INTRODUCTION

Most types of RNA modifications do not change the base-pairing property of the modified nucleotides, with the exceptions of A-to-I and C-to-U RNA editing. A-to-I RNA editing is prevalent and well studied in animal species while in the plant kingdom, C-to-U RNA editing is enriched in the chloroplast and mitochondrial genes (Gray, 2009; Gray & Covello, 1993; Mulligan, Williams, & Shanahan, 1999; Rajasekhar & Mulligan, 1993; Tang & Luo, 2018; Yu & Schuster, 1995). Other types of potentially recoding modifications include the opposite U-to-C substitution (Takenaka, Zehrmann, Verbitsky, Hartel, & Brennicke, 2013), which is much less frequent than C-to-U editing. The factor responsible for C-to-U RNA editing is PPR (pentatricopeptide repeat) protein, which recognizes single-strand RNAs (Yin et al., 2013).

The biological functions of particular C-to-U editing events were consecutively discovered (Takenaka, 2010; Takenaka, Neuwirt, & Brennicke, 2004; Takenaka, Verbitsky, Merwe, Zehrmann, & Brennicke, 2008; Tang, Kobayashi, Suzuki, Matsumoto, & Muranaka, 2010), and systematic identification of these editing sites appeared in recent years with the help of next-generation sequencing (NGS) technique. Bioinformatic tools are also developed to facilitate researchers to accurately identify C-to-U RNA editing sites from the big data (Du, Jia, & Li, 2009; Li et al., 2019; Lo Giudice, Hernandez,
2.1 | Data collection

The reference genome and CDS sequences of *Arabidopsis thaliana* were downloaded from TAIR database. The TAIR 9 version of annotation was used. The mRNA-sequencing data of *Arabidopsis thaliana* (three replicates for roots and shoots) were retrieved from a previous study (Hsu et al., 2016). As described by the paper, the *Arabidopsis* Columbia-0 (abbr. Col-0) seeds (around 1,500 seeds per vessel) were surface sterilized and imbibed at 4°C for 2 days. They were grown hydroponically on fine nylon mesh supported by a customized rack in Magenta vessel GA-7-3 (Sigma; V8380) with filtered sterile liquid media and shaken at 85 rpm under 16-hr light (110–115 μmol m⁻² s⁻¹ from cool white fluorescent bulbs) and 8-hr dark at 22°C (Hsu et al., 2016). The tissues were collected from 4-day old seedlings.

2.2 | NGS data processing

We mapped the mRNA-seq reads to the reference CDS sequence using Bowtie2 (version 2.2) (Langmead & Salzberg, 2012). The uniquely mapped reads were extracted using SAMtools (version 1.5) (Li et al., 2009). The variant calling process is accomplished by SAMtools “mpileup” with default parameters. The output file of mpileup is “vcf” format, which contains one variation site per line. The information for each variation site includes total depth on each site, the reference allele count and each alternative allele count. Sites with more than one variation type are discarded. Variation level = alternative allele count/ total depth. Take C-to-U RNA editing sites for instance, editing level = T/(C + T). To avoid the false-positive variants caused by technical limitations and polymorphism at DNA level, the variant sites with levels between 0.05 and 0.95 and appear in all six samples were regarded as RNA editing events (extremely low level is possibly caused by sequencing errors and extremely high level is likely caused by mutations at DNA level) (Peng et al., 2012; Ramaswami et al., 2013; Yu et al., 2016). Gene expression levels are measured by RPKM (reads per kilobase per million mapped reads). The total number of (unique) reads mapped to each gene is counted. We conducted gene count, the longest isoform of each gene is chosen. If two isoforms have the same length, they are sorted alphabetically. Htseq-count (Anders, Pyl, & Huber, 2015) is used to count the reads for each gene.

2.3 | Conservation and divergence analysis

The CDS sequences of *Arabidopsis thaliana* and *Arabidopsis lyrata* (http://plantsensembl.org/) were translated into protein sequences. We aligned these protein sequences (blastp, version 2.2) (Camacho et al., 2009) between two sibling species to define the best-matched pair of genes (orthologous gene) according to the lowest E-value (E-value < 1e⁻5 by default) and high identity. By this step, we obtained the orthologous gene list between the two species. Next, the CDS sequences were aligned (transalign, version 6.6) (Rice, Longden, & Bleasby, 2000) within each pair of orthologous genes. The orthologous sites are split manually according to the CDS alignment files. Given the aligned CDS of each pair of *Arabidopsis thaliana* and *Arabidopsis lyrata* genes, the non-synonymous and synonymous substitution rates are calculated by yn00 (Xu & Yang, 2013).
2.4 | Differential and non-differential editing sites

For each C-to-U editing site, we calculated the standard deviation (SD) of editing levels across all samples. We ranked the sites with increasing SD and divided them into two halves, termed non-differential sites and differential sites. "sd" was performed in R language with function "sd()." The same idea is applied to the dN dS comparison. We have divided the editing sites into two halves according to the dN values of host genes, or divided them into three equal bins with increasing dN/dS values.

2.5 | Statistical analysis and code availability

All statistical analyses and the graphic work were conducted in R environment (http://www.R-project.org/) (version 3.3.3). The Wilcoxon rank sum tests were performed with command "wilcox.test()" and Fisher’s exact tests were performed with command "fisher.test()."

3 | RESULTS

3.1 | The profiles of C-to-U RNA editome in Arabidopsis thaliana

The mRNA-seq data of Arabidopsis thaliana were retrieved from a previous study (Hsu et al., 2016). After variant calling process (METHODS), in total, 174 variation sites in CDS were identified and 130 of them (75%) are C-to-T changes (Figure 1a), indicating that the C-to-U RNA editing events are reliably detected. The majority (123, 94.6%, Data S1) of these 130 C-to-U editing sites are located in chloroplast or mitochondrial genes (Figure 1b). Interestingly, the context flanking the focal C-to-U editing sites (123 sites) or the background (unedited) cytidines. Only the cytidine sites in the CDS of mitochondria and chloroplast genes are considered. (d) C-to-U editing levels of the 123 mitochondrial and chloroplast editing events. The editing sites in three biological replicates (roots and shoots) are pooled. (e) C-to-U editing levels of 90 nonsynonymous (nsy) and 33 synonymous (syn) editing events. The editing sites in three biological replicates (roots and shoots) are pooled, p-value is calculated by Wilcoxon rank sum test. (f) Fractions of nonsynonymous and synonymous editing sites. Observed (obs.) and expected (exp.) values are shown. The number of expected nonsynonymous and synonymous sites is defined by assuming all cytidines in the CDS in mitochondria or chloroplast genes to be mutated to thymidine. p-value is calculated by Fisher’s exact test.
3.2 | Signals of adaptation

Among these 123 C-to-U editing sites in chloroplast or mitochondrial genes (Data S1), 90 were nonsynonymous (cause AA change) and 33 were synonymous (AA is unchanged). Interestingly, the levels of nonsynonymous C-to-U events are significantly higher than the levels of synonymous sites (Figure 1e, $p < 2.2 \times 10^{-16}$, Wilcoxon rank sum test), suggesting that high level of nonsynonymous C-to-U editing is favored by natural selection. Moreover, given the fact that 73% (90/123) of C-to-U editing sites are nonsynonymous, if we mutate all cytidines to thymidines in chloroplast and mitochondrial genes, the expected fraction of nonsynonymous mutations is significantly lower (Figure 1f, $p < 0.05$, Fisher’s exact test). This pattern again indicates that many nonsynonymous C-to-U editing events are adaptive and maintained by positive selection.

We next investigated the DNA-level conservation of the edited cytidines. We aligned the CDS between *Arabidopsis thaliana* and *Arabidopsis lyrata* to determine whether a cytidine is conserved (noncons). In this plot, the nonsynonymous editing sites in three biological replicates (roots and shoots) are pooled. $p$-value is calculated by Wilcoxon rank sum test. (b) Gene expression levels of the C-to-U edited genes and the unedited genes in mitochondria and chloroplast. RPKM (reads per kilobase per million mapped reads) is used to measure the gene expression level. Three biological replicates (roots or shoots) are pooled. MT, mitochondria; CP, chloroplast; mod, modified (edited) genes; unmod, unmodified (unedited) genes. $p$-value is calculated by Wilcoxon rank sum test. (c) Pearson’s correlation between editing level of each site and the expression level (RPKM) of its host gene. PCC, Pearson’s correlation coefficient. (d) Pearson’s correlation between sequencing depth (coverage) of each editing site and the expression level (RPKM) of its host gene. (e) Comparison between editing levels in conserved genes (low dN) and non-conserved genes (high dN). Editing sites are divided into two halves according to the dN values of host genes. $p$-value is calculated by Wilcoxon rank sum test. (f) Comparison of editing levels in different categories. Editing sites are divided into three equal bins according to the dN/dS values of host genes.
site is positively correlated with the expression level of the host gene (Figure 2c). Since highly expressed genes are usually more conserved and functionally more important, our observation indicates that the C‐to‐U editing events might be evolutionarily adaptive so that they are favored by highly expressed genes.

A potential trick here is that editing events are more likely to be detected in highly expressed genes, so that the result in Figure 2b might be due to detection bias. However, we intend to solve this problem by using the editing level as a variable instead of the occurrence of editing events. The C‐to‐U editing level is positively correlated with gene expression level (Figure 2c). Editing level = edited reads/total reads covering a site (depth). The number of total reads (depth) is intuitively positively correlated with gene expression level (Figure 2c). Editing level is positively correlated with the expression level of the host gene (Figure 2c). Since highly expressed genes are usually more conserved and functionally more important, our observation indicates that the C‐to‐U editing events might be evolutionarily adaptive so that they are favored by highly expressed genes.

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We also investigated the relationship between editing level and the gene conservation level (dN and dN/dS) (METHODS). Even the gene expression level suffers from detection bias, the dN and dS values are purely calculated from genomic sequences and are unaffected by the NGS reads. We divided the editing sites into two halves according to the dN values of host genes. We observed that editing level is generally higher in genes with higher conservation level (Figure 2e, p < .01, Wilcoxon rank sum test). Considering that dN is often negatively correlated with expression level, which makes it difficult to tease apart what factors might contribute to differences in editing between genes. We next divided the editing sites into three equal bins according to the dN/dS values of host genes. We found that the pattern observed for dN also holds for dN/dS (Figure 2f), suggesting that C‐to‐U editing occurs in more conserved genes.

We also stress the reason for discarding the sites with extremely high levels. Here, we display the variation profile between level 0.95 ~ 1 (Figure 3a). The variation types do not enrich C‐to‐T mutations and the variant sites are not enriched in mitochondrial or chloroplast genes. This mutation profile resembles the landscape of DNA mutations, suggesting that these sites are not bona fide RNA modification sites and should not be included in our analyses.
3.3 | Highly edited sites have less variable editing levels

It is conceivable that the editing events and levels could be tissue-specific as previous study revealed (Tseng et al., 2013). The tissue-specific sites could be caused by the tissue-specific expressed genes. Our strict pipeline only included the editing sites appearing in all samples (three root samples and three shoot samples), which consequently discarded the root- or shoot-specific genes or sites. We ask that if highly edited sites are more likely to be beneficial and functional, then they should exhibit consistent editing level to assure their proper function.

Given the fact that the edited genes show highly correlated expression levels between roots and shoots (Figure 3b), our editing level comparison would not suffer from detection bias across different tissues. We rank the editing sites by the editing level difference between the root and shoot samples (standard deviation across the samples, Figure 3c, also see METHODS). Differential sites (diff) have larger difference of editing levels across the samples, and vice versa (Figure 3c,d). We find that the less variable half of sites have significantly higher editing levels than the more variable half of editing sites (Figure 3e, p < .001, Wilcoxon rank sum test). It is possible that the highly edited sites (mostly nonsynonymous sites) are more essential and functional so that their editing levels tend to be “fixed.” We further tested that the host genes of differential or non-differential sites have similar expression levels (Figure 3f), indicating that the variation observed in editing level might not be caused by gene expression difference.

4 | DISCUSSION

According to Edera et al. (2018, their findings could draw to a conclusion that the nonsynonymous editing events are selectively favored, which is similar to ours. However, their multiple angiosperm study revealed another interesting trend that highly edited cytidines were likely to be replaced with thymidines during genome evolution. Although other study also discovered that the loss of editing events in kin species might be due to the changed editing factor (Kawabe et al., 2019), the C-to-T DNA switch found by Edera et al. caught our interest. If the advantage of RNA editing is increasing the protein diversity, then the C-to-T-mutated DNA would actually abolish this diversity. An alternative explanation for the C-to-T DNA switch is, the edited isoform (U/T version) is more advantageous than the unedited isoform (C version), so that those extremely highly edited sites are almost equivalent to a permanent DNA mutation. This idea could be supported by our observation that highly edited sites have less variable editing levels (Figure 3), which possibly reflects the necessity for steady and consistent editing level for these sites. At this stage, why these edited genes do not directly change their DNA sequences before the occurrence of RNA editing remains unknown.

Here, we show what patterns of editing would look like if C-to-U editing is generally selectively neutral, deleterious, or beneficial. Positive selection: in a previous paper studying A-to-I RNA editing events in Arabidopsis thaliana (Liu & Zhang, 2018; Xu & Zhang, 2014) proved the non-adaptive feature of the RNA editing events by showing that (a) the fraction of nonsynonymous editing events is lower than the fraction of synonymous editing; (b) the nonsynonymous editing levels are no higher than the synonymous editing levels. The author concludes that these editing events are slightly deleterious or neutral under the observed patterns. We surmise that if C-to-U editing in Arabidopsis is slightly deleterious or neutral, it should exhibit those patterns as described above.

Our study systematically demonstrates that the C-to-U editing events in Arabidopsis thaliana are overall adaptive and favored by natural selection. Together with previous literatures, the advantage of this editing mechanism might be the amino acid changes conferred by the high-level nonsynonymous editing events. Our work analyzed the C-to-U RNA editome from novel aspects, which broadened our horizon in the Arabidopsis community and should also be appealing to the evolutionary phytologists.

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CONFLICT OF INTEREST

The authors declare they have no conflict of interest.

AUTHOR CONTRIBUTIONS

Lai Wei designed and supervised this research. Duan Chu and Lai Wei analyzed the data. Duan Chu and Lai Wei wrote this article.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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