

The Subtelomere of *Oryza sativa* Chromosome 3 Short Arm as a Hot Bed of New Gene Origination in Rice

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ABSTRACT Despite general observations of non-random genomic distribution of new genes, it is unclear whether or not new genes preferentially occur in certain genomic regions driven by related molecular mechanisms. Using 1.5 Mb of genomic sequences from short arms of chromosome 3 of *Oryza glaberrima* and *O. punctata*, we conducted a comparative genomic analysis with the reference *O. sativa* ssp. *japonica* genome. We identified a 60-kb segment located in the middle of the subtelomeric region of chromosome 3, which is unique to the species *O. sativa*. The region contained gene duplicates that occurred in Asian cultivated rice species that diverged from the ancestor of Asian and African cultivated rice one million years ago (MYA). For the 12 genes and one complete retrotransposon identified in this segment in *O. sativa* ssp. *japonica*, we searched for their parental genes. The high similarity between duplicated paralogs further supports the recent origination of these genes. We found that this segment was recently generated through multiple independent gene recombination and transposon insertion events. Among the 12 genes, we found that five had chimeric gene structures derived from multiple parental genes. Nine out of the 12 new genes seem to be functional, as suggested by Ka/Ks analysis and the presence of cDNA and/or MPSS data. Furthermore, for the eight transcribed genes, at least two genes could be classified as defense or stress response-related genes. Given these findings, and the fact that subtelomeres are associated with high rates of recombination and transcription, it is likely that subtelomeres may facilitate gene recombination and transposon insertions and serve as hot spots for new gene origination in rice genomes.

Key words: comparative genomics; gene duplication; *Oryza sativa*; subtelomere; new genes.

INTRODUCTION

Previous studies have provided evidence of the significant role that novel genetic elements have played in organismal diversification and speciation (Ohno, 1970; Long et al., 2003). Various mechanisms, such as retroposition, exon shuffling, tandem gene duplication, and transposon-mediated gene duplication, have been proposed for the creation of novel genetic elements in numerous organisms (see reviews by Long et al., 2003; Fan et al., 2007a). However, detailed analysis of gene duplication and novel gene evolution in plants is still lacking.

Comparative genomics is a powerful tool to search for gene duplication events across entire genomes, and has been applied in the analysis of several organisms (e.g. Betran et al., 2002; Marques et al., 2005; Zhang et al., 2005; Wang et al., 2006). Comparing closely related species is particularly powerful for the detection of recent gene duplications. For example, the recently released 12 wild *Drosophila* genomic sequences have provided excellent opportunities to decipher gene and

genome duplication at the phylogenetic level within a single genus *Drosophila* (*Drosophila* 12 Genomes Consortium, 2007). However, the few plant species for which genome sequences are available are distantly related, and thus a search for new genes has been hindered by insufficient resolution for short evolutionary time intervals.

The genus *Oryza*, which contains the world's most important food crop—rice (*O. sativa*)—is an ideal plant model system to study detailed gene and genome evolution, due to small

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genome size and the availability of genome sequences from both subspecies of cultivated rice, *japonica* and *indica* (IRGSP, 2005; Yu et al., 2002). Findings from rice research can also inform studies on other cereal crops, such as corn and wheat, which, despite sharing a common ancestor 50 MYA, have genome sizes of six to 38 times larger than rice, respectively. Thus, rice provides the central comparative genomics core for monocot research (Bennetzen, 2007; Paterson et al., 2005; Wing et al., 2005).

The genus *Oryza* is composed of 23 species that are classified into 10 distinct genome types (six diploid and four allotetraploid) (Ge et al., 1999), and the phylogenetic relationships among these genome types are well resolved (Ge et al., 1999; Zhu and Ge, 2005) and cover an approximate 17 million year time span. Such broad diversification over such a relatively short period of time indicates the potential for new gene creation is relatively high.

In 2004, we were funded to create a genus level comparative genomics system for the genus *Oryza* composed of 11 bacterial artificial chromosome (BAC)-based fingerprint/end sequence physical maps, representative of the 10 genome types, aligned to the rice RefSeq (Wing et al., 2005; Ammiraju et al., 2006; Kim et al., 2008). The *Oryza* Map Alignment Project (OMAP) system now provides immediate access to virtually any region of the collective *Oryza* genomes for detailed comparative investigation. As part of an international effort to functionally characterize all rice genes, we are focusing on a detailed analysis of the short arm of chromosome 3 and have used the BAC-based physical maps to select minimum tiling paths of BAC clones across the chromosome 3 short arms of *O. glaberrima* [AA], *O. punctata* [BB], *O. officinalis* [CC], and *O. minuta* [BBCC] for sequencing. As part of an initial pilot project to sequence these short arms, we sequenced and finished ~1.5-Mb BAC tiles from *O. glaberrima* and *O. punctata* and compared these sequences with the *O. sativa ssp. japonica* RefSeq. With such a genomicscale comparison, we were able to identify a recently evolved 60-kb DNA segment in *O. sativa* that contained a number of young genes that originated within the last one million years (Myr).

RESULTS

Comparative Analysis of a 1.5-Mb Region from the Short Arm of Chromosome 3 in *O. sativa ssp. japonica*, *O. glaberrima*, and *O. punctata*

A unique and contiguous 60-kb region from the subtelomeric region of the short arm of chromosome 3 of the Asian culti-

vated rice, *O. sativa ssp. japonica*, was identified by comparing with 1.5-Mb orthologous regions sequenced from the AA genome of African cultivated rice *O. glaberrima* and the BB genome of *O. punctata*. This unique region in *O. sativa* was found to contain 12 candidate genes and one complete 11-kb long *TyGypsy* LTR retrotransposon that was annotated as three independent retrotransposons in The Institute for Genomic Research (TIGR) gene ontology database (Figure 1 and Table 1). To determine if the 60-kb region was unique to *japonica* rice or could be found in its putative progenitor species, *O. nivara* and *O. rufipogon*, we searched BAC end sequence (BES) datasets for these species (Kim et al., 2008) for sequences similar to the 60-kb sequence using BLAST. This analysis identified orthologous BESs to *Os03g01410* in both *O. nivara* and *O. rufipogon*, and *Os03g01420* in *O. nivara* (Table 2), thus providing evidence that at least part of the unique 60-kb sequence could be found in these two wild species. This finding was further supported by analysis of the complete and partial orthologous sequences from *O. nivara* and *O. rufipogon*, respectively, which revealed the presence of both genes 2 and 5 (Yu et al., unpublished data). Since BES datasets and sequence data were not available for two additional AA genome species, *O. barthii* (the wild progenitor of *O. glaberrima*) and *O. glumaepatula* (a wild species from South America), we designed a pair of diagnostic PCR primers to detect the presence or absence of the *O. sativa ssp. japonica* unique 60-kb region at this location on chromosome 3 (Figure 1). The expected PCR amplification band size between primers *Os03g01360F* and *Os03g01530R* for *O. sativa*, as shown in Figure 1, is 65 Kb, which is too big for the regular PCR amplification. If this segment is missing, the expected size of the PCR band would be 2.5–3 kb, which is exactly what we detected using genomic DNA isolated from the *O. glaberrima* control and *O. barthii* and *O. glumaepatula* (Figure 2). We further constructed a synteny map using the chromosome 3 sequences to check whether this 60-kb region was located outside the syntenic chain based on the Chain and Net pipeline of UCSC. As expected, whether using the *O. glaberrima* or *O. punctata* genome as the reference sequence (data not shown), almost the entire region fell in the syntenic gap, which supports the hypothesis that most of the genes in this 60-kb region in *O. sativa* should be very young.

Thus, all data indicated that the origin of this region predates the divergence of the *O. sativa*–*O. rufipogon*–*O. nivara* clade but occurred after the divergence of this clade from the ancestral *O. glaberrima*, *O. barthii*, and *O. glumaepatula* clade

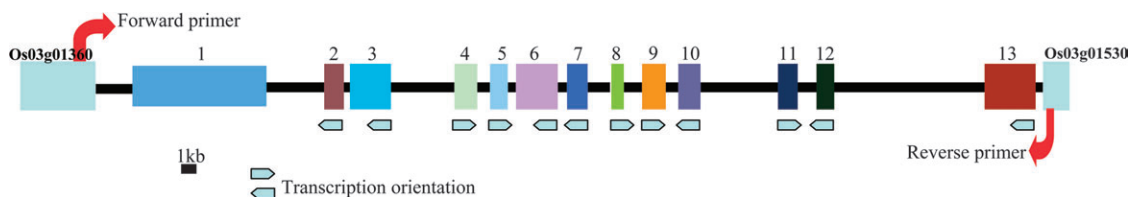


Figure 1. Schematic Sketch of 60-kb Segment in *O. sativa ssp. japonica*.

The primer pair was used to amplify this segment for other AA species. See Table 1 for gene order.

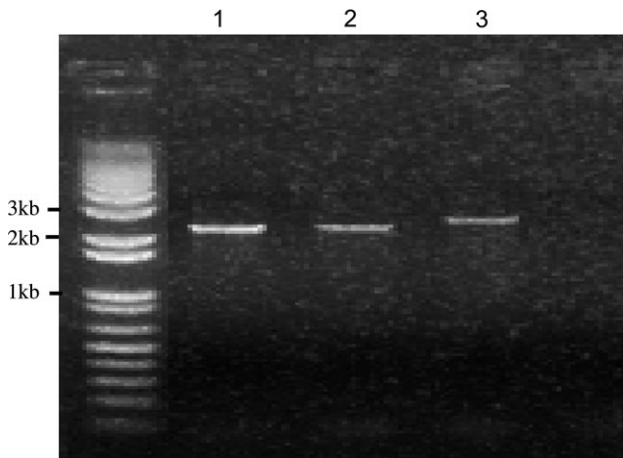
Table 1. The Detailed Information and Ka/Ks Analysis for 13 Annotated Genes.

Gene order	Gene ID (TIGR)	Paralogs (identity and alignment length)	Annotation	Expression (no. of EST)	Ka	Ks	Ka/Ks (LRT <i>p</i> -value)
1	<i>Os03g01380</i> <i>Os03g01390</i> <i>Os03g01400</i>	<i>Os01g17360</i> (97.2%/2328) <i>Os07g35630</i> (95.8%/988) <i>Os11g34420</i> (97.3%/1031)	<i>TyGypsy</i> retrotransposon	No (0)	NA	NA	NA
2	<i>Os03g01410</i>	<i>Os01g72700</i> + de nova (93.8%/1190)	STK kinase	No (0)	0.0250	0.1761	0.142/0.011
3	<i>Os03g01420</i> [#]	<i>Os03g01490</i> (100%/267)	Expression protein	FL-cDNA and EST transcript (65)	0	0	∞
4	<i>Os03g01430</i>	<i>Os03g01480</i> (100%/303)	Hypothetical protein	No (0)	0	0	∞
5	<i>Os03g01436</i>	<i>Os03g01470</i> (99.7%/1173)	Conserved hypothetical protein	FL-cDNA and EST transcript (1) (leaf of seedling 4 min gamma-irradiation, after 6 h)	NA	NA	NA
6	<i>Os03g01442</i> [#]	<i>Os01g69904</i> [#] (97.6%/2647)	Expression protein	FL-cDNA and EST transcript (12)	0.0277	0.0003	99/0.13
7	<i>Os03g01450</i> [*]	<i>Os04g32150</i> (93.6%/643) + <i>Os06g11900</i> (92.1%/478)	Hypothetical protein	EST transcript (3) (leaf of seedling 4 min gamma-irradiation, after 6 h)	NA	NA	NA
8	<i>Os03g01460</i>	No close homology	Hypothetical protein	No (0)	NA	NA	NA
9	<i>Os03g01470</i>	<i>Os03g01436</i>	Expression protein	FL-cDNA and EST transcript (6) (shoot and callus)	NA	NA	NA
10	<i>Os03g01480</i>	<i>Os03g01430</i>	Hypothetical protein	No (0)	0	0	∞
11	<i>Os03g01490</i> ^{*#}	<i>Os03g01420</i> ⁺	Expression protein	FL-cDNA and EST transcript (46)	0	0	∞
12	<i>Os03g01500</i> [*]	<i>Os12g24870</i> (94.0%/1050)	Hypothetical protein	EST transcript (2)	NA	NA	NA
13	<i>Os03g01520</i> ^{*#}	No close homology	Hypothetical protein	EST transcript (3)(2/2 callus)	NA	NA	NA

Notes: * Denotes mis-annotated gene models described in the TIGR rice annotation database. # Denotes genes with multiple alternative splicing isoforms. The cDNA from each parental gene and young gene were aligned with BLASTN using a low complexity filter off. Identity and alignment length were calculated using the chained SearchIO module of BioPerl. NA stands for non-feasible substitution analysis due to retrotransposon, no candidate parental gene, or distinct gene structure.

Table 2. BLAST Analysis Results Using the Unique *O. sativa* ssp. *japonica* 60-kb Segment as a Query against the *O. rufipogon* and *O. nivara* BES Datasets.

Species of BAC	BAC ID (location of BAC)	Identity (%)	Length of alignment (bp)	<i>O. sativa</i> orthologues
<i>O. nivara</i>	OR_BBa0096N24	99.21	629	<i>Os03g01420</i>
<i>O. nivara</i>	OR_BBa0036B16	99.83	590	<i>Os03g01420</i>
<i>O. nivara</i>	OR_BBa0122N16	97.79	768	<i>Os03g01420</i>
<i>O. nivara</i>	OR_BBa0059M20	98.94	661	<i>Os03g01410</i>
<i>O. rufipogon</i>	OR_CBa0020A22	98.97	677	<i>Os03g01410</i>

**Figure 2.** The PCR Amplification Using Primers Spanning in the Flanking Region of the 60-kb *O. sativa* Specific Segment in Three *Oryza* AA Genome Species.

(1) *O. glaberrima*; (2) *O. barthii*; (3) *O. glumaepatula*.

(Figure 3). The phylogenetic distribution of this region suggests that the genes encoded in the segment originated around 1 MYA or later.

Gene Structure and Origination of Annotated Genes

Twelve candidate genes, two embedded in PackMULEs (*Os03g01410* and *Os03g01520*) and one embedded in a *Helitron* (*Os03g01442*), and one complete *TyGypsy* retrotransposon were annotated in the 60-kb segment in *O. sativa* (Figure 1 and Table 1). Among the genes, 11 were classified as hypothetical or expressed genes. Only *Os03g01410*, contained within a PackMULE, could be assigned a putative function, based on TIGR ontology assignments, and was proposed to encode a 'STK kinase' (Table 1). In order to detect the origin of these 12 genes, we searched the *O. sativa* ssp. *japonica* genome and were able to identify candidate parental genes for 10 genes, and all the paralogs had high overall DNA sequence identity (greater than 93%), which is suggestive of being derived from recent duplication and transpositional events (Table 1). The two genes (*Os03g01460* and *Os03g01520*) without paralogs elsewhere in the *O. sativa* genome may have originated de novo, or the parental genes from which they were

derived may have subsequently been deleted, rearranged, or degraded.

By thoroughly comparing the gene structures of the 10 pairs of paralogs in *O. sativa* ssp. *japonica*, two distinct classes were revealed. First, five genes (*Os03g01410*, *Os03g01420*, *Os03g01450*, *Os03g01490*, and *Os03g01500*) had chimeric gene structures in which part of the parental gene was combined with additional sequences. Specifically, *Os03g01410* was a chimera composed of part of the *Os01g72700* gene and a 60-bp newly recruited sequence at the 3' end (Figure 4A and Supplemental Figure 1). *Os01g72700* was annotated as an ATP binding protein and its corresponding mRNA has been identified in several different rice tissues. *Os03g01420* and *Os03g01490* shared 450 bp of partial homologous coding sequence and appeared to have been created by an inverse tandem duplication event (Figures 4B and 5, and Supplemental Figure 2). However, we could not locate the homologous sequence for the remaining coding sequences for these two genes. Gene *Os03g01450* was found to have originated from three separate sequences. *Os04g32150* and *Os06g11900* recombined to form the first three exon of *Os03g01450*, and then combined with the flanking sequence to generate the entire gene structure (Figure 4C). The first intron of *Os03g01500* was very large (>10 kb) and appears to have originated from three independent sequences composed of: (1) the first exon (270 bp) and almost all of the first intron (780) of gene *Os12g24870*, (2) part of the first exon of *Os01g55880* (110 bp), and (3) 9 kb of flanking sequence. However, the remaining sequence of *Os03g01500* had no paralogous sequence identified (Figure 4D).

In the second class, an internal duplication within the 60-kb unique region appears to have created a second copy of three genes. The respective gene pairs are *Os03g01420*–*Os03g01490*, *Os03g01430*–*Os03g01480* and *Os03g01436*–*Os03g01470*, and a high level of sequence identity (99.89%) for this 8.5-kb region suggests this was a very recent duplication event, as shown in Figure 5. However, even though the tandem duplication was recent, two paralogous genes showed noteworthy divergence regarding gene structure. Specifically, the *Os03g01420*–*Os03g01490* gene pair had similar alternative isoforms (an alternative second exon), but contained distinct 3' terminal ends, of which *Os03g01490* was 36 bp longer (Figure 4B). Finally, the *Os03g01436* and *Os03g01470* gene pair was

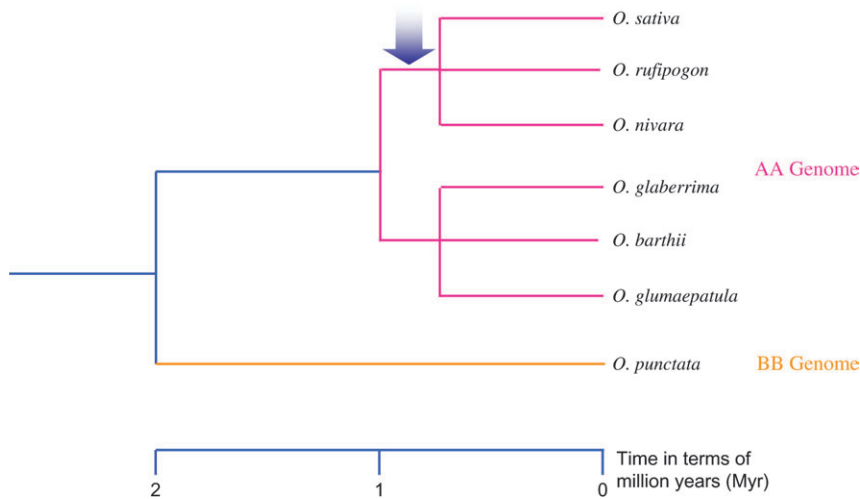


Figure 3. The Phylogenetic Relationship of AA Genome *Oryza* Species Rooted by BB Genome Species *O. punctata*.

The blue arrow indicates the origination point of this 60-kb segment.

found to have used the sense and anti-sense strands of one nearly identical DNA segment, respectively, encoding two totally distinct transcribed genes with open reading frames (ORFs) of more than 150 amino acid residues each.

Substitution Test between Paralogs

We investigated the functionality of six of the 12 *O. sativa* candidate genes that had clearly identifiable paralogues by calculating the K_a and K_s values (Table 1). For three of the remaining six candidate genes (*Os03g01460*, *Os03g01500*, and *Os03g01520*), we were not able to identify their corresponding parental genes (*Os03g01460* and *Os03g01520*) or coding sequence (*Os03g01500*) as described above. K_a and K_s values for the final three genes (*Os03g01436*, *Os03g01470*, and *Os03g01450*) could not be calculated because they had totally different ORFs. That is, paralogous genes *Os03g01436* and *Os03g01470* used sense and antisense strands as ORFs, respectively, and *Os03g01450* shared some sequences with *Os04g32150* and *Os06g11900*, but their gene structures were different compared to either of them with non-alignable ORFs.

K_s values for two of the candidate genes, *Os03g01410* and *Os03g01442*, were relatively small, with a value of 0.176 and 0.0003, respectively, and zero for paralogous gene pairs (*Os03g01480* vs *Os03g01430* and *Os03g01420* vs *Os03g01490*) due to their nearly identical sequences. Using a synonymous substitution rate of 0.65/100 Myr/synonymous site, as estimated for the *Adh* loci of grasses (Gaut et al., 1996), we estimated that the gene duplication events for all five paralogous duplicate genes ranged from 14 Myr (*Os03g01410*) to 0.02 Myr (*Os03g01442*) (Table 1).

Os03g01410 showed a low K_a/K_s value (0.14) that significantly deviated from neutrality ($= 0.5$) with $p \sim 0.011$ under the LRT test, indicating this gene is functionally constrained under purifying selection (Table 1). By contrast, *Os03g01442* had an excess number of non-synonymous substitutions, with four replacement substitutions and zero synonymous substitutions.

The LRT test gave a marginally significant p -value of 0.13, given an omega (K_a/K_s) = 1 under neutrality. Such an excess of non-synonymous substitutions suggests the possibility of adaptive evolution of gene *Og03g01442* after gene duplication.

Expression Analysis

Eight of 12 *O. sativa* candidate genes appeared to be transcribed, as evidenced by the presence of either EST and/or FL-cDNA sequence in Genbank (Table 1 and Supplemental Figure 3). In all eight cases, at least two transcript sequences could be found in Genbank. All but two (*Os03g01436* and *Os03g01470*) of these eight genes have multiple exons with conventional splice junctions. Finally, mRNA accumulation of three genes (*Os03g01420*, *Os03g01442* and *Os03g01490*) appeared to be fairly high in vivo, as revealed by the presence of 65, 12, and 46 independent EST sequences in Genbank, respectively, as well as Massively Parallel Signature Sequencing (MPSS) expression signatures (Nakano et al., 2006).

The presence of abundant ESTs allowed us to analyze tissue-specific profiles of mRNA accumulation. As shown by the UniGene Profile Viewer (Wheeler et al., 2008), both *Os03g01420* (with Chi square $p \sim 2 \times 10^{-9}$) and *Os03g01490* ($p \sim 8 \times 10^{-6}$) were highly associated with ESTs derived from callus tissue (Supplemental Figure 4A and 4B). *Os03g01442* mRNA was also found in callus tissue, but only a few copies were detected, which could not ensure a robust statistical test (Supplemental Figure 4C). However, MPSS data derived from mRNA isolated from various rice tissues, 6 h post inoculation with *Xanthomonas oryzae* pv. *oryzae*, revealed that mRNA from the *Os03g01442* gene was present in leaf tissue only with a normalized tag count of 34. Interestingly, although the numbers of ESTs were quite low, mRNA accumulation for genes *Os03g01470* and *Os03g01520* were only detected in callus tissue and mRNA accumulation for genes *Os03g01436* and *Os03g01450* were only detected in leaf tissue subjected to gamma radiation. In total, except for *Os03g01500*, whose ESTs data lack specific tissue information, all other seven genes appear to be involved in manipulated

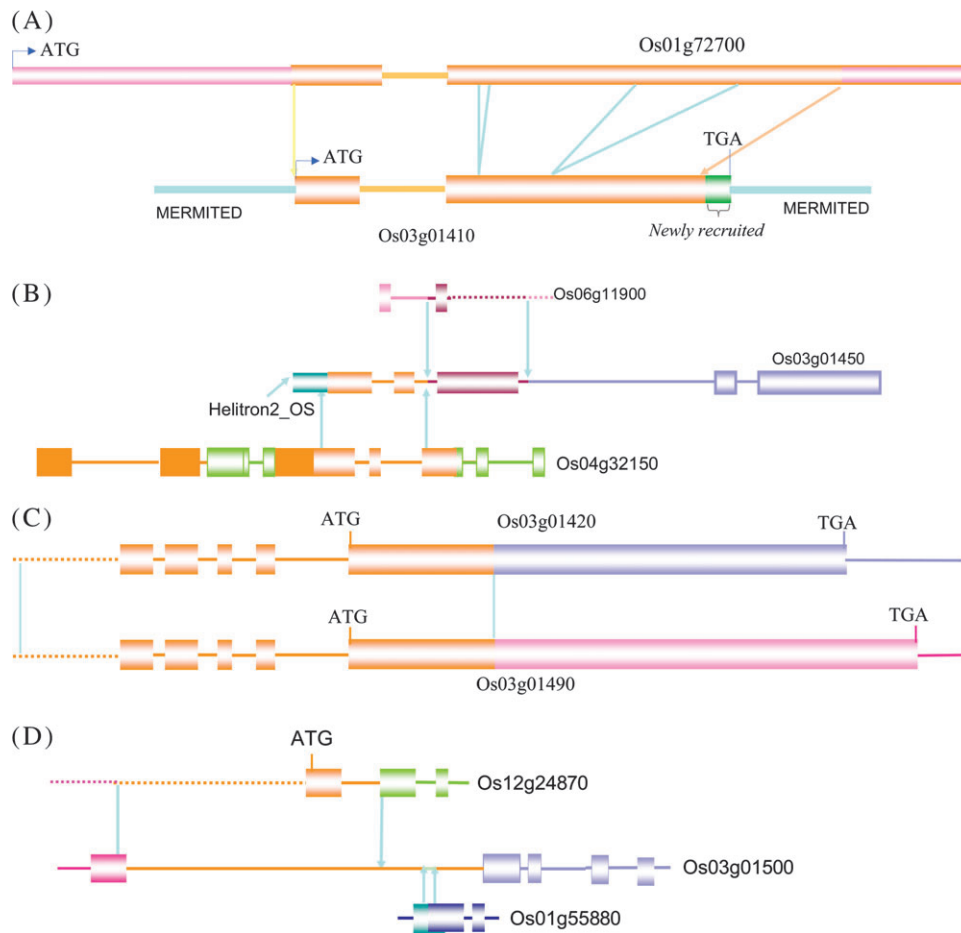


Figure 4. The Schematic Sketch of Five Chimeric Gene Origination Patterns.

- (A) *Os03g01410*.
- (B) *Os03g01450*.
- (C) *Os03g01490* and *Os03g01420*.
- (D) *Os03g01500*.

The homologous regions are shown in the same color. Solid boxes stand for exon, and solid lines indicate introns. The genes showing both start and stop codons have complete gene structure, otherwise only partial genes were drawn.

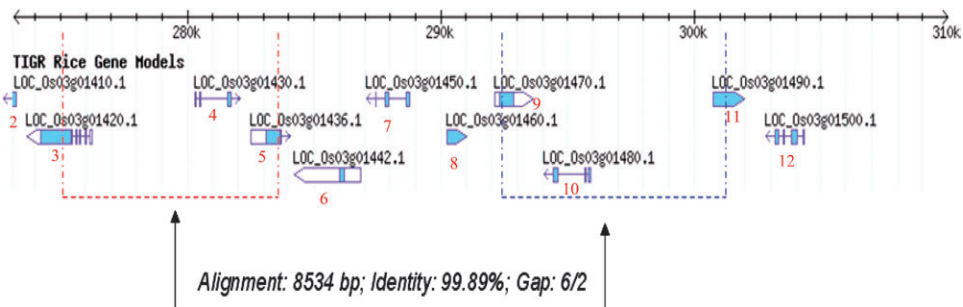


Figure 5. A Region Showing the Recent Tandem Inverse Duplication (in Dash Line Boxes).

tissue (callus), general defense, resistance responses, or various other responses that may have caused the cell death. In contrast, all paralogs showed no or less mRNA accumulation from defense response-derived tissues (see Supplemental Figure 4D–4G).

We further conducted an experimental profiling for two of the genes (*Os03g01442* and *Os03g01450*), for which we also generated polymorphism data. The mRNA accumulation profile of *Os03g01442* was found to be quite different when

compared to its parental gene. RT-PCR results show that *Os03g01442* mRNA could be detected at moderate levels in leaf and root tissues but not in stems or flowers (Figure 6). In contrast, we did not detect any mRNA in these tissues for *Os01g69904*, the paralogous counterpart of *Os03g01442* (Figure 6). No mRNA was detected for *Os03g01450* under normal growth conditions (Figure 6), which is consistent with MPSS data showing that *Os03g01450* tags could be detected in leaf tissue subjected to gamma radiation.

Population Genetics Analysis

To determine the pattern of DNA variation in this 60-kb segment in *O. sativa*, we investigated the nucleotide polymorphism spectrum of annotated protein genes for *Os03g01442* and *Os03g01450* across 30 different *O. sativa ssp. japonica* accessions that represented broad geographical distribution (Table 3). We observed a relatively low polymorphism rate in the two genes tested, consistent with previous studies (Caicedo et al., 2007). Furthermore, all neutrality tests (Tajima's D, Fu and Li's D, and Fay and Wu's H) with both negative values, and a coalescence simulation test revealed a biased frequency spectrum that deviated significantly from neutrality for *Os03g01442*. *Os03g01450* showed the same trend with a negative Tajima's D, although the *p*-value was marginally significant, at 0.04 (Table 3).

DISCUSSION

Chimeric genes can be generated through DNA-level recombination or retroposition-targeting mechanisms (Arguello et al., 2007). Chimeric genes derived from multiple parental loci, due to their potential to evolve novel functions, have provided

a very useful system to study gene evolution and its role in species diversification. For DNA-level recombination, several molecular mechanisms (homologous and non-homologous) have been observed to recombine different genic and non-genic regions to create chimeric genes (Roth and Wilson, 1988; Stankiewicz and Lupski, 2002). In retroposition-based chimeric gene formation, retroposed copies can recruit target sequences to form novel gene structures. In a previous effort to systematically detect retroposed genes in the rice genome (*O. sativa ssp. indica*), Wang et al. (2006) detected extensive origination activities that led to the formation of a large number of chimeric retroposed genes. Subsequent studies confirmed that the majority of these retroposed genes are also present in the *O. sativa ssp. japonica* genome (Fan et al., unpublished). In this study, consistently with previous findings, we annotated 12 putative young genes and a single TyGypsy retrotransposon in a contiguous 60-kb sequence in *O. sativa ssp. japonica*, five of which appeared to be chimeric genes created by DNA-level recombination. We found two genes (*Os03g01450* and *Os03g01500*) that were created by a combination of two genes, and one gene (*Os03g01410*) that was formed from a paralogous gene and a flanking sequence. The homologous gene pair, *Os03g01420* and *Os03g01490*, appears to contain chimeric gene structures with shared partial homologous sequences. An excess of chimeric gene formation appears to be a phenomenon of the grass species. This is in contrast to data from *Arabidopsis* and other dicot species, where no chimeric retroposed genes were identified (Fan et al., 2007b) and few chimeric gene structures, formed through DNA-level exon shuffling, have been reported (Drea et al., 2006; Domon and Steinmetz, 1994). The higher rate of chimeric gene formation through gene duplication and the generation of a larger number of functional genes in rice

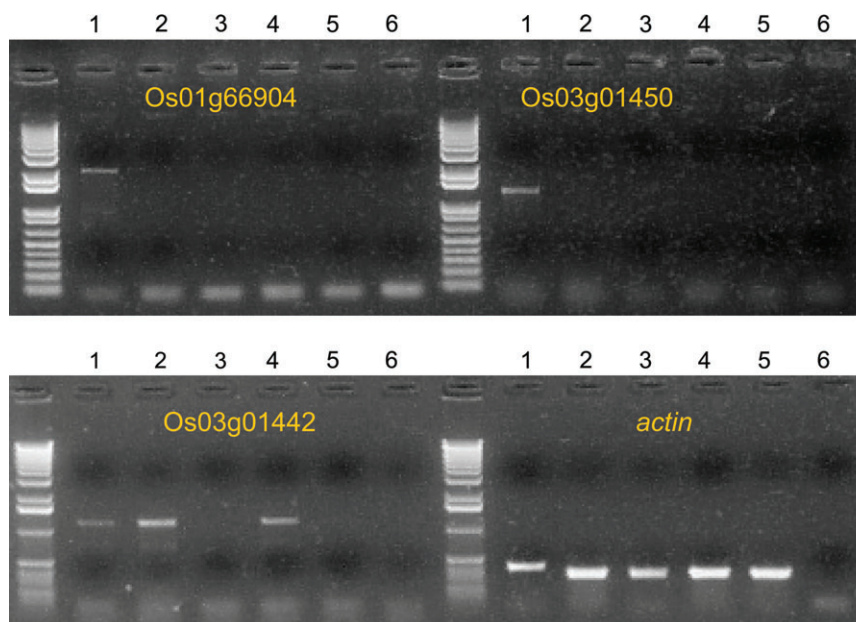


Figure 6. Expression Analysis of *Os3g01442*, *Os03g01450*, *Os01g69904*, and Transcript Internal Control *actin* (with 35 Cycles of PCR). (1) Nipponbare genomic DNA; (2) Nipponbare leaf cDNA; (3) Nipponbare stem cDNA; (4) Nipponbare root cDNA; (5) Nipponbare flower cDNA; (6) Negative control.

Table 3. Levels of Polymorphism of Two Genes in *O. sativa* ssp. *japonica* and Neutrality Tests on the Site Frequency Spectrum.

Summary statistic	<i>Os03g01450</i>	<i>Os03g01442</i>
<i>N</i>	30	30
<i>L</i>	820	1341
<i>S</i>	5	7
Π	0.00063	0.00035
Θ	0.00154	0.00132
Tajima's <i>D</i>	-1.62, $p = 0.04^*$	-2.17, $p = 0.001^{**}$
Fu and Li's <i>D</i> *	-1.48, $p = 0.15$	-3.60, $p = 0.003^{**}$
Fay and Wu's <i>H</i>	0.48, $p = 0.52$	-7.27, $p = 0.003^{**}$

Note: The Fay and Wu's *H* was calculated using paralogous sequences as outgroup. *N*, population size; *L*, gene length (bp); *S*, the number of segregation sites. * The significance as $P < 0.05$; ** the significance as $P < 0.01$.

may demonstrate that the diversification of the grass species is a mirror of their broad ecological adaptation and morphological complexity.

Our analysis of the automated gene models found in the TIGR rice annotation database in combination with the EST/cDNA evidence suggests that at least eight of the gene models need to be manually re-annotated. Specifically, the gene model for *Os03g01450* is in total conflict with the splicing structure revealed by EST sequences (Supplemental Figure 3E). Moreover, it constitutes a sense/antisense pair with *Os03g01470* and shares at least 700 base pairs of sequence. Similarly, the gene structures of *Os03g01500* and *Os03g01520* also form a non-exonic overlapping gene pair (Supplemental Figure 3D). Lastly, four genes (*Os03g01420*, *Os03g01442*, *Os03g01490*, and *Os03g01520*) have alternative splice sites, thereby encoding two splicing isoforms each.

Although the species tree indicates the evolutionary age of this 60-kb region should be around 1 Myr, estimations of gene duplication events based on *Ks* ranged from 0.02 to 14 Myr. Such a conflict might be caused by the following factors. First, there may be a greater variation for substitution rates among genes than previously thought, so the rate we used derived from *Adh* may not be appropriate for other genes. An interesting alternative would be that, in some cases, such as with *Os03g01410* embedded within a PackMULE, a gene might have actually emerged 14 MYA and then moved to its current location more recently. This latter hypothesis predicts that one would find many homologous sequences in non-syntenic regions in several of the wild relatives of rice. It will be possible to test this idea more systematically when complete genomic sequences from the other *Oryza* species become available.

Both comparative genomics and experimental analysis showed that this 60-kb segment in *O. sativa* ssp. *japonica* started evolving after the divergence of Asian (*O. sativa*, *O. rufipogon*, and *O. nivara*) and African species (*O. glaberrima* and *O. barthii*) about 1 MYA. Given this segment contained 12 relatively young putative genes in *O. sativa*, the most straightforward explana-

tion for its origination is a single segmental duplication. This hypothesis seems highly unlikely for a number of reasons. First, as shown above, an inverse tandem duplication occurred, which contributed three pairs of genes. Second, candidate parental genes of the 12 genes were distributed across the entire genome (i.e. chromosomes 1, 3, 4, 6, and 12). Third, the overall identity between paralogous duplicates fluctuated from 93 to 100%, which points to different evolutionary ages, corresponding to 0.02 Myr ($Ks = 0.0003$) and 14 Myr ($Ks = 0.17$). Fourth, three of the candidate genes are embedded in transposable elements, PackMULEs and *Helitrons*, known to be associated with new gene formation and movement. Fifth, the region contains a complete *TyGypsy* LTR retrotransposon that was estimated to have inserted about 0.5 MYA. Finally, we found that the orthologous regions in *O. nivara* and *O. rufipogon* were smaller than in *O. sativa* and contained fewer genes (Yu et al., unpublished data), which suggests that the evolution of this segment is still ongoing through recurrent recombination and transposition via transposable elements.

Thus, multiple independent duplications and TE-mediated transpositions are a plausible explanation for the origination of all these new genes in this region. Our hypothesis is supported by the location of these genes and retrotransposons. This region is located between 250 and 310 Kb at the subtelomeric region of chromosome 3 in *O. sativa* ssp. *japonica*. It has been reported that subtelomeric regions, usually around 500 Kb near the tip of each chromosome, have much higher recombination activity and tend to be gene-rich and highly transcriptionally active (Mizuno et al., 2006). Thus, frequent recombination renders it possible for such a short region to accumulate more duplicated sequences in a short time span, and the local environment makes these duplicates more likely to maintain or evolve a new transcriptional activity, like recruiting new exons or forming sense/antisense gene pairs.

What mechanism(s) is implicated in these numerous recombination and transposition events? A straightforward possibility is repeat element-mediated homologous recombination, given that subtelomeres and telomeres are known to be enriched with transposon elements, which have been reported to be important for the creation of new genes in the *Drosophila* genomes (Anderson et al., 2008) and plant genomes (Hudson et al., 2003; Wang et al., 2006; Hollister and Gaut, 2007; Gaut and Ross-Ibarra, 2008). One of the genes within this 60-kb region (*Os03g01442*) is embedded within a *Helitron*, an autonomous DNA transposon. Consistent with this, its parental gene, *Os01g69904*, is also located adjacent to one *Helitron* repeat. *Helitrons* are known to be able to shuffle pre-existing genes in plants (Lal et al., 2003; Kapitonov and Jurka, 2001, 2007; Hollister and Gaut, 2007). Furthermore, the highly conserved sequences found in the flanking regions of *O. glaberrima*, *O. barthii*, and *O. glumaepatula* (Supplemental Figure 5) and the presence of apparent transposon insertion sites in the flanking region of *O. sativa* (Supplemental Figure 6) further support our conclusion that recombination and/or transposition via transposable elements were highly active

mechanisms that led to the recent creation of young genes in *O. sativa*.

It has been reported that the cultivated rice genome encodes 898 functional retrogenes (Wang et al., 2006), which is in contrast to the prevailing view that plant genomes contain only a few retrogenes. Moreover, about 100 of the 898 retrogenes are very young, as suggested by their *Ks* values, which are smaller than 0.1 (less than 10 Myr). In our study, we identified 12 candidate young genes and one retrotransposon in *O. sativa*. Except for retrotransposons, whose transcriptional activity might be repressed by the host genome, nine of 12 genes showed evidence of functionality based on EST/cDNA/MPSS data and/or evolutionary analyses. Remarkably, some (e.g. *Os03g01436* and *Os03g01450*) appear to be involved in defense responses. The emergence of new genes related to stress and defense could be rationalized by the fact that rice has broad ecological adaptation and is under large selective pressures to protect itself against natural disasters and predators. Here, we only analyzed about the first 10% of the short arm of chromosome 3, and found a number of functional genes that originated recently through independent recombination and transpositional events. If the subtelomeric region of chromosome 3 is representative of the remaining 23 chromosome arms, then it is highly likely that we will identify many more young genes supporting our hypothesis that subtelomere serves as a hot bed for gene origination and evolution by recruiting DNA-level duplicated genes in rice.

We performed further detailed analyses for two of the identified 12 genes in the segment. The polymorphism spectrum of *Os03g01450* showed a slight deviation from neutrality as revealed by a marginally significant *p*-value (0.04) using the Tajima's *D* test (Table 3). By contrast, both expression and population genetic analyses implied that *Os03g01442* may be subject to positive selection during its origination and fixation. A differential mRNA accumulation pattern was observed for *Os03g01442*, with no mRNA detected in stems and flowers and moderate mRNA levels in leaves and roots. In contrast, no transcript was detected for *Os03g01442*'s parental copy *Os01g69904* in the same tissues. Furthermore, the biased frequency spectrum with an excess of both rare alleles and high frequency polymorphisms in *Os03g01442* also suggested the possibility that positive selection is acting on this gene. Although further analysis should be conducted in this region to rule out the possibility of demographic effects for the biased polymorphism spectrum (with a broader sample base, more young genes and flanking sequences), this case analysis provides an example that selection may be the acting force to drive the fixation of young genes in rice.

METHODS

Sequencing 1.5-Mb BAC Tiles from *O. glaberrima* and *O. punctata*

We selected two ~1.5-Mb minimum tiling paths of overlapping BAC clones from *O. glaberrima* and *O. punctata* from the

short arms of chromosome 3 utilizing previously described BAC libraries and BAC fingerprint/end-sequenced physical maps (Ammiraju et al., 2006; Kim et al., 2007, 2008). Each BAC clone was shotgun-sequenced, finished and sequence-validated using standard procedures as previously described (IRGSP, 2005), such that the final finished sequences had an error rate of less than one base in 10 000. Overlapping BAC sequences from each species were then manually assembled into ~1.5-Mb pseudomolecules and used for further analysis.

Searching the *O. sativa ssp. japonica* Specific Sequence by Comparative Analysis

We performed genomic pairwise comparison between *O. sativa ssp. japonica* genome and 1.5-Mb *O. glaberrima* chromosome 3 short arm sequences. The annotation and coding sequences (CDS) of *O. sativa ssp. japonica* were downloaded from TIGR (www.tigr.org/tdb/e2k1/osa1/). A MegaBLAST of 1.5-Mb *O. glaberrima* sequence to the CDS of *O. sativa ssp. japonica* was conducted. We searched orthologous sequences between two species; meanwhile, we also paid attention to the unique sequence found only in *O. sativa ssp. japonica*, but absent in *O. glaberrima*. We further searched the *O. sativa ssp. japonica* sequence to sequence of *O. punctata*. In such effort, we found an *O. sativa ssp. japonica* unique segment bearing sequence in length of 60 kb.

Sequence Analysis of the *O. sativa* Specific 60-kb Segment

To determine if the 60-kb region identified in *O. sativa ssp. japonica* was unique to the *japonica* genome, we used two approaches. To probe the wild Asian species, we used BLAST to identify any BESs from the *O. rufipogon* and *O. nivara* BAC libraries that were similar to the 60-kb *japonica* sequence, then detected those BACs that locate in the subtelomeric region of the chromosome 3 short arm using the Finger Printed Contigs (FPC)-based physical map and SyMAP synteny browser (Kim et al., 2008; Soderlund et al., 2006). For the African and South American species, we performed PCR amplification on genomic DNA isolated from *O. glaberrima*, *O. barthii*, and *O. glumeatulata*, using a pair of primers specific to *O. glaberrima* and *O. sativa ssp. japonica* located in the flanking regions of the 60-kb sequence (see Figure 1). Based on the TIGR rice annotation database (Release 5), we were able to identify 12 genes and one retrotransposon in this segment. In order to understand the evolution pattern and history of these genes, we implemented a robust strategy to search for their candidate parental genes (paralogs) in the *O. sativa ssp. japonica* genome. In brief, we searched each gene together with its 10K flanking sequence against the whole genome using BLASTN. We then mapped the 1.5-Mb chromosome 3 sequence of *O. sativa ssp. japonica* to the rest of the *japonica* RefSeq using the ChainSelf pipeline developed by UCSC. We manually checked the hits generated by both methodologies and retrieved the best hits to serve as the parental genes.

We compared both CDS and genomic sequence for both paralogs to gain insight into gene structure and origin, and we further calculated the Ka/Ks ratio using maximum likelihood algorithm using the PAML package (Yang, 2007). The significance of Ka/Ks that deviated from neutrality (= 0.5) were tested using the LRT (Emerson et al., 2004). Specifically, we aligned protein sequences of the parental gene and the daughter gene with MUSCLE (Edgar, 2004) and converted the protein alignment into the codon-based nucleotide alignment with the Pal2nal script (Suyama et al., 2006). We then used codeml with fixed and free omega models to test whether any of the young genes detected were under selective constraint, namely Ka/Ks was significantly smaller than 0.5 (Yang, 2007).

We re-annotated four incorrect TIGR gene models using the UniGene rice EST/mRNA dataset. ESTs were mapped to the genome by BLAT (Kent, 2002) and all ambiguous or low-quality mappings were discarded (Zhang et al., 2006). Then, we remapped these hits to the genome using SIM4 (Florea et al., 1998) and GeneSeqer (Schlueter et al., 2003) with the rice scoring matrix in order to refine the entire splicing structure. Gene structures with the highest identity and the most standard splicing junction were retained.

DNA Extraction, PCR Amplification, and Sequencing

Total genomic DNA was extracted from fresh leaves of a single plant using the Qiagen DNeasy kit following the manufacturer's protocol. PCR reactions were performed using Invitrogen Taq polymerase, with annealing temperature adjusted based on the length of fragments with 1 kb min^{-1} . Double-stranded PCR products were purified using either the Qiagen PCR purification or Qiagen miniprep Gel purification kits. Purified PCR products were sequenced using the ABI-3730XL 96-capillary automated DNA sequencer. Sequences were edited and assembled. Clustal X was used to align sequences for further analyses (Thompson et al., 1997). Manual adjustments were made where necessary.

Expression Analysis

As mentioned above, we mapped the latest UniGene rice EST/mRNA dataset to the complete genome, which consists of more than 1 000 000 entries. The expression profiles for two genes were further investigated using reverse transcription (RT)-PCR in different tissues grown under normal conditions. Total RNA was extracted from leaf, root, stem, and entire flower bud using a Qiagen total RNA extraction kit. cDNA were generated using the Invitrogen RT-PCR kit and full description of RT-PCR was described previously. The constitutively expressed gene *actin* was used as internal control to quantify the density of cDNA.

Population Genetics Analysis

We sampled the worldwide collection of *O. sativa ssp. japonica* accessions to generate a nucleotide frequency spectrum for population genetics analysis. Most *O. sativa ssp. japonica*

accessions were chosen from a wide range of Asia, with a few samples from Africa. Basic population genetic analysis was implemented in DnaSP (Rozas et al., 2003). Sequence diversity was quantified as nucleotide diversity (π) (Nei, 1987) and Watterson's θ (1975). Tests of deviation from neutrality were conducted using Tajima's D (1989), Fu and Li's D (1993), and Fay and Wu's H (2000) tests. We further used coalescent simulation to assess the significance of the statistic for the all parameters generated. The neutral coalescent process was simulated using 2000 replicates, with the number of segregating sites set to that observed in the data.

SUPPLEMENTARY DATA

Supplementary Data are available at *Molecular Plant Online*.

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REFERENCES

- Ammiraju, J.S., et al. (2006). The *Oryza* bacterial artificial chromosome library resource: construction and analysis of 12 deep-coverage large-insert BAC libraries that represent the 10 genome types of the genus *Oryza*. *Genome Res.* **16**, 140–147.
- Anderson, J.A., Song, Y.S., and Langley, C.H. (2008). Molecular population genetics of *Drosophila* subtelomeric DNA. *Genetics* **178**, 477–487.
- Arguello, J.R., Fan, C., Wang, W., and Long, M. (2007). Origination of chimeric genes through DNA-level recombination. *Genome Dyn.: Protein and Gene Evolution* **3**, 131–146.
- Bennetzen, J.L. (2007). Patterns in grass genome evolution. *Curr. Opin. Plant Biol.* **10**, 176–81.
- Betran, E., Thornton, K., and Long, M. (2002). Retroposed new genes out of the X in *Drosophila*. *Genome Res.* **12**, 1854–1859.
- Caicedo, A.L., et al. (2007). Genome-wide patterns of nucleotide polymorphism in domesticated rice. *PLoS Genet.* **3**, 1745–1756.
- Domon, C., and Steinmetz, A. (1994). Exon shuffling in anther-specific genes from sunflower. *Mol. Gen. Genet.* **244**, 312–317.
- Drea, S.C., Lao, N.T., Wolfe, K.H., and Kavanagh, T.A. (2006). Gene duplication, exon gain and neofunctionalization of OEP16-related genes in land plants. *Plant J.* **46**, 723–735.
- Drosophila 12 Genomes Consortium** (2007). Evolution of genes and genomes on the *Drosophila* phylogeny. *Nature* **450**, 203–218.

- Edgar, R.C. (2004). MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res.* **32**, 1792–1797.
- Emerson, J.J., Kaessmann, H., Betran, E., and Long, M. (2004). Extensive gene traffic on the mammalian X chromosome. *Science* **303**, 537–540.
- Fan, C., Emerson, J.J., and Long, M. (2007a). The origin of new gene. In *Evolutionary Genomics and Proteomics*, M. Pagel and A. Pomiankowski, eds (Sunderland, Massachusetts, USA: Sinauer Associates, Inc.), pp. 27–44.
- Fan, C., Vibranovski, M., Chen, Y., and Long, M. (2007b). A microarray-based genomic hybridization method for identification of new genes in plants: case analyses of *Arabidopsis* and *Rice*. *J. Integ. Plant Biol.* **49**, 915–926.
- Fay, J., and Wu, C.-I. (2000). Hitchhiking under positive Darwinian selection. *Genetics* **155**, 1405–1413.
- Florea, L., Hartzell, G., Zhang, Z., Rubin, G.M., and Miller, W. (1998). A computer program for aligning a cDNA sequence with a genomic DNA sequence. *Genome Res.* **8**, 967.
- Fu, Y., and Li, W.-H. (1993). Statistical tests of neutrality of mutations. *Genetics* **133**, 693–709.
- Gaut, B.S., and Ross-Ibarra, J. (2008). Selection on major components of angiosperm genomes. *Science* **320**, 484–486.
- Gaut, B.S., Morton, B.R., McCaig, B.C., and Clegg, M.T. (1996). Substitution rate comparisons between grasses and palms: Synonymous rate differences at the nuclear gene *Adh* parallel rate differences at the plastid gene *rbcl*. *Proc. Natl Acad. Sci. U S A.* **93**, 10274–10279.
- Ge, S., Sang, T., Lu, B.R., and Hong, D.Y. (1999). Phylogeny of rice genomes with emphasis on origins of allotetraploid species. *Proc. Natl Acad. Sci. U S A.* **96**, 14400–14405.
- Hollister, J.D., and Gaut, B.S. (2007). Population and evolutionary dynamics of *Helitron* transposable elements in *Arabidopsis thaliana*. *Mol. Biol. Evol.* **24**, 2515–2524.
- Hudson, M.E., Lisch, D.R., and Quail, P.H. (2003). The FHY3 and FAR1 genes encode transposase-related proteins involved in regulation of gene expression by the phytochrome A-signaling pathway. *Plant J.* **34**, 453–471.
- International Rice Genome Sequencing Project (2005). The map based sequencing of the rice genome. *Nature* **436**, 793–800.
- Kim, H., et al. (2008). Construction, alignment and analysis of twelve framework physical maps that represent the ten genome types of the genus *Oryza*. *Genome Biol.* **9**, R45.
- Kim, H., San Miguel, P., Nelson, W., Collura, K., Wissotski, M., Walling, J.G., Kim, J.P., Jackson, S.A., Soderlund, C., and Wing, R.A. (2007). Comparative physical mapping between *O. sativa* (AA genome type) and *O. punctata* (BB genome type). *Genetics* **176**, 379–390.
- Kapitonov, V.V., and Jurka, J. (2001). Rolling-circle transposons in eukaryotes. *Proc. Natl Acad. Sci. U S A.* **98**, 8714–8719.
- Kapitonov, V.V., and Jurka, J. (2007). Helitrons on a roll: eukaryotic rolling-circle transposons. *Trends Genet.* **23**, 521–529.
- Kent, W.J. (2002). BLAT—the BLAST-like alignment tool. *Genome Res.* **12**, 656–664.
- Lal, S.K., Giroux, M.J., Brendel, V., Vallejos, C.E., and Hannah, L.C. (2003). The maize genome contains a helitron insertion. *Plant Cell.* **15**, 381–391.
- Long, M., Betran, E., Thornton, K., and Wang, W. (2003). The origin of new genes: glimpses from the young and old. *Nat. Rev. Genet.* **4**, 865–875.
- Marques, A.C., Dupanloup, I., Vinckenbosch, N., Reymond, A., and Kaessmann, H. (2005). Emergence of young human genes after a burst of retroposition in primates. *PLoS Biol.* **3**, e357.
- Mizuno, H., Wu, J., Kanamori, H., Fujisawa, M., Namiki, N., Saji, S., Katagiri, S., Katayose, Y., Sasaki, T., and Matsumoto, T. (2006). Sequencing and characterization of telomere and subtelomere regions on rice chromosomes 1S, 2S, 2L, 6L, 7S, 7L and 8S. *Plant J.* **46**, 206–217.
- Nakano, M., Nobuta, K., Vemaraju, K., Tej, S.S., Skogen, J.W., and Meyers, B.C. (2006). Plant MPSS databases: signature-based transcriptional resources for analyses of mRNA and small RNA. *Nucleic Acids Res.* **34**, D731–D735.
- Nei, M. (1987). *Molecular Evolutionary Genetics* (New York: Columbia University Press).
- Ohno, S. (1970). *Evolution by Gene Duplication* (Berlin: Springer).
- Paterson, A.H., Freeling, M., and Sasaki, T. (2005). Grains of knowledge: genomics of model cereals. *Genome Res.* **15**, 1643–1650.
- Roth, D., and Wilson, J. (1988). Illegitimate recombination in mammalian cells. In *Genetic Recombination*, R. Kucherlapati and G.R. Smith, eds (Washington, DC, USA: American Society of Microbiology), pp. 621–653.
- Rozas, J., Sanchez-DelBarrio, J.C., Messeguer, X., and Rozas, R. (2003). DnaSP, DNA polymorphism analyses by the coalescent and other methods. *Bioinformatics* **19**, 2496–2497.
- Schlueter, S.D., Dong, Q., and Brendel, V. (2003). GeneSeqer@PlantGDB: gene structure prediction in plant genomes. *Nucleic Acids Res.* **31**, 3597–3600.
- Soderlund, C., Nelson, W., Shoemaker, A., and Paterson, A. (2006). SyMAP: a system for discovering and viewing syntenic regions of FPC maps. *Genome Res.* **16**, 1159–1168.
- Stankiewicz, P., and Lupski, J.R. (2002). Molecular–evolutionary mechanisms for genomic disorders. *Curr. Opin. Genet. Dev.* **12**, 312–319.
- Suyama, M., Torrents, D., and Bork, P. (2006). PAL2NAL: robust conversion of protein sequence alignments into the corresponding codon alignments. *Nucleic Acids Res.* **34**, W609–W612.
- Tajima, F. (1989). Statistical methods for testing the neutral mutation hypothesis by DNA polymorphism. *Genetics* **123**, 585–595.
- Thompson, J., Gibson, T.J., Plewniak, F., Jeanmougin, F., and Higgins, D.G. (1997). The Clustal X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. *Nucleic Acids Res.* **24**, 4876–4882.
- Wang, W., et al. (2006). High rate of chimeric gene origination by retroposition in plant genomes. *Plant Cell.* **18**, 1791–1802.
- Watterson, G. (1975). On the number of segregating sites in genetic models without recombination. *Theor. Popul. Biol.* **7**, 256–276.
- Wheeler, D.L., et al. (2008). Database resources of the National Center for Biotechnology Information. *Nucleic Acids Res.* **36**, D13–D21.
- Wing, R.A., et al. (2005). The *Oryza* Map Alignment Project: the golden path to unlocking the genetic potential of wild rice species. *Plant Mol. Biol.* **59**, 53–62.
- Yang, Z. (2007). PAML 4: phylogenetic analysis by maximum likelihood. *Mol. Biol. Evol.* **24**, 1586–1591.

- Yu, J., et al. (2002). A draft sequence of the rice genome (*Oryza sativa* L. ssp. *indica*). *Science* **296**, 79–92.
- Zhang, Y., Liu, X.S., Liu, Q.R., and Wei, L. (2006). Genome-wide in silico identification and analysis of cis natural antisense transcripts (cis-NATs) in ten species. *Nucleic Acids Res.* **34**, 3465–3475.
- Zhang, Y., Wu, Y., Liu, Y., and Han, B. (2005). Computational identification of 69 retroposons in *Arabidopsis*. *Plant Physiol.* **138**, 935–948.
- Zhu, Q., and Ge, S. (2005). Phylogenetic relationships among A-genome species of the genus *Oryza* revealed by intron sequences of four nuclear genes. *New Phytol.* **167**, 249–265.