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Genetic studies on resistance to Rice tungro bacilliform virus and evolution of its related endogenous viruses in the genus Oryza [an abstract of dissertation and a summary of dissertation review]

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Rice, *Oryza sativa* L., is one of the important cereal crops that provides a staple food for half of the world’s population. Rice viral diseases constitute a major obstacle to rice production. Rice is known to be infected by more than 30 viruses, but 25 viruses do not have direct economic effects in terms of rice production. Rice tungro disease (RTD) caused by Rice tungro bacilliform virus (RTBV) negatively affects rice production in South Asia and Southeast Asia. Plants have developed multilayered defense mechanisms against microbial pathogens such as gene silencing of viral small RNA (vsRNA) or host micro RNA (miRNA). Regarding the anti-viral defense against (RTBV), I considered three factors namely, endogenous RTBV like (eRTBVL) sequence, vsRNA and rice miRNA, in this study.

In Chapter 2, I focused on eRTBVL sequences, which have been identified in the rice genome. The distribution of eRTBVL sequences differs between the two genomes; the *O. sativa* genome contains more than 100 copies of eRTBVLs, while the *O. glaberrima* genome has only a few copies. Differences in the eRTBVL copy number in these genomes might be associated with the different reactions of these two species to RTBV. Firstly, I confirmed innate vulnerability to RTBV of *O. glaberrima* carrying a few eRTBVL copies, then the possible function of eRTBVL-derived small RNAs were examined by the comparison between *O. sativa* and *O. glaberrima*. Small RNA sequencing analysis indicated that the expression of small RNAs from eRTBVL might be associated with the phenotypic differences between the two genotypes after RTBV infection. I could suggest two possible ideas to explain the interaction between the susceptibility to RTBV and the presence of eRTBVL. One is that the constitutive expression of small RNAs from eRTBVL promotes the silencing machinery mediated
by vsRNA in RTBV-infected plants. The other possibility is that the small RNAs from eRTBVL respond immediately after RTBV infection to inhibit the replication of the virus at an initial stage of the infection.

In Chapter 3, the resistant (TW16) and susceptible (TN1) genotypes against RTBV were used for RTBV infection test and small RNA sequencing. To examine the RNAi-based defense machinery in rice during RTBV infection, I characterized vsRNA and rice-derived miRNA by small RNA sequencing and compared the expression of small RNAs between TN1 and TW16. A total of 11 crucial miRNAs related to RTBV resistance were identified in this study. The expression analysis of AGO genes, vsRNA and rice miRNA indicated that the resistant plants against RTBV might interfere with the cleavage of target RNAs by down-regulation of miRNAs and protect the rice growth from viral disease symptom. The identified miRNAs might be valuable candidates to study novel insight into the defense mechanisms of resistant genotypes against RTBV.

Chapter 4 describes evolutionary patterns of the local chromosomal regions during the Oryza AA-genome speciation through the comparisons of eRTBVL-D sequences among the Oryza AA-genome species. eRTBVL-D is known as the oldest eRTBVL family that was integrated into the Oryza AA-genome before speciation. The eRTBVL-D sequences were available for the evolutionary study of the local-chromosomal regions. I found the swerve phylogenetic relationships from the known Oryza speciation and chromosomal rearrangements at the eRTBVL-D loci. I described the functional implications of eRTBVL-D loci by the evolutionary analyses for natural selection. The results of Tajima’s D test indicated that natural selection might not have acted on eRTBVL-D as the central factor. Hence, eRTBVL could exist as “genomic fossils”. In this chapter, I proposed that eRTBVL-D loci could be a useful resource as genetic neutral markers, because they have rarely selective effect. The analysis of eRTBVL-D provided novel evolutionary insight for local chromosomal regions during the rice AA-genome speciation.

Taken together, this study proposed the possible RTBV-viral mechanism in the resistant plants and the evolutionary history of the related endogenous viruses. I expect that my study gives an insight into how to develop resistant rice cultivars against Tungro disease.