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Research Article

Genetic innovations: Transposable element recruitment and *de novo* formation lead to the birth of orphan genes in the rice genome

Running Title: Origin of orphan genes in the rice genome

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Abstract

Orphan genes are genetic innovations that lack homologs in other lineages.

Orphan genes can rapidly originate and become substantially functional, yet the mechanisms underlying their origins are still largely unknown in plants. Here, we

investigated the origin of orphan genes in the *Oryza sativa* ssp. *japonica*

‘Nipponbare’ genome via genome-wide comparisons with ten closely related

Oryza species. We identified a total of 37 orphan genes in the Nipponbare genome

that show short sequence lengths, elevated GC content, and absence of introns.

Interestingly, half of the identified orphan genes originated via a distinctive

mechanism that involved the generation of new coding sequences via independent

and rapid divergence within the inserted transposable element. Our results provide

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valuable insight into genetic innovations in the model rice genome that formed on a very short time scale.

Key words: comparative genomics, origin, orphan gene, transposable element.

1 Introduction

Orphan genes are newly born genes, present in only one species or in a specific taxonomic clade but are evidently absent in other closely related taxa (Fischer & Eisenberg, 1999; Boffelli et al., 2004). Compared to other protein-coding genes, orphan genes usually possess several atypical genetic characteristics, such as small coding regions (Lipman et al., 2002), fast evolutionary rates (Domazet-Loso & Tautz, 2003), uncharacterized functional domains (Daubin & Ochman, 2004), and variable GC content (Arendsee et al., 2014; Palmieri et al., 2014; Sun et al., 2015; Xu et al., 2015). Although the functions of the majority of orphan genes are largely unknown, these genes appear to play an important, and sometimes even essential, roles in species-specific adaptation. For example, orphan genes can encode phylum-specific morphology

in *Hydra* (Khalturin et al., 2009), confer *Fusarium* resistance in wheat (Perochon et al., 2015), participate in early human brain development (Zhang et al., 2011), and participate in primary metabolic pathways to allow *Arabidopsis thaliana* adapt to environmental changes (Jones et al., 2016).

The identification and classification of orphan genes largely depends on two factors. Both the evolutionary distance between the target taxa/clade and its nearest sequenced relatives as well as the quality of genome sequence is used (Tautz & Domazet-Loso, 2011). Due to limited data availability, previous studies have mainly focused on the identification of orphan genes, specific to large taxonomic groups, which may span long evolutionary time scales. For example, 270 orphan genes were found in primates, based on comparisons with nonprimate species (Toll-Riera et al., 2009). In plants, 1,789 orphan genes were identified in Brassicaceae that lack homology in non-Brassicaceae plant species (Donoghue et al., 2011). However, having a long evolutionary history may actually reduce the power of analyses, because this may make it more difficult to identify orphan genes

since the homology relationship might be masked by increased sequence divergence. Recently, with the increasing number of available genomes from closely related species, it has become feasible to identify orphan genes on small evolutionary time scales. These short time-scales analyses can benefit to understand the origin and evolutionary processes of newly born genes.

Several excellent models have been proposed to explain the origin of orphan genes. For instance, a duplication-divergence mechanism is considered the main mechanism for the birth of orphan genes in zebrafish (Yang et al., 2013) and *Drosophila* (Zhou et al., 2008). Transposable elements (TEs) exaptation is another vital mechanism to consider in the origin of orphan genes in primates (Toll-Riera et al., 2009) and likewise in silkworm (Sun et al., 2015). Other studies have indicated that the majority of orphan genes could arise *de novo* from noncoding sequences (Wissler et al., 2013; Zhang et al., 2019). Recently, Prabh & Rödelsperger (2019) also proposed a mixed origin mechanism, incorporating the notion that an orphan gene could generate through both duplication-divergence

and the *de novo* mechanism. In plants, a previous study indicated the existence of four distinct original mechanisms for the formation of orphan genes in *A. thaliana*, including overlap with conserved gene loci, duplication followed by sequence divergence, TE exaptation, and *de novo* formation (Donoghue et al., 2011). However, the origin and dynamics of orphan genes in other plants were less studied.

The Asian cultivated rice *Oryza sativa* ssp. *japonica* is one of the youngest taxa (< 0.55 myr), and has a more comprehensively assembled and annotated genome. To date, at least 13 genomes of *Oryza* belonging to different lineages have been released, which provide an excellent opportunity to systematically identify orphan genes within these genomes (Stein et al., 2018). In this study, we attempted to find which original mechanism play an important role in the newly born genes. To answer this question, we investigated origin mechanisms for newly born orphan genes in the *Oryza sativa* ssp. *japonica* 'Nipponbare' using genome-wide comparisons with ten closely related *Oryza* species.

2 Material and Methods

2.1 Data sources

To investigate the origin of orphan genes, we selected the model rice plant *Oryza sativa* ssp. *japonica* as the focal taxon. Genomic data, including nucleotide and amino acid sequences of protein genes and the genome sequence of *O. sativa* ssp. *japonica* ‘Nipponbare’ were downloaded from TIGR release 7 (Kawahara et al., 2013). We also downloaded the genomic data for *O. sativa* ssp. *indica* ‘Shuhui 498’ (R498) from MBKbase (Du et al., 2017) and for nine other *Oryza* species (*O. barthii*, *O. brachyantha*, *O. glaberrima*, *O. glumaepatula*, *O. longistaminata*, *O. meridionalis*, *O. nivara*, *O. punctata*, and *O. rufipogon*) from the Gramene database version 52 (<http://www.gramene.org>) to use as outgroups. The genomic data of *O. sativa* ssp. *japonica* ‘Nipponbare’ were extensively compared with the data for three close relatives (*O. nivara*, and *O. rufipogon*, and *O. sativa* ssp. *indica*). All of the genomic data used in this study are summarized in Table S1.

2.2 Identification of orphan genes

To reliably identify orphan genes in the *O. sativa* ssp. *japonica* ‘Nipponbare’ (hereafter, the Nipponbare) genome, we performed genome-wide pairwise comparisons between Nipponbare and other ten *Oryza* genomes (outgroups). The identification of the orphan genes on long evolutionary time scales was mostly based on the use of amino acid alignments (Cai et al., 2006; Toll-Riera et al., 2009; Johnson & Tsutsui, 2011), which largely rely on genome annotation quality. Critically speaking, genome annotation artifact within the outgroup may affect orphan gene identification. Therefore, we implemented strict procedures to reduce annotation artifacts from the outgroups (Fig. 1 for details).

Initially, all of the 55 986 coding sequences (CDSs) annotated in the Nipponbare genome were used as queries to search against CDSs in the outgroup using BLAT(at the threshold of 80% sequence coverage) (Kent, 2002) and to search against genomes in the outgroup with BLASTN (default task: Megablas, E value ≤ 0.01). Adopting this approach enabled us to filter any homologous genes

that were perhaps misannotated in the outgroups. We also performed a TBLASTN search in the NCBI NR database (up until 2018/05/07) using amino acid sequences as queries to exclude any genes with homologs in other plants.

Moreover, we implemented a similar procedure which has been described as the “flanking gene method” (Freeling et al., 2008). This procedure utilizes neighboring syntenic genes to identify the presence or absence of putative orphan genes in the three closest relatives. Five to ten upstream and downstream genes from the orphan genes in Nipponbare were used for synteny analysis via BLASTP and TBLAST alignments (E value $\leq 10^{-5}$).

2.3 Characteristics of orphan genes

To better understand whether orphan genes possess special features, we analyzed three important genic characteristics of the orphan genes. Except for the identified orphan genes, all other protein-coding genes in the Nipponbare genome were considered nonorphan genes. Comparisons between orphan genes and nonorphan genes were performed using custom Perl scripts, including calculations of the protein length, intron number, and GC content at the CDS level. The

Mann-Whitney test was also applied to assess the differences between orphan and nonorphan genes (Bauer, 1972).

2.4 Functional signatures

Expression signatures of orphan genes were first determined via searches against publicly available full-length cDNA (FL-cDNA), expressed sequence tag (EST) (Pontius et al., 2003), and microarray data (<http://rice.plantbiology.msu.edu>). We also searched for all functional domains in orphan genes via the InterProScan programs at the European Bioinformatics Institute website (version 75.0) (EBI, <http://www.ebi.ac.uk/Tools/InterProScan>). Additionally, expression level were examined on the basis of 284 high-quality RNA-Seq datasets, deposited in the Rice Expression Database (<http://expression.ic4r.org/index>). An orphan gene was considered to be expressed at a threshold of fragments per kilobase of transcript per million fragments mapped (FPKM) greater than 0 in at least two samples.

To verify orphan genes expression levels, reverse transcription-polymerase chain reaction (RT-PCR) experiments were performed on eight randomly selected orphan genes. Primer information is provided in Table S2, and the house-keeping gene *Tubulin* was selected as an internal control. Total RNA was extracted from mixed rice tissue samples using an EastepTM Super Total RNA Extraction Kit (Promega, Beijing, China). Based on concentration and quality assays, performed with a NanoDrop ND 1000, 2 µg of total RNA was used to synthesize the first-strand cDNA with the GoScriptTM Reverse Transcription System (Promega, Beijing, China). Finally, high-quality cDNA was used as templates in the RT-PCRs. The RT-PCR products were resolved on 3% agarose gels via electrophoresis at 120 V for 30 minutes.

2.5 Tracing the birth of orphan genes

Based on syntenic analysis of the flanking genes in Nipponbare and its close relatives, we compared multiple sequences in the syntenic regions of the genomes of Nipponbare and its three close relatives via the MUSCLE program (Edgar, 2004) using the codon alignment option with the Nipponbare CDSs as the

template. Any orphan genes that had partial homologs in the syntenic regions of its close relatives were considered as noncoding homologous sequences. If an orphan gene lacked noncoding homologous sequences in the close relatives, we searched for TEs via CENSOR program against the RepBase database (Kohany et al., 2006) in the syntenic regions of Nipponbare and its close relatives .

3 Results

3.1 Identification of thirty-seven functional orphan genes in the Nipponbare genome

We implemented a strict pipeline on the basis of both homology and synteny to search for newly born orphan genes (< 0.55 myr) in Nipponbare after it diverged from *O. rufipogon* (Fig. 1). We identified 38 genes without any homologs in the ten *Oryza* outgroups via extensive comparisons. Among these genes, one gene (*LOC_Os04g34130*) was removed from further analysis because the sequence was identical to that of the *Escherichia coli* *ECs4062* gene, identified through the NCBI NR database (Fig. S1), implying possible sequence

contamination in the Nipponbare genome. Therefore after exclusion, a total of 37 orphan genes were identified in the Nipponbare genome (Table 1).

Syntenic information was further used to verify results of the gene homology searches. Due to the fact that orphan genes inherently lack phylogenetic conservation, the “flanking gene method” (Freeling et al., 2008) was used to deduce the syntenic locations of the orphan genes in the close relatives.

Twenty-seven orphan genes were anchored to corresponding syntenic regions in at least one close relative (Table S3). In these cases, all the syntenic regions in the close relatives exhibited continuous sequences without any gaps, excluding the improbable orphan genes attributed to poor genome assemblies in the outgroup. Furthermore, no intact open reading frames (ORFs) were detected in the syntenic regions, suggesting these orphan genes were truly absent even in the close relatives.

Analysis of the functionality of orphan genes provides evidence for its authenticity. Functionality of all 37 identified orphan genes was supported by at

least one piece of evidence, either ESTs, FL-cDNA sequences, microarray data, RNA-seq experimental data, or Interpro Protein Domain Alignments (Tables 1, S4). We found that 35 orphan genes contain intrinsically disordered protein domains by MobiDB-lite (Necci et al., 2017) analysis at the InterPro website (Table S5). We also successfully validated expression patterns of two orphan genes from among eight randomly selected orphan genes via RT-PCR experiments (Fig. S2).

3.2 Orphan genes exhibited genic features distinct from those of Nonorphan genes

To investigate whether the identified orphan genes had distinct properties, we compared three genic features (protein length, intron number, and GC content) between the orphan genes and nonorphan genes. Peptide lengths of the orphan genes were significantly shorter than those of the nonorphan genes (Fig. 2A). The median length of the nonorphan proteins (336 amino acids) is approximately three times longer than that of the orphan proteins (105 amino acids) (Mann-Whitney test, p -value = $6.458e-13$). Orphan genes contained fewer introns (with a median

of 0) than did the nonorphan genes (with a median of 2) (Mann-Whitney test, p -value = $6.458e-13$) (Fig. 2B). The GC content at the CDS level of the orphan genes (with a median of 74.23%) was notably higher than that of the nonorphan genes (with a median of 54.10%) (Mann-Whitney test, p -value= $2.585e-14$) (Fig. 2C).

3.3 The majority of orphan genes originated from a TE-mediated mechanism followed by rapid divergence

To obtain clues about the underlying processes involved in the birth of orphan genes, we compared multiple sequences in the syntenic regions of Nipponbare and of its three close relatives using the MUSCLE program (Edgar, 2004). We did not detect any homologous sequences in the close relatives for 19 Nipponbare orphan genes. In order to trace origin mechanisms for these 19 untraceable orphan genes, we searched for the TEs in the syntenic regions of the genomes of Nipponbare and of its close relatives. These searches were motivated by previous observations which suggest orphan genes are derived from TEs in primates (Toll-Riera et al., 2009) and in silkworms (Sun et al., 2015).

Determination of TE-derived sequences was the key evidence suggesting a TE-mediated origin of the orphan genes. We checked whether the TEs were intact with complete target site duplications (TSDs) in the syntenic regions. A typical example (*LOC_Os01g72920*) of how an orphan gene could be formed from TEs is shown in Fig. 3. *LOC_Os01g72920* is an orphan gene located on chromosome 1: 42288473–42289606, where it is embedded in an LTR-18C_OS-LTR (RepBase ID) retrotransposon. We further confirmed that this retrotransposon was recently inserted into the Nipponbare genome with “TTATG” as the TSD sequence, as it was not found in the syntenic regions of three close relatives (*O. sativa* ssp. *indica* ‘Shuhui 498’, and *O. nivara*). Sequences around the TE were conserved among Nipponbare and its close relatives. Therefore, specifically inserting LTR-18C_OS-LTR retrotransposon resulted in the formation of the orphan gene *LOC_Os01g72920*. The first exon of *LOC_Os01g72920* overlapped with the LTR-18C_OS-LTR retrotransposon, and the second exon combined the TSD sequence “TTATG” with the retrotransposon sequence.

In total, 19 of the identified Nipponbare orphan genes (51%) were derived from TE insertion events. TE-mediated orphan genes can be sorted into two groups (Fig. 4A): Fourteen orphan genes were generated by recent TE insertion events that only occurred in the Nipponbare genome (e.g., *LOC_Os01g72920*) (Figs. 4A, S3A-S3I, S3P-S3R; Table 2), and five orphan genes were formed by ancient TE insertion events when TEs were inserted in the common ancestor of Nipponbare and its close relatives (Figs. 4A, S3J-S3M, S3O; Table 2).

Eleven different TEs participated in the origin of 19 orphan genes (Table 2).

TE expansion can duplicate orphan genes after its origin. If an orphan gene duplicated through TEs, we hypothesized that we should be able to detect associated gene paralogs, and all those paralogs should be related to one TE type. We found nine orphan genes associated to three TEs that fit this hypothesis. In detail, five orphan genes (*LOC_Os04g11940*, *LOC_Os05g48540*, *LOC_Os08g36270*, *LOC_Os08g44980*, and *LOC_Os11g30450*) were paralogs (Fig S4). All paralogs appear to be associated with ENSPM4 (Table 2), so we

speculated that these five genes formed by ENSPM4 expansion, after one had initially originated. Similarly, *LOC_Os05g42940* and *LOC_Os06g16530* appear to have formed through SPMLIKE transposon expansion (Fig S5), and *LOC_Os08g09680* and *LOC_Os11g35600* appear to have formed through SZ-67LTR expansion (Fig S6). Although three orphan genes (*LOC_Os06g51300*, *LOC_Os08g26960* and *LOC_Os07g26890*) appear to be related to SZ-67LTR insertion, they have completely different coding sequences. This suggests that these three orphan genes originated independently after TE insertion. In summary, 13 independent origin events and 3 TE expansion events account for 19 Nipponbare orphan genes (Table 2).

3.4 *De novo* origination served as another primary mechanism for orphan gene formation

The identification of noncoding orthologous sequences in the syntenic region of close relatives can provide strong evidence for *de novo* origination. The power of sequence similarity searches depends on the size of the query genes (Ruiz-Orera & Mar Albà, 2018). As a result, orthologous sequences of short genes

may be missed when using Megablast to rule out similar hits in the genomes of close relatives. By comparing multiple sequences of syntenic regions in Nipponbare and close relatives via the MUSCLE program, highly divergent or small homologous non-coding sequences can be identified in the close relatives.

In this study, eight orphan genes in Nipponbare (22%) were confidently defined as *de novo* origination events because noncoding orthologous sequences existed in the close relatives. During the process of *de novo* origination, it is crucial to obtain a start codon and to remove internal stop codons via mutations, such as indels (i.e., frameshift) and point mutations. For instance, noncoding sequences in Nipponbare were transformed into the orphan gene *LOC_Os04g22510* via three critical enabling mutations: 1) start codon acquisition; 2) insertion of a single “T” at base pair position 46, which resulted in a frameshift that removed a stop codon at base pair position 50; and 3) conversion of a premature “TAG” stop codon into “TCG” (which encodes serine) at base pair positions 293–295 via a point mutation (Fig. S7). In summary, of the eight *de*

novo-formed orphan genes, four appear to have formed through start codon acquisition, and indel and point mutations; two appear to have formed via start codon acquisition and indels; one appears to have formed via start codon acquisition alone, and one via enabling indels alone (Table 3).

4 Discussion

In this study, we identified 37 recently originated orphan genes in the Nipponbare genome (Table 1). Orphan genes are sometimes considered as *de novo* genes originating from ancestral noncoding sequences (Chen et al., 1997; Knowles & McLysaght, 2009; Li et al., 2010; Murphy & McLysaght, 2012; Xie et al., 2012). In fact, *de novo* formation is just one way by which orphan genes can form, as many orphan genes are derived from other distinct evolutionary processes, such as the duplication-divergence mechanism (Schlotterer, 2015; Moyers & Zhang, 2016), TE exaptation (Toll-Riera et al., 2009), loss of homologous genes in related species (Zhao et al., 2015), repetition of low-complexity short peptides (Chen et al., 1997; Cheng & Chen, 1999), and horizontal gene transfer from fast-evolving donors (Keeling & Palmer, 2008;

Husnik & McCutcheon, 2018). An orphan gene can also originate through a combination of several origin mechanisms, such as the mixed origin mechanism in nematodes (Prabh & Rödelsperger, 2019). Zhang et al. (2019) recently reported that *de novo* genes contributed to the rapid evolution of *Oryza* protein diversity over 15 myr. In order to provide more details about the origination of orphan genes, we investigated the origins of the youngest orphan genes in the model plant *O. sativa* ssp. *japonica* 'Nipponbare' (< 0.55 myr).

Identification pipelines and genome annotation may affect the number of orphan genes identified. In this study, because we utilized a strict filtering procedure (excluding genes that have any MEGABLAST hits in the outgroup) to identify Nipponbare orphan genes, we exclude the *de novo* gene reported in Zhang et al. (2019) which highlighted highly similarity in non-coding sequences across the relatives. The strict filtering procedure used in our study could lead to an underestimation in the number of *de novo* genes. Genome annotation of focal genome is another important factor which might affect the identification of orphan

genes (Denton et al., 2014; Prabh & Rödelsperger, 2016). For instance, among 37 Nipponbare orphan genes in our analysis (MSU-RAP annotation), we found that only eight orphan genes (22%) were formed by de novo, but these genes were not annotated in Zhang et al. (2019) study (OGE/IOMAP annotation).

In our study, the majority of the orphan genes (51%) were derived from TE-related sequences, indicating that TE could serve as the main genetic material for the origin of orphan genes in the Nipponbare genome. We identified a distinctive TE-mediated mechanism which differs from previously reported mechanisms involved in the birth of orphan genes. Previous studies indicated that TEs can contribute to novel genes formation via three distinct mechanisms. First, segments of TE sequences could be recruited as dependent exons or as parts of exons in host genes via a processes, known as, TE exonization or TE exaptation (Fig. 4B) (Nekrutenko & Li, 2001; Long et al., 2003; Shapiro, 2005; Chen et al., 2013; Long et al., 2013). Second, several TEs, such as MULEs, can capture host genes or gene fragments, such as pack MULE model in the Jiang et al. (2004),

referring to as “TE pack duplication” (Fig. 4C). This TE-mediated mechanism often generates duplicated genes, and some captured gene fragments which can become functional (Jiang et al. 2004; Flagel & Wendel, 2009; Panchy et al., 2016).

Third, conserved TE sequences, such as those encoding transposases, can be domesticated via evolution to harbor specific traits in a process called TE domestication (Fig. 4D) (Kapitonov & Jurka, 2005; Jangam et al., 2017). However, none of these TE-related mechanisms are absolutely consistent with our results.

In our analysis, orphan genes could origin independently through rapid divergence, after the TE insertions, a process that we consider independent mutations within inserted TEs (Fig. 4E). Ten Nipponbare orphan genes had no identifiable paralogs in the genome and appear to be associated with different TEs, which indicates that these orphan genes formed independently after the TE insertions. But once an orphan gene is formed, it will be duplicated through TE expansion (e.g., TE pack duplication mechanism), for example, nine orphan genes were formed by three TEs (ENSPM4, SPMLIKE, and SZ-67LTR) expanded,

similar to TE pack duplication process (Fig. 4C). In these nine genes, it is difficult to distinguish which three are parent genes because they have no comparative homologs in the close relatives.

The identification of independent origin through rapid divergence within inserted TEs that explain Nipponbare orphan gene origination was based on a short-timescale analysis of TE insertion events in the target taxon and its closely related species. Previous studies on orphan genes have mainly focused on detecting TE-related orphan gene sequences in target taxon/taxa by BLAST searches against a TE database (Toll-Riera, et al., 2009; Donoghue et al., 2011; Wissler et al., 2013; Yang et al., 2013; Sun et al., 2015), without comparisons to its close relatives. Therefore, these analysis failed to reveal a detailed picture of the TE insertions or the emergence of TE-related orphan genes. In addition to searching for intact transposons in the syntenic regions of the focus genome and those of its close relatives, we also speculated on the relative times of TE insertion events and the origins of the orphan genes. This novel analysis led us to infer that

the orphan genes can form independently through rapid divergence in the inserted TEs. The fact that the orphan genes originated from independent divergence within inserted TE. The origination of the orphan genes from independent divergence after TE insertion is unlikely to be Nipponbare specific. This mechanism might also be present in other taxa that have not yet been characterized.

The emergence of orphan genes from TE-related sequences may be supported by the high TE mutation rates. Historically, TEs were considered "junk" sequences in genomes (Ohno, 1972). However, TEs are highly mutagenic due to their high genomic abundance, which can easily promote chromosomal rearrangements (Schrader & Schmitz, 2019). TE insertions can drastically affect the evolution of the surrounding genes by altering their genetic structure and/or regulatory sequences. Most TEs remain silent, evolving in a neutral fashion, however, a proportion appear to gain adaptive roles (Arkhipova, 2018). TE-derived proteins have been abundantly and recurrently domesticated, and they

are now considered an important adaptive mechanism for evolutionary innovation (Jangam et al., 2017; Schrader & Schmitz, 2019). TEs have very high mutation rates, a trait that has been confirmed in the model plant *A. thaliana* at the population level (Li et al., 2018), which is also consistent with the previous hypothesis of that high mutation rate will present along with insertion (Tian et al., 2008)

The most prevalent TE-associated mechanism for orphan gene formation in Nipponbare (51%) is comparable to the prevalence of this mechanism in primates (53%) (Toll-Riera et al., 2009). Nevertheless, the frequency of the TE-associated mechanism for orphan genes formation in the Nipponbare genome is substantially higher than that in model plant *A. thaliana* (9.73%) (Donoghue et al., 2011). This difference could be due to the relative abundances of TE content in their genomes. The TE content in the *A. thaliana* genome is only 10% (*Arabidopsis* Genome Initiative, 2000) but reaches up to 48.6% in the Nipponbare genome (International Rice Genome Sequencing, 2005). The conclusion that a TE-mediated mechanism

is the dominant driver of Nipponbare orphan gene formation is indeed inconsistent with the conclusions of previous studies on *Drosophila* (Zhou et al., 2008) and *A. thaliana* (Donoghue et al., 2011). These studies indicated that duplication-divergence was the primary mechanism for orphan gene formation based on comparisons with distant reference genomes. However, duplication require much long evolutionary time scales to accumulate sufficient nucleotide substitutions. Thus, this model was insufficient to explain the youngest genes formed on a short time scale which we identified in Nipponbare genome (< 0.55 myr). The dominance of different mechanisms for orphan gene origination in various species suggests that different mechanisms might play different roles in each species.

Shorter protein length and fewer introns in the Nipponbare orphan genes were consistent with the findings of studies on orphan genes in both animals (Zhang et al., 2007; Murphy & McLysaght, 2012; Wissler et al., 2013; Yang et al., 2013; Palmieri et al., 2014; Mayer et al., 2015) and plants (Lin et al., 2010; Xu et

al., 2015). This suggest that these two characteristics are conserved among the orphan genes in all eukaryotes. The elevated GC content of the Nipponbare orphan genes was consistent with previous reports in plants such as *A. thaliana* (Arendsee et al., 2014), *C. sinensis* (Xu et al., 2015), and Poaceae (Campbell et al., 2007). High GC content could cause intrinsically disordered proteins (Basile et al., 2017). In our study, we also found that almost all (95%) Nipponbare orphan genes have the disordered protein properties, which in agreement with previous results (Bornberg-Bauer et al., 2015; Wilson et al., 2017). Short protein lengths and high GC content made it difficult for us to design primers and to perform RT-PCR experiments. Even though we get some expression evidences, it remains unclear which orphan gene began to function after origination because nonfunctional proteins are tend to pervasively transcript and translate (Ruiz-Orera & Mar Albà, 2018; Prabh & Rödelsperger, 2019). Although future studies will be required to determine which orphan genes are fixed in the population and associated with species-specific functions, our finding provide clear insight into genetic innovations in the rice genome.

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References

- Arabidopsis* Genome Initiative. 2000. Analysis of the genome sequence of the flowering plant *Arabidopsis thaliana*. *Nature* 408: 796.
- Arendsee ZW, Li L, Wurtele ES. 2014. Coming of age: Orphan genes in plants. *Trends in Plant Science* 19: 698-708.
- Arkhipova IR. 2018. Neutral theory, transposable elements, and eukaryotic genome evolution. *Molecular Biology and Evolution* 35: 1332-1337.
- Basile W, Sachenkova O, Light S, Elofsson A. 2017. High GC content causes orphan proteins to be intrinsically disordered. *PLoS Computational Biology* 13: e1005375.

Bauer DF. 1972. Constructing confidence sets using rank statistics. *Journal of the*

American Statistical Association 67: 687-690.

Boffelli D, Nobrega MA, Rubin EM. 2004. Comparative genomics at the

vertebrate extremes. *Nature Reviews Genetics* 5: 456-465.

Bornberg-Bauer E, Schmitz J, Heberlein M. 2015. Emergence of de novo proteins

from 'dark genomic matter' by 'grow slow and moult'. *Biochemical Society*

Transactions 43: 867-873.

Cai JJ, Woo PCY, Lau SKP, Smith DK, Yuen KY. 2006. Accelerated evolutionary

rate may be responsible for the emergence of lineage-specific genes in

ascomycota. *Journal of Molecular Evolution* 63: 1-11.

Campbell MA, Zhu W, Jiang N, Lin H, Ouyang S, Childs KL, Haas BJ, Hamilton

JP, Buell CR. 2007. Identification and characterization of lineage-specific

genes within the Poaceae. *Plant Physiology* 145: 1311-1322.

Chen LB, DeVries AL, Cheng HC. 1997. Evolution of antifreeze glycoprotein

gene from a trypsinogen gene in Antarctic notothenioid fish. *Proceedings of the National Academy of Sciences USA* 94: 3811-3816.

Chen SD, Krinsky BH, Long MY. 2013. New genes as drivers of phenotypic evolution. *Nature Reviews Genetics* 14: 745-745.

Cheng CHC, Chen LB. 1999. Evolution of an antifreeze glycoprotein. *Nature* 401: 443-444.

Daubin V, Ochman H. 2004. Bacterial genomes as new gene homes: The genealogy of ORFans in *E-coli*. *Genome Research* 14: 1036-1042.

Denton JF, Lugo-Martinez J, Tucker AE, Schrider DR, Warren WC, Hahn MW. 2014. Extensive error in the number of genes inferred from draft genome assemblies. *PLoS Computational Biology* 10: e1003998.

Domazet-Loso T, Tautz D. 2003. An evolutionary analysis of orphan genes in *Drosophila*. *Genome Research* 13: 2213-2219.

Donoghue MTA, Keshavaiah C, Swamidatta SH, Spillane C. 2011. Evolutionary

origins of Brassicaceae specific genes in *Arabidopsis thaliana*. *BMC*

Evolutionary Biology 11: 47.

Du H, Yu Y, Ma Y, Gao Q, Cao Y, Chen Z, Ma B, Qi M, Li Y, Zhao X, Wang J,

Liu K, Qin P, Yang X, Zhu L, Li S, Liang C. 2017. Sequencing and *de novo*

assembly of a near complete indica rice genome. *Nature Communications* 8:

15324.

Edgar RC. 2004. MUSCLE: Multiple sequence alignment with high accuracy and

high throughput. *Nucleic Acids Research* 32: 1792-1797.

Flagel LE, Wendel JF. 2009. Gene duplication and evolutionary novelty in plants.

New Phytologist 183: 557-564.

Fischer D, Eisenberg D. 1999. Finding families for genomic ORFans.

Bioinformatics 15: 759-762.

Freeling M, Lyons E, Pedersen B, Alam M, Ming R, Lisch D. 2008. Many or most

genes in *Arabidopsis* transposed after the origin of the order Brassicales.

Genome Research 18:1924-1937.

Husnik F, McCutcheon JP. 2018. Functional horizontal gene transfer from bacteria to eukaryotes. *Nature Reviews Microbiology* 16: 67-79.

Jangam D, Feschotte C, Betran E. 2017. Transposable element domestication as an adaptation to evolutionary conflicts. *Trends in Genetics* 33: 817-831.

Jiang N, Bao ZR, Zhang XY, Eddy SR, Wessler SR. 2004. Pack-mule transposable elements mediate gene evolution in plants. *Nature* 431: 569-573.

Johnson BR, Tsutsui ND. 2011. Taxonomically restricted genes are associated with the evolution of sociality in the honey bee. *BMC Genomics* 12: 164.

Jones DC, Zheng WG, Huang S, Du CL, Zhao XF, Yennamalli RM, Sen TZ, Nettleton D, Wurtele ES, Li L. 2016. A clade-specific *Arabidopsis* gene connects primary metabolism and senescence. *Frontiers in Plant Science* 7: 983.

Kapitonov VV, Jurka J. 2005. RAG1 core and V(D)J recombination signal

sequences were derived from Transib transposons. *PLoS Biology* 3:

998-1011.

Kawahara Y, de la Bastide M, Hamilton JP, Kanamori H, McCombie WR, Ouyang

S, Schwartz DC, Tanaka T, Wu JZ, Zhou SG, Childs KL, Davidson RM, Lin

HN, Quesada-Ocampo L, Vaillancourt B, Sakai H, Lee SS, Kim J, Numa H,

Itoh T, Buell CR, Matsumoto T. 2013. Improvement of the *Oryza sativa*

Nipponbare reference genome using next generation sequence and optical

map data. *Rice* 6: 4.

Keeling PJ, Palmer JD. 2008. Horizontal gene transfer in eukaryotic evolution.

Nature Reviews Genetics 9: 605-618.

Kent WJ. 2002. BLAT—the BLAST-like alignment tool. *Genome Research* 12:

656-664.

Khalturin K, Hemmrich G, Fraune S, Augustin R, Bosch TCG. 2009. More than

just orphans: are taxonomically-restricted genes important in evolution?

Trends in Plant Science 25: 404-413.

This article is protected by copyright. All rights reserved.

Knowles DG, McLysaght A. 2009. Recent *de novo* origin of human protein-coding genes. *Genome Research* 19: 1752-1759.

Kohany O, Gentles AJ, Hankus L, Jurka J. 2006. Annotation, submission and screening of repetitive elements in Repbase: RepbaseSubmitter and Censor. *BMC Bioinformatics* 7: 474.

Li CY, Zhang Y, Wang Z, Zhang Y, Cao C, Zhang PW, Lu SJ, Li XM, Yu Q, Zheng X, Du Q, Uhl GR, Liu QR, Wei L. 2010. A human-specific *de novo* protein-coding gene associated with human brain functions. *PLoS Computational Biology* 6: e1000734.

Li ZW, Hou XH, Chen JF, Xu YC, Wu Q, Gonzalez J, Guo YL. 2018. Transposable elements contribute to the adaptation of *Arabidopsis thaliana*. *Genome Biology and Evolution* 10: 2140-2150.

Lin HN, Moghe G, Ouyang S, Iezzoni A, Shiu SH, Gu X, Buell CR. 2010. Comparative analyses reveal distinct sets of lineage-specific genes within *Arabidopsis thaliana*. *BMC Evolutionary Biology* 10: 41.

This article is protected by copyright. All rights reserved.

Lipman DJ, Souvorov A, Koonin EV, Panchenko AR, Tatusova TA. 2002. The relationship of protein conservation and sequence length. *BMC Evolutionary Biology* 2: 20.

Long M, Betran E, Thornton K, Wang W. 2003. The origin of new genes: Glimpses from the young and old. *Nature Reviews Genetics* 4: 865-875.

Long MY, VanKuren NW, Chen SD, Vibranovski MD. 2013. New gene evolution: Little did we know. *Annual Review of Genetics* 47: 307-333.

Mayer MG, Rodelsperger C, Witte H, Riebesell M, Sommer RJ. 2015. The orphan gene *dauerless* regulates dauer development and intraspecific competition in nematodes by copy number variation. *PLoS Genetics* 11: e1005146.

Moyers BA, Zhang JZ. 2016. Evaluating phylostratigraphic evidence for widespread *de novo* gene birth in genome evolution. *Molecular Biology and Evolution* 33: 1245-1256.

Murphy DN, McLysaght A. 2012. *De novo* origin of protein-coding genes in

murine rodents. *PLoS One* 7: e48650.

Nekrutenko A, Li WHS. 2001. Transposable elements are found in a large number of human protein-coding genes. *Trends in Genetics* 17: 619-621.

Ohno S. 1970. *Evolution by gene duplication*. Berlin Heidelberg: Springer-Verlag.

Palmieri N, Kosiol C, Schlotterer C. 2014. The life cycle of *Drosophila* orphan genes. *Elife* 3: e01311.

Panchy N, Lehti-Shiu M, Shiu SH. 2016. Evolution of gene duplication in plants. *Plant Physiology* 171: 2294-2316.

Perochon A, Jia JG, Kahla A, Arunachalam C, Scofield SR, Bowden S, Wallington E, Doohan FM. 2015. TaFROG encodes a Pooideae orphan protein that interacts with SnRK1 and enhances resistance to the mycotoxigenic fungus *Fusarium graminearum*. *Plant Physiology* 169: 2895-2906.

Pontius JU, Wagner L, Schuler GD. 2003. UniGene: A unified view of the transcriptome. In: Bethesda MD editors. *National Center for Biotechnology*

Information [online]. Available from

<http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=books>.

Prabh N, Rödelsperger C. 2016. Are orphan genes protein-coding, prediction artifacts, or non-coding RNAs? *BMC Bioinformatics* 17: 226.

Prabh N, Rödelsperger C. 2019. De novo, divergence, and mixed origin contribute to the emergence of orphan genes in *Pristionchus* nematodes. *G3: Genes, Genomes, Genetics* g3: 400326.

Ruiz-Orera J, Verdaguer-Grau P, Villanueva-Cañas JL, Messeguer X, Albà MM. 2018. Translation of neutrally evolving peptides provides a basis for de novo gene evolution. *Nature Ecology & Evolution* 2: 890.

Schlotterer C. 2015. Genes from scratch - the evolutionary fate of *de novo* genes. *Trends in Genetics* 31: 215-219.

Schrader L, Schmitz J. 2019. The impact of transposable elements in adaptive evolution. *Molecular Ecology* 28: 1537-1549.

This article is protected by copyright. All rights reserved.

Shapiro JA. 2005. A 21st century view of evolution: Genome system architecture, repetitive DNA, and natural genetic engineering. *Gene* 345: 91-100.

Stein JC, Yu Y, Copetti D, Zwickl DJ, Zhang L, Zhang C, Chougule K, Gao D, Iwata A, Goicoechea JL, Wei S, Wang J, Liao Y, Wang M, Jacquemin J, Becker C, Kudrna D, Zhang J, Londono CEM, Song X, Lee S, Sanchez P, Zuccolo A, Ammiraju JSS, Talag J, Danowitz A, Rivera LF, Gschwend AR, Noutsos C, Wu CC, Kao SM, Zeng JW, Wei FJ, Zhao Q, Feng Q, El Baidouri M, Carpentier MC, Lasserre E, Cooke R, Rosa Farias DD, da Maia LC, Dos Santos RS, Nyberg KG, McNally KL, Mauleon R, Alexandrov N, Schmutz J, Flowers D, Fan C, Weigel D, Jena KK, Wicker T, Chen M, Han B, Henry R, Hsing YC, Kurata N, de Oliveira AC, Panaud O, Jackson SA, Machado CA, Sanderson MJ, Long M, Ware D, Wing RA. 2018. Genomes of 13 domesticated and wild rice relatives highlight genetic conservation, turnover and innovation across the genus *Oryza*. *Nature Genetics* 50: 285-296.

-
- Sun W, Zhao XW, Zhang Z. 2015. Identification and evolution of the orphan genes in the domestic silkworm, *bombyx mori*. *Febs Letters* 589: 2731-2738.
- Tautz D, Domazet-Loso T. 2011. The evolutionary origin of orphan genes. *Nature Reviews Genetics* 12: 692-702.
- Tian DC, Wang Q, Zhang PF, Araki H, Yang SH, Kreitman M, Nagylaki T, Hudson R, Bergelson J, Chen JQ. 2008. Single-nucleotide mutation rate increases close to insertions/deletions in eukaryotes. *Nature* 455: 105-108.
- Toll-Riera M, Bosch N, Bellora N, Castelo R, Armengol L, Estivill X, Alba MM. 2009. Origin of primate orphan genes: A comparative genomics approach. *Molecular Biology and Evolution* 26: 603-612.
- Wilson BA, Foy SG, Neme R, Masel J. 2017. Young genes are highly disordered as predicted by the preadaptation hypothesis of de novo gene birth. *Nature Ecology & Evolution* 1: 0146.
- Wissler L, Gadau J, Simola DF, Helmkamp M, Bornberg-Bauer E. 2013.

Mechanisms and dynamics of orphan gene emergence in insect genomes.

Genome Biology and Evolution 5: 439-455.

Xie C, Zhang YE, Chen JY, Liu CJ, Zhou WZ, Li Y, Zhang M, Zhang R, Wei L, Li

CY. 2012. Hominoid-specific *de novo* protein-coding genes originating from long non-coding RNAs. *PLoS Genetics* 8: e1002942.

Xu YT, Wu GZ, Hao BH, Chen LL, Deng XX, Xu Q. 2015. Identification,

characterization and expression analysis of lineage-specific genes within sweet orange (*Citrus sinensis*). *BMC Genomics* 16: 995.

Yang LD, Zou M, Fu BD, He SP. 2013. Genome-wide identification,

characterization, and expression analysis of lineage-specific genes within zebrafish. *BMC Genomics* 14: 65.

Zhang G, Wang H, Shi J, Wang X, Zheng H, Wong GK, Clark T, Wang W, Wang

J, Kang L. 2007. Identification and characterization of insect-specific proteins by genome data analysis. *BMC Genomics* 8: 93.

Zhang L, Ren Y, Yang T, Li GW, Chen JH, Gschwend AR, Yu Y, Hou GX, Zi J,

Zhou R, Wen B, Zhang JW, Chougule K, Wang MH, Copetti D, Peng ZY,

Zhang CJ, Zhang Y, Ouyang YD, Wing RA, Liu SQ, Long MY. 2019. Rapid

evolution of protein diversity by *de novo* origination in *Oryza*. *Nature*

Ecology & Evolution 3: 679-690.

Zhao Y, Tang L, Li Z, Jin JP, Luo JC, Gao G. 2015. Identification and analysis of

unitary loss of long-established protein-coding genes in Poaceae shows

evidences for biased gene loss and putatively functional transcription of

relics. *BMC Evolutionary Biology* 15: 66.

Zhang YE, Landback P, Vibranovski MD, Long M. 2011. Accelerated recruitment

of new brain development genes into the human genome. *PLoS Biology* 9:

e1001179.

Zhou Q, Zhang G, Zhang Y, Xu S, Zhao R, Zhan Z, Li X, Ding Y, Yang S, Wang

W. 2008. On the origin of new genes in *Drosophila*. *Genome research* 18:

1446-1455.

Tables
Table 1 Information on orphan genes in the Nipponbare genome

Orphan gene	Interpro Domain	FL-cDN A	ESTs	RNA-se q	Microarra y	RT-PCR
LOC_Os01g505 60	+			+		
LOC_Os01g729 20	+					
LOC_Os03g178 30	+					
LOC_Os03g604 19	+	+	+	+		
LOC_Os04g119 40	+			+		
LOC_Os04g225 10	+			+	+	+
LOC_Os05g429 40	+			+		
LOC_Os05g466 50	+			+		
LOC_Os05g485 40	+					
LOC_Os06g165 30	+			+		

LOC_Os06g198 80	+			+
LOC_Os06g339 10	+			
LOC_Os06g381 90				+
LOC_Os06g443 90	+			+
LOC_Os06g513 00	+	+		+
LOC_Os07g262 40	+			+
LOC_Os07g267 70	+			
LOC_Os07g268 90	+			+
LOC_Os08g053 30	+			+
LOC_Os08g096 80	+			
LOC_Os08g264 60	+		+	+
LOC_Os08g269 60	+			
LOC_Os08g362 70	+			
LOC_Os08g449	+			+

80

LOC_Os09g132 60	+		+	+	
LOC_Os09g356 40	+		+		
LOC_Os10g095 60	+		+	+	
LOC_Os11g069 50	+		+		
LOC_Os11g304 50	+		+		
LOC_Os11g356 00	+		+		
LOC_Os11g442 00	+		+	+	
LOC_Os11g442 70	+		+	+	
LOC_Os11g442 80			+	+	
LOC_Os12g098 80	+		+	+	
LOC_Os12g110 60	+		+	+	+
LOC_Os12g332 50	+		+		
LOC_Os12g432 00	+		+		

+, evidence present.

Table 2 Origin of orphan genes in Nipponbare mediated by transposable element insertion

Orphan genes	TE Class	TE IDs in RepBase	TE insertion time	Origin event
LOC_Os04g11940	DNA/CACTA	ENSPM4	recent	1
LOC_Os05g48540				
LOC_Os08g36270				
LOC_Os08g44980				
LOC_Os11g30450				
LOC_Os05g42940	DNA/CACTA	SPMLIKE	recent	1
LOC_Os06g16530				
LOC_Os08g09680	RNA/LTR	SZ-67LTR	recent	1
LOC_Os11g35600				
LOC_Os06g198	DNA/CACTA	SPMLIKE-B_O	recent	1

80	A	S		
LOC_Os01g729 20	RNA/LTR	LTR-18C_OS-L TR	recent	1
LOC_Os05g466 50	RNA/LTR	LTR-18_OS-LT R	recent	1
LOC_Os06g443 90	RNA/LTR	MuDR-N18C_O S	recent	1
LOC_Os08g264 60	RNA/LTR, LINE	COPIA1-LTR_ OS, LINE1-11_OS	recent	1
LOC_Os11g069 50	RNA/LTR	LTR-18B_OS-L TR	ancient	1
LOC_Os12g332 50	RNA/LTR	LTR-18K_OS-L TR	ancient	1
LOC_Os06g513 00	RNA/LTR	SZ-67LTR	ancient	1
LOC_Os07g268 90	RNA/LTR	SZ-67LTR	ancient	1
LOC_Os08g269 60	RNA/LTR	SZ-67LTR	ancient	1

Table 3 Crucial mutations that transformed noncoding sequences into coding sequences in the *de novo* process

Orphan gene	Start codon acquisition	Enabling indels	Enabling point mutations
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LOC_Os03g178 30	+	+	+
LOC_Os03g604 19	+	+	+
LOC_Os04g225 10	+	+	+
LOC_Os06g339 10	+	+	-
LOC_Os08g053 30	+	-	-
LOC_Os09g356 40	+	+	+
LOC_Os11g442 00	+	+	-
LOC_Os12g432 00	-	+	-

+, present; -, absent.

Figure legends

Fig. 1. The computational pipeline for the identification of orphan genes in Nipponbare based on similarity and synteny. The topology of the phylogeny referred to the published high-resolution tree in Stein et al. (2018). Orphan genes were identified in the Nipponbare genome using 10 other *Oryza* genomes as outgroups. The flanking genes of the Nipponbare orphan genes were used to detect syntenic regions in three close relatives, including *O. rufipogon*, *O. sativa* ssp. *indica* ‘Shuhui 498’, and *O. nivara*.

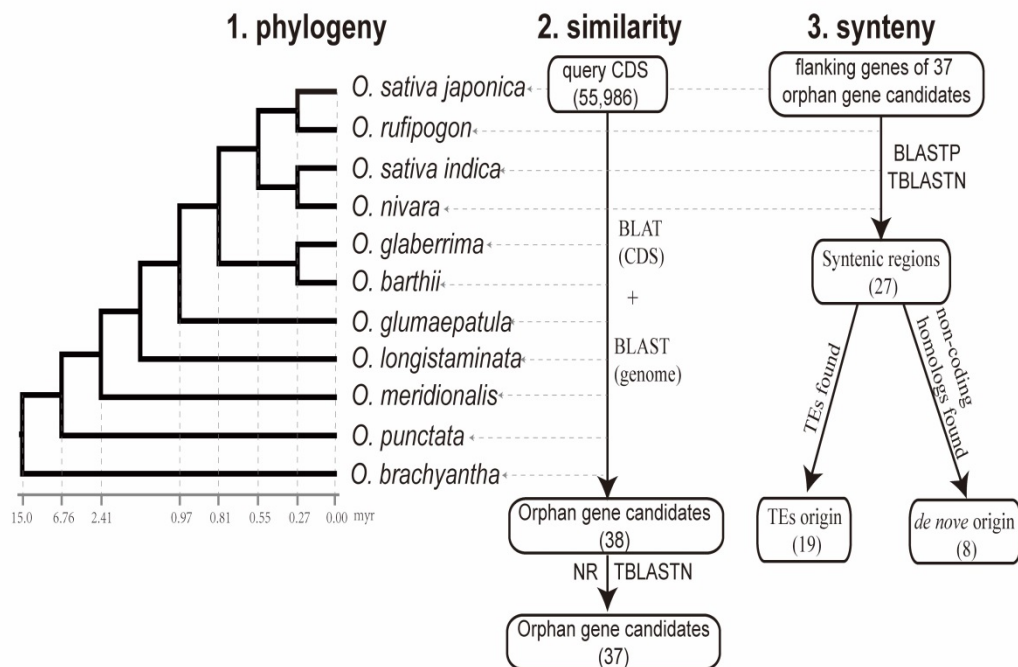


Fig. 2. Comparison of the genic features conserved between orphan genes and nonorphan genes in Nipponbare using a Wilcoxon rank sum test ($p < 0.05$). **A,** The protein-coding length of the orphan genes was much shorter than that of nonorphan genes. **B,** The intron number in orphan genes was significantly lower than that of nonorphan genes. **C,** The GC content of the orphan genes at the CDS level was significantly higher than that of nonorphan genes. Abbreviations: AAs, amino acids.

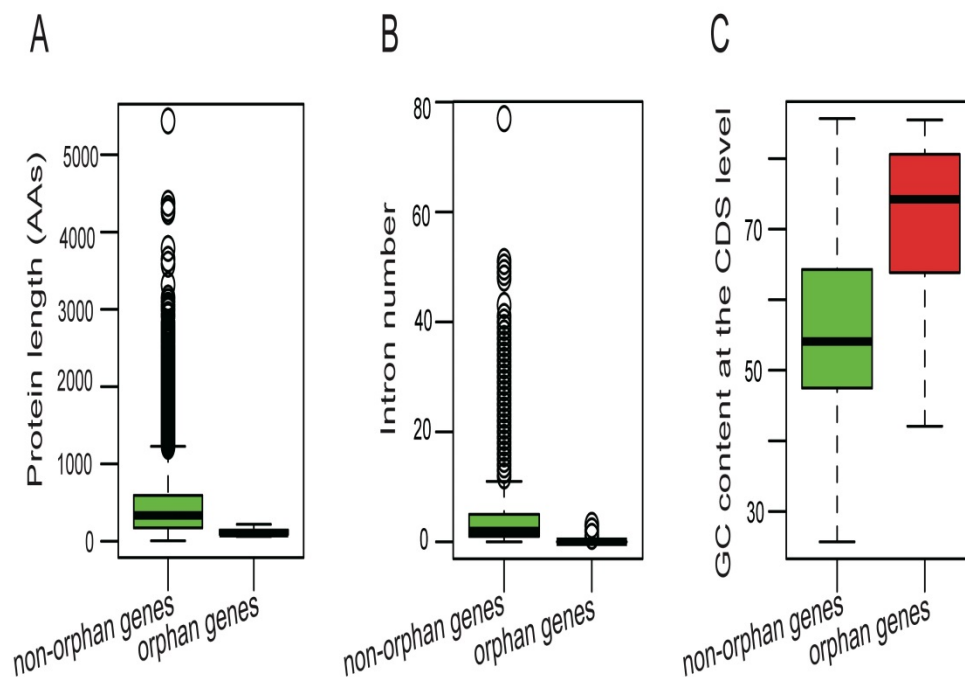


Fig. 3. Orphan gene LOC_Os01g72920 originated via a transposable element (TE)- mediated mechanism. Panel A shows an enlarged view of the 5' junction of the TE insertion. Panel B shows an enlarged view of the 3' junction of the TE insertion. Panel C illustrates the syntentic relationship between Nipponbare and its close relatives. The gray lines in the panel D represent syntentic chains among the four genomes. Boxes marked in the same color represent orthologous genes. The red box represents the orphan gene. Abbreviations: NIP, *O. sativa* ssp. *japonica* 'Nipponbare'; R498, *O. sativa* ssp. *indica* 'Shuhui 498'; NIVA, *O. nivara*.

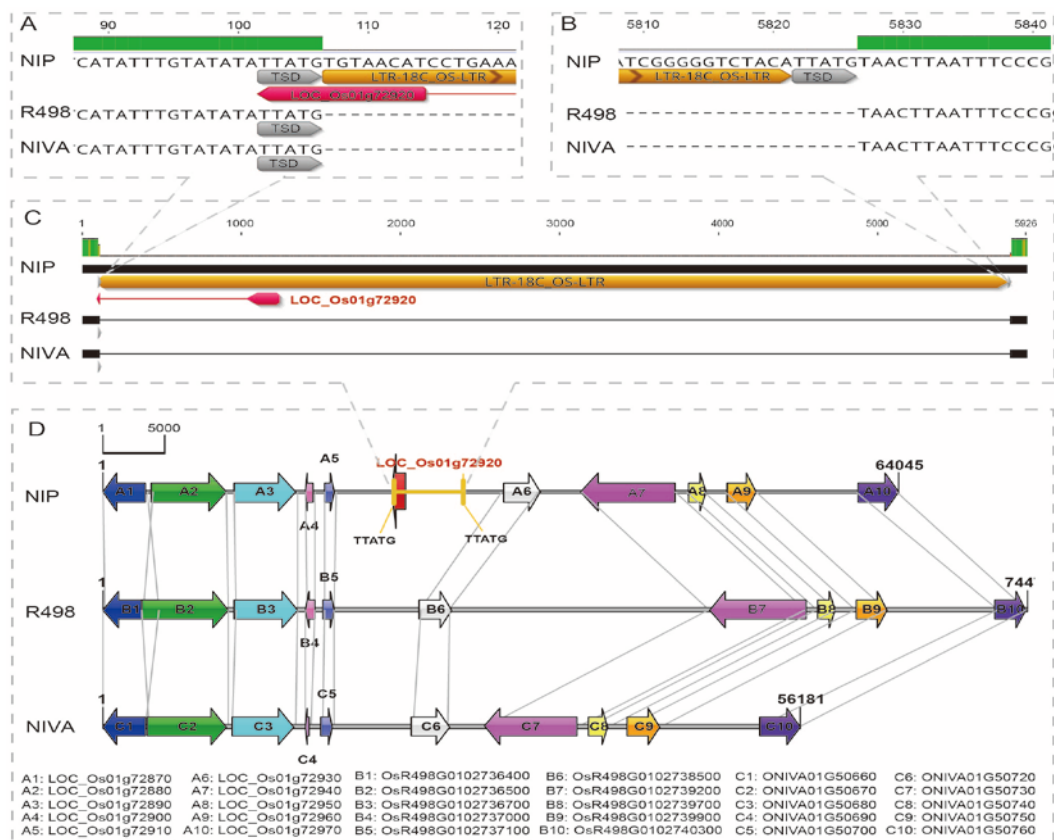


Fig. 4. Summary of new genes originated via the transposable element (TE)-mediated mechanism. **A,** Different TE insertion events associated with orphan genes in Nipponbare are shown on the branches of the phylogenetic tree. **B,** TE exonization, a host gene recruits a segment of a TE sequence as a dependent exon or as part of an exon of the host gene, also called TE exaptation, referring to the description of Chen et al. (2013). **C,** The pack duplication model: genes or gene fragments are captured by TEs to generate duplications, referring to the pack-MULE model of Jiang et al. (2004). **D,** TE domestication: conserved TE sequences, such as those encoding transposases, can be domesticated via evolution to harbor specific traits, referring to the description of Jangam et al. (2017). **E,** Independent origin through rapid divergence, orphan genes originate independently through rapid divergence within inserted TEs. Dashed line, empty position of a closely related outgroup in the alignment; Orange boxes/lines, TEs; Red boxes: orphan genes.

