High Occurrence of Functional New Chimeric Genes in Survey of Rice Chromosome 3 Short Arm Genome Sequences

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High Occurrence of Functional New Chimeric Genes in Survey of Rice Chromosome 3 Short Arm Genome Sequences

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Abstract

In an effort to identify newly evolved genes in rice, we searched the genomes of Asian-cultivated rice Oryza sativa ssp. japonica and its wild progenitors, looking for lineage-specific genes. Using genome pairwise comparison of approximately 20-Mb DNA sequences from the chromosome 3 short arm (Chr3s) in six rice species, O. sativa, O. nivara, O. rufipogon, O. glaberrima, O. barthii, and O. punctata, combined with synonymous substitution rate tests and other evidence, we were able to identify potential recently duplicated genes, which evolved within the last 1 Myr. We identified 28 functional O. sativa genes, which likely originated after O. sativa diverged from O. glaberrima. These genes account for around 1% (28/3,176) of all annotated genes on O. sativa’s Chr3s. Among the 28 new genes, two recently duplicated segments contained eight genes. Fourteen of the 28 new genes consist of chimeric gene structure derived from one or multiple parental genes and flanking targeting sequences. Although the majority of these 28 new genes were formed by single or segmental DNA-based gene duplication and recombination, we found two genes that were likely originated partially through exon shuffling. Sequence divergence tests between new genes and their putative progenitors indicated that new genes were most likely evolving under natural selection. We showed all 28 new genes appeared to be functional, as suggested by KS/Ks analysis and the presence of RNA-seq, cDNA, expressed sequence tag, massively parallel signature sequencing, and/or small RNA data. The high rate of new gene origination and of chimeric gene formation in rice may demonstrate rice’s broad diversification, domestication, its environmental adaptation, and the role of new genes in rice speciation.

Key words: chimera, comparative genomics, gene duplication, new gene, Oryza.

Introduction

The genetic fundamental of organismal biodiversity is considerably relied on origination of new genetic elements. Myriad examples have provided evidence supporting newly evolved gene involvement in adaptive changes (Long and Langley 1993; Zhang et al. 2002; Long et al. 2003; Jones et al. 2005; Des Marais and Rausher 2008; Fan, Emerson, et al. 2008; Zhou et al. 2008; Heinen et al. 2009; Parker et al. 2009; Chen et al. 2010; Ding et al. 2010; Potrzebowski et al. 2010; Charrier et al. 2012; Yeh et al. 2012). Understanding the molecular mechanisms involved in the formation of new genes is progressing rapidly, although many details of these mechanisms and their interactions await further investigation. As reviewed previously (Long et al. 2003; Kaessmann et al. 2009; Cardoso-Moreira and Long 2012; Ranz and Parsch 2012), the major mechanisms of new gene origination include but not limited to tandem gene duplication, exon shuffling, retroposition, mobile elements, horizontal gene transfer, gene fusion/fission, de novo origination, or a combination of two or more of the mechanisms (Wang et al. 2000; Bachtrog and Charlesworth 2003; Jones and Begun 2005). Systematical comparative genomic analysis using Drosophila genomes revealed that DNA-based gene duplication and retroposition played major roles in the formation of new genes (Yang et al. 2008; Zhou et al. 2008). Because of...
the limitation of genome sequence data and genetic resources, we do not know yet about the prospect of new gene formation in the plant kingdom as much as in animals, though a few recent studies have demonstrated that many similarities exist between plants and animals (Zhang et al. 2005; Wang et al. 2006; Fan et al. 2008; Zhu et al. 2009; Sakai et al. 2011).

To understand the molecular processes and mechanisms governing the evolution of new genes and their functions, we must search for genes that originated recently and study their origination patterns and functions. The methods of detecting new genes have evolved dramatically with the advancement of experimental and computational technology and massive DNA sequence data generated in both model and nonmodel organisms. Early discoveries of new genes were largely based on the detection of a single gene by chance. Phylogenetic comparisons of genetic signals (e.g., fluorescence in situ hybridization and genomic southern blotting) have also been used as an efficient and reliable way to identify new protein-coding genes in Drosophila and mammals at a larger scale (Betran et al. 2002; Wang et al. 2004; Marques et al. 2005). This was not the case in plants, due to technical challenges, for example, difficulty of cytogenetic analysis for plant chromosomes and low efficiency and high false-positive rate of genomic southern blotting analysis to identify gene duplication events. In plants, previous works have also provided a tool that used array-based comparative genomic hybridization to identify potential new genes in the closely related Arabidopsis species (Fan et al. 2007). However, the most effective technique for finding duplications and further identifying new genes would be a genomic sequence comparison based on the availability of genome sequences. Similar efforts have been applied in the analysis of several other genomes and yielded a fair amount of information contributing to our understanding of the evolution of genes and genomes (Stein et al. 2003; Chimpanzee Sequencing and Analysis Consortium 2005; Marques et al. 2005; Yu et al. 2005; Clark et al. 2007; Jun et al. 2009; Liti et al. 2009; Marques-Bonet and Eichler 2009; Marques-Bonet et al. 2009; Green et al. 2010; Jensen and Bathtro 2010; Gan et al. 2011; Hu et al. 2011; Kim et al. 2011; Locke et al. 2011; Zhang et al. 2011; Scally et al. 2012). Moreover, comparing closely related species, as demonstrated in the Drosophila melanogaster subgroup (Yang et al. 2008; Zhou et al. 2008), provided more powerful strategy for identifying gene duplication events across the entire genome and for revealing the extent and pattern of new gene origins.

As part of an international effort to characterize the functions of all rice genes (Zhang et al. 2008), sequences of chromosome 3 short arm (Chr3s) using the bacterial artificial chromosome (BAC)-based physical maps to select minimum tilling paths of BAC clones in most Oryza species have been finished and are publically available. Therefore, these genomic sequence data provide an opportunity to decipher gene and genome evolution at the phylogenetic level within a single genus using comparative genomics approaches. The genus Oryza is composed of 23 species that diverged over a relatively short time period approximately 15–20 Ma with broad diversification and largely solved phylogenetics (Ge et al. 1999; Zhu and Ge 2005; Ammiraju et al. 2008; Tang et al. 2010). Oryza sativa ssp. japonica and O. glaberrima are Asian- and African-cultivated rice species, respectively. Phylogenetically, O. sativa ssp. japonica and O. glaberrima belong to the AA genome type in the genus Oryza, which diverged roughly from 0.5 to 1 Ma (Ammiraju et al. 2008; Tang et al. 2010). Species of O. punctata belongs to the BB genome type and is used as outgroup of the AA genome Oryza species for phylogenetic analysis. AA and BB genome type species diverged at around 2–5 Ma (fig. 1) (Ammiraju et al. 2008; Tang et al. 2010). Through the genome sequence comparisons between Asian rice species (including O. sativa, O. nivara, and O. rufipogon) and African rice species (including O. glaberrima, O. barthii, and O. punctata), this study aimed to identify Chr3s potential new genes, which recently originated in O. sativa and/or its wild species progenitors, O. nivara and O. rufipogon.

Materials and Methods

Searching O. sativa ssp. japonica-Specific New Genes by Comparative Genome Analysis

Sequence data of Chr3s in O. glaberrima, O. punctata, and O. barthii, O. nivara and O. rufipogon were downloaded from Gramene (http://www.gramene.org/). Chr3s sequences of O. sativa ssp. indica were downloaded from 2003/10/7 BGI version (ftp://ftp.genomics.org.cn/pub/ricedb/rice_update_data/genome/9311). The whole-genome sequences of O. glaberrima were downloaded from http://www.iplantcollaborative.org/. We performed genome pairwise comparisons between O. sativa ssp. japonica Chr3s coding sequences (CDSs) and other five species Chr3s genome sequences. The annotation and CDSs of O. sativa ssp. japonica were downloaded from Michigan State University (MSU) Rice Genome Annotation Project (RGAP, MSU V7) (http://rice.plantbiology.msu.edu/downloads.shtml). To search for the O. sativa-specific new genes, the first step was to identify the Chr3s orthologous genes among six species. We used two criteria to define the orthologous genes. First, we conducted a BLAT (Kent 2002) search for Chr3s orthologous genes by aligning genome sequences of O. glaberrima, O. sativa ssp. indica, O. barthii, O. punctata, O. nivara, and O. rufipogon against the CDSs of O. sativa ssp. japonica. We had two requirements: the alignment of the orthologous sequence needed to cover over 95% of the length of the O. sativa ssp. japonica CDSs and must be located in the synteny region of all the genomes. Whether an O. sativa ssp. japonica gene was considered in the synteny region was defined by the presence of at least two flanking genes in the 30-kb DNA fragment.
containing the gene hit in other genomes. Second, the orthologous sequences were defined as two sequences with reciprocal best hits of each other. We conducted the reciprocal searches using BLAT and defined a pair of sequences from two genomes having the best hit against each other as “reciprocal” best hits. We descendingly sorted the hits according to the BLAT alignment score and then BLAT identity score (http://genome.ucsc.edu/FAQ/FAQblat.html#blat4 for methods to compute these two scores). We then defined the ones ranking in the first as the “best” hits. After we identified the orthologous genes, we filtered them out, and picked the remaining annotated genes, which are only present in O. sativa ssp. japonica and/or the other three Asian rice species (O. sativa ssp. indica, O. rufipogon, O. nivara) but are absent in all the African rice species O. glaberrima, O. barthii, and O. punctata (fig. 1). We further BLAT CDSs of O. sativa ssp. japonica-specific genes to the entire O. glaberrima genome and identified their homologous regions in O. glaberrima. The results were then BLAT back to all CDSs of O. sativa ssp. japonica. We only selected O. sativa ssp. japonica genes, which did not have reciprocal BLAT best hits in O. glaberrima genome as O. sativa ssp. japonica new gene candidates. These genes likely originated after the divergence between Asian rice species and African rice species about 1 Ma. We further estimated the average rates of synonymous substitution ($K_s$) using gKaKs pipeline with Yn00 method for all Chr3s orthologous genes earlier identified between O. sativa ssp. japonica and O. glaberrima (Zhang et al. 2013).

To determine the origination pattern of these recently evolved new genes in O. sativa ssp. japonica, we searched for their paralogs in the O. sativa ssp. japonica genome. To identify paralogous gene pairs, we BLAT the CDSs of the candidate genes against all the CDSs of O. sativa ssp. japonica with the match length of the paralogous gene pair more than 100 bp and mismatch length/(mismatch length + match length) less than 0.1. We picked up only the paralogous gene pairs with $K_s$ less than 0.0192, which is the average $K_s$ of the orthologous gene pairs between O. sativa ssp. japonica and O. glaberrima corresponding to 1 Myr divergence time. We further removed the genes with “retrotransposon protein” and “transposon protein” terminology in their annotations to define the list of O. sativa ssp. japonica new gene candidates. Next, to test whether these O. sativa lineage-specific new genes were ancient duplicate genes that lost in African Oryza species, we applied reciprocal BLASTP searches to identify whether these new gene candidates contain orthologous copies in other distantly related species. We BLASTP protein sequences of these new gene candidates to all proteins in Uniprot (http://www.uniprot.org/), which includes SwissProt and TrEMBL data. If a new gene candidate had hits in other species, we BLASTP these hits back to all O. sativa ssp. japonica proteins (http://www.gramene.org/Multi/blastview). If this best hit from BLASTP search was the new gene, we deleted this new gene candidate. We also used RepeatMasker (RepeatMasker libraries version: rm-20120418) to scan the transposons existing in CDSs of new gene candidates.

**Sequence Divergence and Phylogenetic Analysis**

We calculated the ratio of nonsynonymous substitution and synonymous substitution rates ($K_a/K_s$ donated as “$\omega$”) using maximum likelihood algorithm (codeml) implemented in the PAML package (Yang 2007). The significance of $\omega$ that deviated from neutrality ($\omega = 1$) was tested using the likelihood ratio test (LRT). We aligned the sequences of paralogous/orthologous gene pairs using bl2seq (Altschul et al. 1997). We used codeml to calculate the $\omega$ value between the two
sequences (Yang and Nielsen 2000). We then used codeml with two models (α fixed at 1 and α varying freely) to test whether any of the identified new genes were statistically under natural selection (Yang 2007). Phylogenetic analysis of the gene tree was performed using Neighbor Joining algorithm implemented in PAUP (Swofford 2002). The CDSs of the gene family were aligned using ClustalW (Larkin et al. 2007). The bootstrap analysis with 1,000 replicates was used to assess the robustness of the branches.

To address whether α < 1 is due to that the parental gene is under strong purifying selection and the new gene is a pseudogene evolving neutrally, we applied PAML branch model to calculate α values for the branch leading to new genes. We first downloaded the recently completed whole-genome sequences of O. glaberrima, O. barthii, and O. punctata from http://www.iplantcollaborative.org. We identified the orthologous sequences of parental genes from the three outgroup species using ortholog search approach described earlier. We aligned only homologous region for all sequences using MAFFT (Katoh et al. 2005) and Perl scripts. We estimated α for the foreground branch leading to the O. sativa ssp. japonica lineagespecific new gene and for background branches leading to the parental genes and their orthologous genes in outgroup species (O. glaberrima, O. barthii, and O. punctata). We used a two-ratio model allowing different α in foreground and background branches with PAML codeml. The significant level of foreground branch α was tested using LRT compared with the null hypothesis of a model where foreground α fixed to 1 and background α varied freely (Yang 2007).

Expression Analysis

The expression of identified new genes was determined by the presence of full-length cDNA (FL-cDNA), expressed sequence tag (EST; Pontius et al. 2003), RNA sequencing transcriptome data (RNA-seq) (He et al. 2010; Zemach et al. 2010; Davidson et al. 2012), massively parallel signature sequencing (MPSS) (Nakano et al. 2006), and small RNA sequencing signatures (Nobuta et al. 2007). RNA-seq data, which were processed by RGAP, were downloaded from http://rice.plantbiology.msu.edu/expression.shtml. The transcription abundance was reported in fragments/kilobase of transcript/million fragments mapped (FPKM) across 11 libraries including leaves—20 days, postemergence inflorescence, pre-emergence inflorescence, anther, pistil, seed-5 DAP, embryo-25 DAP, endosperm-25 DAP, seed-10 DAP, shoots, and seedling four-leaf stage (supplementary table S1, Supplementary Material online). The significant level of foreground branch α was tested using LRT compared with the null hypothesis of a model where foreground α fixed to 1 and background α varied freely (Yang 2007).

Identification of New Chimeric Genes

After we compared the new genes with their paralogs, we detected that many new genes have formed chimerical gene structures with flanking sequences or other gene sequences. If the flanking or other gene sequences that a new gene recruited in the CDS are larger than 30 bp, we considered it as a new chimeric gene. To identify whether a new chimeric genes has transcription evidence for the chimerical CDS structure, we mapped EST, full-length cDNA, and RNA-seq sequences to the junctions of chimeras. We obtained RNA-seq raw data from NCBI Sequence Read Archive (SRA: SRR352184.sra, SRR352187.sra, SRR352189.sra, SRR352190.sra, SRR352192.sra, SRR352194.sra, SRR352204.sra, SRR352206.sra, SRR352207.sra, SRR352209.sra, SRR352211.sra, SRR042529.sra, SRR034580.sra, SRR034581.sra, SRR034582.sra, SRR034583.sra) from http://sra.dnanexus.com/dispatch_many. We preprocessed the RNA-seq data with quality control using trim_galore (Version 0.2.5) (http://www.bioinformaticsbabraham.ac.uk/projects/trim_galore/) before mapping.
We removed duplications existing in aligned reads due to polymerase chain reaction using picard-tools-1.79 (http://picard.sourceforge.net/) after mapping. Given the length of the RNA-seq reads ranging from 35 to 40 bases, we extracted 32-bp DNA sequences of upstream and downstream flanking regions at the breakpoints of a chimeric gene. We then mapped the RNA-seq reads to the extracted flanking DNA sequences with Tophat v2.0.7 (Trapnell et al. 2009). Finally, we checked whether any RNA-seq reads aligned on these flanking sequences and crossed the chimerical breakpoints. We applied similar approach to map the EST sequence data to the extracted chimera breakpoint flanking DNA sequences with BLAT. We also checked whether these chimeric genes have FL-cDNA through browsing http://rice.plantbiology.msu.edu/cgi-bin/gbrowse/rice/.

Results

Identification of Potential New Gene Candidates for O. sativa ssp. japonica

Three steps were carried out to detect the potential new genes that recently originated in O. sativa ssp. japonica and its wild progenitors. First, the comparative genomic analysis of Chr3s pseudomolecules among six species identified 862 annotated genes only present in O. sativa ssp. japonica and/or its progenitors. Second, from the 862 gene candidates, we filtered out the gene candidates, which had reciprocal best hits in the O. glaberrima whole-genome sequence. This yielded 753 O. sativa ssp. japonica-specific gene candidates. Third, we BLAT these 753 candidates to all the CDSs of O. sativa ssp. japonica to find the best-hit paralogs and then calculated the Ks between O. sativa ssp. japonica-specific gene and its paralog. On the basis of the average Ks = 0.0192 of 1,797 Chr3s orthologous genes between O. sativa and O. glaberrima, we inferred that the paralogous pairs with Ks less than 0.0192 were potential new genes that likely originated after the divergence of O. sativa and O. punctata (~2–5 Ma). Both LOC_Os03g24980 and LOC_Os03g24990 that originated after the divergence of O. sativa and O. glaberrima (~0.5–1 Ma) appeared to be chimeric. LOC_Os03g24990 was possibly generated by DNA-level recombination of LOC_Os03g24960 and its target flanking sequence. LOC_Os03g24980 recruited exons of LOC_Os03g24970 and local sequences as its intron (supplementary fig. S2W–X, Supplementary Material online).

For the remaining 23 new genes, 21 were apparently generated through the single-gene DNA level recombination–mechanism gene duplication (supplementary fig. S2, Supplementary Material online). Comparing gene DNA sequences and exon–intron structure between new genes and parental genes, we observed four general patterns of DNA-based recombination and duplications for new gene origination in O. sativa Chr3s: 1) the new gene recruited partial parental gene sequences to form a new chimerical gene structure (fig. 2A), for example, LOC_Os03g04190, LOC_Os03g04230, LOC_Os03g04270, LOC_Os03g09130, LOC_Os03g11860, LOC_Os03g15110, LOC_Os03g18650, LOC_Os03g21310, LOC_Os03g25950, and LOC_Os03g29140. 2) The new gene recruited partial parental gene sequences formed an intact nonchimeric gene (fig. 2B), for example, LOC_Os03g02130, LOC_Os03g03050, LOC_Os03g04760, LOC_Os03g07690, LOC_Os03g15060, and LOC_Os03g24630. 3) The new gene adopted the entire parental gene sequences and both genes shared the same exon–intron gene structure (fig. 2C), for example, LOC_Os03g07090,LOC_Os03g32526, and LOC_Os03g33920. 4) The new gene recruited the entire parental gene sequences but formed a different exon–intron gene structure (fig. 2D), for example, LOC_Os03g12480 and LOC_Os03g16320.

The origination patterns of these new genes were revealed by the location, gene structure, and sequences comparison between new genes and their paralogous progenitors in O. sativa ssp. japonica. A 30-kb telomeric region containing three functional new genes was generated through a segmental duplication of an unmapped annotated region in O. sativa genome (supplementary figs. S1 and S2A–C, Supplementary Material online). Four adjacent annotated genes, LOC_Os03g24960, LOC_Os03g24970, LOC_Os03g24980, and LOC_Os03g24990, are located in the middle of Chr3s within a13-kb fragment, which is unique to AA genome rice species. By identifying the paralogs of these four genes, we concluded that these genes originated through segmental gene duplication followed by tandem duplication. LOC_Os04g30860 and LOC_Os04g30870 appeared to be the most closely related parental genes given their structure, sequence similarity, and phylogenetic analysis (supplementary figs. S3 and S4, Supplementary Material online).

A partial segment of the region between these two genes was involved in a segmental duplication, which possibly gave rise to LOC_Os03g24960 and LOC_Os03g24970 after the divergence of O. sativa and O. punctata (~2–5 Ma). Both LOC_Os03g24980 and LOC_Os03g24990 that originated after the divergence of O. sativa and O. glaberrima (~0.5–1 Ma) appeared to be chimeric. LOC_Os03g24990 was possibly generated by DNA-level recombination of LOC_Os03g24960 and its target flanking sequence. LOC_Os03g24980 recruited exons of LOC_Os03g24970 and local sequences as its intron (supplementary fig. S2W–X, Supplementary Material online).
LOC_Os03g12580 was formed from shuffling the first exon of LOC_Os06g01010 and its flanking sequences (supplementary fig. S2P, Supplementary Material online).

Chimeric gene formation appears to be very common in new rice genes. Among 28 O. sativa new genes that we observed, 14 new genes are chimerical. The chimerical CDS structure of a new gene is mostly formed by recruiting entire or partial parental gene sequences and DNA sequences from the insertion site (fig. 3A). However, we did find one new gene, LOC_Os03g09130, which was developed from two genes and an insertion of a DNA fragment (fig. 3B). We further examined the transcription of chimerical CDS structure using the expression data. Using RNA-seq data, we found eight chimeric genes that contain RNA-seq reads covering all the breakpoints and three chimeric genes that have RNA-seq reads covering some breakpoints. Using EST data, we identified three chimeric genes that have EST sequences covering all the breakpoints and one chimeric gene that has EST sequence covering some breakpoints. Furthermore, five chimeric genes have FL-cDNA (supplementary table S5, Supplementary Material online). In summary, the chimerical CDS structure for all 14 chimeric genes was confirmed by RNA-seq, EST, and/or FL-cDNA sequence data.

Evolution Pattern of O. sativa ssp. japonica New Genes

We calculated $\omega$ values to gain insight into the evolution of O. sativa ssp. japonica new genes (supplementary table S6, Supplementary Material online). Because all new genes originated and evolved very recently (<1 Ma), we observed very low number and rates of both synonymous and nonsynonymous substitution (supplementary table S6, Supplementary Material online). Nineteen of the 28 paralogous pairs showed no synonymous substitution and/or nonsynonymous substitution. For the remaining nine paralogs, four of them had $\omega$ values less than neutrality, and five had $\omega$ values greater than 1 (supplementary table S6, Supplementary Material online). Furthermore, LRTs for the sequence divergence of the majority of 32 paralogous genes did not show significant deviation from neutrality. This was likely due to the recent

Table 1
The New Genes, Paralogs, and Creation Mechanisms

<table>
<thead>
<tr>
<th>New Gene</th>
<th>Annotation</th>
<th>Paralogs</th>
<th>Possible Formation Mechanisms</th>
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<tbody>
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<td>Os03g01008</td>
<td>Expressed protein</td>
<td>ChrSy.fgenesh.mRNA.80</td>
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<td>Gene duplication</td>
</tr>
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<td>Conserved hypothetical protein</td>
<td>Os06g36630</td>
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</table>
gene duplication that has not yet accumulated enough substitutions to give adequate statistical power.

Based on branch-specific $\omega$ analysis, six new genes have branch-specific $\omega$ less than 0.5. One new gene has branch-specific $\omega$ ranging between 0.5 and 1. Nine new genes have branch-specific $\omega$ more than 1 ranging from 1.92140 to 999.000. Moreover, LRTs showed that four new genes (LOC_Os03g12480, LOC_Os03g21310, LOC_Os03g24990, and LOC_Os03g32526) have branch-specific $\omega$ significantly smaller than 1 (supplementary table S7, Supplementary Material online).

Expression of New Genes in *O. sativa* ssp. *japonica*

All 28 new *O. sativa* genes appeared to be transcribed, as evidenced by the presence of RNA-seq, EST and/or FL-cDNA sequence, and/or small RNA/MPSS sequencing signature (table 2). Sixteen of the 28 new genes had at least two evidences of expression (table 2). Three genes, LOC_Os03g01014, LOC_Os03g01490, and LOC_Os03g07270, had high mRNA enrichment in RNA-seq data (supplementary table S1, Supplementary Material online). Among them, the expression of the two genes including LOC_Os03g01014 and LOC_Os03g07270 was enriched in different tissues: LOC_Os03g01014 was highly expressed in leaves. LOC_Os03g07270 was mainly transcribed in preinflorescence, pistil, seed, and embryo (supplementary table S1, Supplementary Material online). Accumulation of mRNA from two genes (LOC_Os03g01020 and LOC_Os03g01490) appeared to be fairly high in vivo, as revealed by the presence of 9 and 40 independent EST sequences in GenBank, respectively (supplementary table S2, Supplementary Material online).
online). Two genes, \textit{LOC\textsubscript{03}g01020} and \textit{LOC\textsubscript{03}g01490}, expressed substantial enrichments in MPSS (supplementary table S3, Supplementary Material online). Eight genes, \textit{LOC\textsubscript{03}g11860}, \textit{LOC\textsubscript{03}g29140}, \textit{LOC\textsubscript{03}g12850}, \textit{LOC\textsubscript{03}g25950}, \textit{LOC\textsubscript{03}g02340}, \textit{LOC\textsubscript{03}g02130}, \textit{LOC\textsubscript{03}g24630}, and \textit{LOC\textsubscript{03}g24980}, appeared to be enriched in small RNA sequencing signatures (supplementary table S4, Supplementary Material online). Moreover, these eight genes showed transcription of small RNA signatures in different tissues and developmental stages (supplementary table S4, Supplementary Material online). To compare the general pattern of small RNA expression signatures between new genes and regular functional genes, we randomly picked up 500 functional genes and found that 82.2% of the 500 genes show small RNA expression signature, thus the small RNA signature was higher in regular functional genes than in the new genes.

**Discussion**

**High Rate of New Gene Origination in Rice Genome**

*Oryza sativa* ssp. *japonica* Chr3s contains approximately 3,100 annotated CDSs including hypothetical and transposable element (TE)-related genes. In our effort to systematically search for potential new genes, which recently evolved in *O. sativa* ssp. *japonica*, we were able to identify 28 new genes, which account for 1% of total genes on Chr3s. However, it is likely that we underestimated or possibly overestimated the true number of new genes in *O. sativa* ssp. *japonica*. These values may be underestimates of the true number of new genes considering two reasons. First, we filtered out all TE-related genes (“retrotransposon protein” and “transposon protein”) after the unique *O. sativa* ssp. *japonica* genes were found. Second, we used the average \( K_s \) value of orthologous genes between *O. sativa* ssp. *japonica* and *O. glaberrima* as a cutoff value to define the age of the paralogous duplication event. It is likely that some new genes evolved quickly and that the substitution rate may be elevated. These criteria could possibly ignore some new genes based on their high synonymous substitution rate. Meanwhile, the number of new genes that we identified might be overestimates of the true number of new genes given two possibilities. First, although *O. sativa* ssp. *japonica* new genes do not have orthologs in *O. glaberrima*, it is possible to have orthologs present outside of Chr3s in other rice species due to chromosomal rearrangement (e.g., segmental duplication and transposition). Second, the low \( K_s \) values, which can be resulted from gene conversion and locally reduced mutation rate, may not truly reflect the age of duplications. Therefore, considering both situations, we estimated that *O. sativa* ssp. *japonica*-specific new genes would account for 0.8–2% of total annotated genes in the entire rice genome. RGAP annotated a total of 56,797 genes including putative, expressed, hypothetical, and TE-related genes (http://rice.plantbiology.msu.edu/riceInfo/info.shtml#Genes). Therefore, we deduced that the rice genome (a total of ~57,000 genes) might have 500–1,000 new genes (0.0088–0.017/gene/Myr), which evolved around 1 Ma after *O. sativa* ssp. *japonica* split from *O. glaberrima*. This new gene origination rate (per gene per Myr) in rice
and could drive the rapid occurrence of new genes in rice genome. Though genus Oryza stands as a small group in the plant kingdom containing only 23 species, the diversity and ecological adaptability of rice, which is found in a wide range of habitats from forest, savanna, and mountainsides to river and lakes, is remarkable and could drive the rapid occurrence of new genes in rice genome. (Wang et al. 2006). A previous study reported that cultivated rice (O. sativa ssp. indica) genome encodes 898 functional retroposed genes, of which 380 were predicted to have chimerical protein sequence structures (Wang et al. 2006). Because the most recent divergent time can better record the recent evolutionary events, our observation provided additional solid evidence for the high rate of new gene origination. Consistent with previous finding, we annotated a total of 28 new genes on O. sativa ssp. indica Chr3s, 14 (50%) of which appeared to be chimeric genes generated by segmental duplication and DNA-level recombination. Our current study revealed a high rate of chimeric gene origination as: 14 x 20 = 280 chimeric genes/Myr/genome. The higher rates of chimeric gene formation and the generation of a large number of functional genes in rice again demonstrated the broad diversification and adaptation of the grass species. Both our previous and current studies all demonstrated that rice genomes displayed an accelerated gene origination rate and generated a high number of chimeric gene structures that held potential to evolve novel functions (Wang et al. 2006; Fan et al. 2008). However, these findings are in contrast to the recently reported lower gene origination rate, which may result from extremely conservative genome annotation (Sakai et al. 2011). Conservative annotation is an approach that has been widely used in functional genomics and molecular functional analysis but may not fit the need for evolutionary genomic study. In practice, new evolutionary changes, including new genes, are seriously underestimated by this approach (Zhang et al. 2012).

Previous studies in Drosophila have demonstrated that repetitive elements could facilitate recombination to generate high occurrences of chimeric genes (Yang et al. 2008). In rice, the abundance of Pack-MULEs could capture fragment(s) of genomic DNA sequence while also rearranging and fusing with target sequence to generate a large amount of new reading frame and chimerical transcripts (Jiang et al. 2004). Therefore, mechanisms such as these could be responsible for the chimeric gene formation in rice genome.

### Table 2
Expression of New Genes in Oryza sativa

<table>
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<tr>
<th>Locus</th>
<th>RNA-Seq Data</th>
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Note: –+, present; –, absent.

**New Gene Originated as Chimera in Rice Genome**

Chimeric genes represent a class of genes that originated from multiple parental sources in coding and/or noncoding (regulatory site) sequences. Because of their unique origination, chimeric genes are unlikely to retain their parental characteristics and thus evolve novel functions. By surveying previous new genes detected in other organisms, it can be concluded that chimeric new genes account for a high percentage of total new genes identified in a variety of organisms ranging from mammals (Paulding et al. 2003; Sayah et al. 2004; Parker et al. 2009), to flies (Long and Langley 1993; Jones et al. 2005; Nozawa et al. 2005) and plants (Long et al. 1996; Wang et al. 2006; Fan et al. 2008). A recent investigation systematically searched through new genes using the Drosophila genome comparisons and found 30% of the new genes in the D. melanogaster species complex recruited various genomic sequences and formed chimeric gene structures. These findings suggest structure innovation is important to the generation of new genes (Zhou et al. 2008). This is similar to what was reported previously in the genomic analysis of O. sativa ssp. indica (Wang et al. 2006). A previous study reported that cultivated rice (O. sativa ssp. indica) genome encodes 898 functional retroposed genes, of which 380 were predicted to have chimerical protein sequence structures (Wang et al. 2006). Because the most recent divergent time can better record the recent evolutionary events, our observation provided additional solid evidence for the high rate of new gene origination. Consistent with previous finding, we annotated a total of 28 new genes on O. sativa ssp. indica Chr3s, 14 (50%) of which appeared to be chimeric genes generated by segmental duplication and DNA-level recombination. Our current study revealed a high rate of chimeric gene origination as: 14 x 20 = 280 chimeric genes/Myr/genome. The higher rates of chimeric gene formation and the generation of a large number of functional genes in rice again demonstrated the broad diversification and adaptation of the grass species. Both our previous and current studies all demonstrated that rice genomes displayed an accelerated gene origination rate and generated a high number of chimeric gene structures that held potential to evolve novel functions (Wang et al. 2006; Fan et al. 2008). However, these findings are in contrast to the recently reported lower gene origination rate, which may result from extremely conservative genome annotation (Sakai et al. 2011). Conservative annotation is an approach that has been widely used in functional genomics and molecular functional analysis but may not fit the need for evolutionary genomic study. In practice, new evolutionary changes, including new genes, are seriously underestimated by this approach (Zhang et al. 2012).
Supplementary Material
Supplementary tables S1–S7 and figures S1–S4 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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Literature Cited


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