (Watterson, 1975). Assessment of significant deviation from neutrality was simulated using a coalescence approach. Since *OB_t01153* has fewer polymorphic sites, it cannot be used for statistical analysis if we analyzed it individually; therefore, we pooled the two sequence data sets together to perform population genetic and recombination analyses. Using the combined polymorphism data, we further performed three powerful recombination detection methods (Piganeau et al., 2004), which are the most efficient and sensitive methods for detecting recombination and gene conversion events: (1) *maxichi*, maximum χ^2 recombination test using Maynard Smith's method (Smith, 1992); (2) *LDr2*, tests the correlation between the measure of linkage disequilibrium, r^2 (Hill and Robertson, 1966), and the distance between sites; and (3) *geneconv*, detects gene conversion events using Sawyer's method (Sawyer, 1989).

Pfam Analysis of *Cen8* Genes

Protein sequences of seven conserved *Cen8* genes were used to find matching protein family at <http://pfam.sanger.ac.uk/>. The search was performed using HMM (hidden Markov model) model and E-value of 1.0 as threshold.

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: AC240787, AC237093, AC223444, AC240789, AC237092, AC223443, AC240788, AC223442, AC240786, AC223445, AC237091, AC223441, AC223438, AC240777, AC223440, AC249775, AC223439, AC240778, AC237085, AC237086, and AC240776.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Boxplot of Ks and Ka Values between *O. sativa* and *O. glaberrima*.

Supplemental Figure 2. Boxplot of Ks and Ka Values between *O. sativa* and *O. brachyantha*.

Supplemental Figure 3. Boxplot of Ks and Ka Values between *O. glaberrima* and *O. brachyantha*.

Supplemental Figure 4. Gene Structure and Polymorphism Distribution in One *O. brachyantha* *Cen8* Gene (*OB_t0083*) and One Gene Fragment (*OB_t01153*).

Supplemental Table 1. List of *O. sativa* *Cen8* Genes.

Supplemental Table 2. Primers Used for RT-PCR.

Supplemental Table 3. Expression Profile of Seven Active *O. brachyantha* *Cen8* Genes.

Supplemental Table 4. Expression of *O. glaberrima* *Cen8* Genes.

Supplemental Table 5. ChIP Analysis.

Supplemental Table 6. Summary of *O. glaberrima* SNP Data.

Supplemental Table 7. Statistical Test of Polymorphism Substitution Pattern in *OB_t00833*.

Supplemental Table 8. Statistical Test of Polymorphism Distribution in *O. brachyantha* *Cen8* Genes.

Supplemental Table 9. Maximum χ^2 Recombination Test.

Supplemental Table 10. Relationship between LD and Distance (*LDr2* Test).

Supplemental Table 11. Sawyer's Gene Conversion Detection (*Geneconv* Test).

Supplemental Table 12. Pfam Search Results for Seven Conserved *Cen8* Genes.

Supplemental Table 13. Summary of BACs and GenBank Accessions Used for *Cen8* Sequencing in *O. glaberrima* and *O. brachyantha*.

Supplemental Table 14. Sampling of *O. glaberrima* Accessions.

Supplemental Table 15. *O. brachyantha* Population Accession Sampling.

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AUTHOR CONTRIBUTIONS

C.F., J.J., and R.A.W. designed the research. C.F. and J.G.W. performed the research. C.F., J.G.W., J.Z., and C.D.H. analyzed data. C.F., J.G.W., C.D.H., J.J., and R.A.W. wrote the article. J.J. and R.A.W. are joint senior authors who contributed equally.

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Conservation and Purifying Selection of Transcribed Genes Located in a Rice Centromere
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