A Region on Chromosome 7 Related to Differentiation of Rice (Oryza sativa L.) Between Lowland and Upland Ecotypes

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Due to global population expansion and climate change impacts, the development of a stable yielding variety that adapts well to unfavorable conditions for rice cultivation, can contribute to sustainable and stable production in rice (Oryza sativa L.). Understanding genetic differentiations to ecotypes for rice cultivations, such as upland, rainfed lowland, and irrigated lowland, is very important to develop the breeding materials for adapting to each environmental condition. The upland landrace variety basically has low tiller/panicle numbers and a large panicle, and the plant architecture is different from that of the lowland variety. The tiller and panicle numbers have been considered as one of the most difficult traits for genetic changes artificially in rice breeding. A low tiller recessive gene ltn2 originated from a New Plant Type variety, IR 65600-87-2-23, harboring segments from an upland variety, Ketan Lumbu (Tropical Japonica Group), was found on chromosome 7, and the other QTLs for culm length, culm weight, panicle length, panicle weight, seed fertility, harvest index, and soil surface rooting were also detected in the same chromosome region. These low tiller genes and the other QTLs were estimated to play an important role in developing the architecture for upland rice. A low tiller recessive gene ltn2 originated from a New Plant Type variety, IR 65600-87-2-23, harboring segments from an upland variety, Ketan Lumbu (Tropical Japonica Group), was found on chromosome 7, and the other QTLs for culm length, culm weight, panicle length, panicle weight, seed fertility, harvest index, and soil surface rooting were also detected in the same chromosome region. These low tiller genes and the other QTLs were estimated to play an important role in developing the architecture for upland rice. Some QTLs for root growth angle, DRO3 and qSFR7, were also found in the same chromosome region from upland varieties categorized into the Tropical Japonica Group, and the QTLs may also be relevant to upland adaptation together with other traits. Previous studies using high throughput re-sequencing (whole genome variation data) of a large batch of rice accessions could identify the ecotype differentiated genomic regions (EDRs) and Ecotype differentiated genes (EDGs) such as Os07g0449700, a type response regulator, which is critical in upland adaptation in the same region of chromosome 7. Two selective loci, E3735 and E4208, for upland and lowland differentiation, and their corresponding genes Os07g0260000 and Os07g0546500 were also detected on chromosome 7 by drought-responding EST-SSRs. These findings indicate that the region on chromosome 7 is highly possible to related to the plant shoot and root architecture in the upland rice variety that has an important role and differentiates between upland and lowland ecotypes.

Keywords: differentiation, upland, lowland, ecosystems, plant architecture, rice (Oryza sativa L.)
INTRODUCTION

Climate change is the main causal element or factor of biotic and abiotic stresses, which have negative effects on global food production including rice (Oryza sativa L.) (Raza et al., 2019). Recently, one study estimated a 15% decrease of rice yield in irrigated conditions of developing countries and a 12% increase in rice price as a result of climate change by 2050 (IFPRI, 2009). By contrast, current estimates showed that rice production had to increase by over 20% before 2030 to satisfy the demands of the world’s growing population (Purevdorj and Kubo, 2005; FAO RMM, 2018) and to avoid food crises.

The development of a stable yielding variety that adapts well to unfavorable conditions for rice cultivation, such as rainfed lowland and upland ecosystems, can contribute to sustainable rice production and ensure global supply. The respective characteristic traits of rice varieties have been modified under both natural and artificial selections, which lead to phenotypical adaptation to the respective environment and genetic differentiation at the same times between ecotypes (Lyu et al., 2014). Under the upland or rainfed lowland ecosystems, rice cultivation needs to reduce water use in rice production and increase the water use efficiency, and from an environmental perspective, emission of methane is substantially lower in this condition (Xia et al., 2014). Understanding the genetic differentiation among upland, rainfed lowland, and irrigated lowland rice is very important for rice breeding in unfavorable conditions. Especially, rice varieties between upland and irrigated lowland differ significantly in phenotypical and physiological traits. Particularly, the upland variety basically has low tiller/panicle numbers, tick and long culm, and a large panicle, and the plant architecture is different from that of lowland varieties.

Several studies have been conducted to fine the genetic factor(s) for differentiation of ecotypes between lowland and upland. Ishikawa et al. (1992) found that the allele of isozyme gene Pgd-1 locating chromosome 11 (Wu et al., 1988) was related with the differentiation between Japanese upland varieties and Japanese lowland ones. Zheng et al. (2000) detected three QTLs, qPRN7 for root penetration ability, qRN7 for root numbers, and qRPI for root penetration index on chromosome 7 in a wax-petroleum layer system simulated to Asian upland soil. Thick root and high penetration ability are important for drought resistance in upland soil. Two deeper root QTL, DRO3 by Uga et al. (2015) and qDRD7 by Luo et al. (2015) which regulate root-growth-angle of upland rice were also detected on chromosome 7. Uddin et al. (2016a; 2016b) described qPN7 for panicule number and a gene for low tiller number, ltn2, which constrict plant architecture in upland on chromosome 7. Monkada et al. (2001) and Lafitte et al. (2004) reported qDTH7, qSF7 and qGW7, which extended the heading date and increased the spikelet fertility and per grain weight in upland. Xu et al. (2020) recently reported upland specific QTLs, qHD7, qGY7, and qHI, which plays important role in reproduction and yields in upland. Three candidate genes; Os07g0449700 by Lyu et al. (2014), Os07g0260000 and Os07g0546500 by Xia et al. (2014), were also reported as ecotype differentiation genes on chromosome 7 which regulate the root system and plant height in upland rice varieties and help with drought tolerance or resistance. Thus, some chromosome regions were estimated to related with the genetic differentiation between lowland and upland ecosystems, and many QTLs genes for agronomic traits differentiating between ecotypes in lowland and upland have been detected (Table 1).

| QTL or gene detected | Physical position (Mb) of the closest marker* | Regulatins/Functions | Origin | Ecotype | Reference |
|----------------------|---------------------------------------------|----------------------|--------|---------|-----------|
| DRO3                 | 24.2                                        | Root angle           | Kinandang Patong | Upland | Uga et al. (2015) |
| qSOR1                | 25.6                                        | Surface rooting      | Gemedjah Beton  | Lowland | Li et al. (2012)  |
| qPV7                 | 24.5                                        | Panicle No.          | IR65660-87-2-2-3 | Upland | Uddin et al. (2016a) |
| qRN7 and ltn2        | 25.1                                        | Low tiller No.       | Azucena         | Upland | Lafitte et al. (2004) |
| qSF7                 | 26.0                                        | Spikelet fertility & Root depth | B61144F-MR-6 | Upland | Tomita and Fukuta (2019) |
| qGW7                 | 26.0                                        | Weight per Grain & Root depth | IR65660-87-2-2-3 | Upland | Xi et al. (2020) |
| qSRF7                | 23.6                                        | Soil Surface rooting | IR65660-87-2-2-3 | Upland | Uddin et al. (2016a) |
| qHD7                 | 25.5                                        | Heading date         | B61144F-MR-6    | Upland | Luo et al. (2015)  |
| qGY7                 | 25.5                                        | Grain yield          | IRAT 109        | Upland | Li et al. (2011)  |
| qRDR7                | 26.0                                        | Deep rooting         | IRAT 109        | Upland | Li et al. (2011)  |
| qBR7                 | 19.0                                        | Root thickness       | IRAT 109        | Upland | Li et al. (2005)  |
| qBN7                 | 25.9                                        | Root thickness       | IRAT 109        | Upland | Li et al. (2005)  |
| qDTH7                | 24.8                                        | Heading date         | O. rufipogon (RGC #105491) | Upland | Monkada et al. (2001) |
| qPRN7                | 17.8                                        | Penetrated root numbers | Azucena         | Upland | Zheng et al. (2000) |
| qTR7                 | 14.4                                        | Total root numbers  | Panel of upland varieties | Upland | Lyu et al. (2014) |
| qRPI                 | 28.4                                        | Root penetration index | Panel of upland varieties | Upland | Xia et al. (2014) |
| Os07g0449700         | 16.1                                        | Root and shoot development | Panel of upland varieties | Upland | Lyu et al. (2014) |
| Os07g0260000 Os07g0546500 | 9.1   | Drought tolerance    | Panel of upland varieties | Upland | Xia et al. (2014) |

*Physical position of the closest markers were retrieved from the genome database in Gramene (http://www.gramene.org) and Oryzabase (https://shigen.nig.ac.jp/oryzabase/), DRO, deeper rooting; SOR, soil surface rooting; BRT, basal root thickness; RDR, ratio of deep rooting; PN, panicule number; TN, tiller number; HI, harvest index; GW, grain weight; SF, spikelet fertility; GW, grain weight; DTH, days to heading; DH, heading date; RP, root penetration; RN, root number; PRN, penetrated root number.

**TABLE 1** | List of genes and QTLs on chromosome 7 for ecotype differentiation.
Interestingly, most of the research findings indicated that a region on chromosome 7 was related to the variations of unique traits between the two ecotypes, such as drought tolerance/avoidance and upland adaptation. This article review examines the involvement of chromosome 7 in the genetic differentiation of upland and lowland varieties, and then discusses how this information can be used in breeding a rice variety that adapts well to unfavorable conditions for sustainable production in rice.

A LOW TILLER GENE CONTRIBUTING TO PLANT ARCHITECTURE

Plant architecture with a low tiller and high-density grain is desirable in limited water conditions to maintain a proper plant density, to avoid episodic drought, and to keep the stable yield production based on the minimizing of reproductive tillers under the serious conditions, such as rainfed lowland and upland (Vergara et al., 1990; Farooq et al., 2011). The control for numbers of tiller and panicle may be the important breeding target for increasing the adaptability and seed production in the growing stages from productive to reproductive stages of rice against upland and serious environmental conditions.

Several artificial mutant genes for tiller numbers, tdr2 (Hasegawa et al., 2005) and rcn9 (Jiang et al., 2006) on chromosome 1, OsTB1 on chromosome 3 (Takeda et al., 2003), HTD1 on chromosome 4 (Zou et al., 2005), and D3 (Ishikawa et al., 2005), MOCl (Li et al., 2003), and rcn8 (Jiang et al., 2006) on chromosome 6, have been found in rice. These genes are associated with branching mechanisms of the tiller. As the natural variations, many QTLs for numbers of panicles and tillers have been reported. Lin et al. (1996) found them on chromosomes 2, 4, 5, and 6; Wu et al. (1998) were on chromosomes 1, 2, 3, 4, 5, 6, 7, 8 and 12; Nagata et al. (2002) were on chromosome 1; Hittalmani et al. (2003) were on chromosomes 1, 3, 4, and 12; Miyamoto et al. (2004) were on chromosomes 2, 5, 6, and 8; Liu et al. (2008; 2010; 2012) were on chromosomes 1, 2, 3, 4, 6, 7, 8, 9, and 12, and Bian et al. (2013) were on chromosomes 1, 2, 4, 5, 6, 7, 8, and 10. A low tiller-number gene, Ltn was found, on chromosome 8 in a Japanese Japonica Group variety, Aikawa 1 (Fujita et al., 2010). Thus, several genes and many QTLs have been identified almost rice chromosomes except for chromosome 11, as the genetic factors for controlling the tiller and panicle numbers in rice. These QTLs for tiller or panicle numbers by Yan et al. (1998), Liu et al. (2008; 2010; 2012), and Bian et al. (2013) were detected in the similar region of long arm on chromosome 7.

International Rice Research Institute (IRRI) developed the New Plant Type (NPT) varieties characterizing as the low tiller numbers and a high productive tiller ratio by introducing the chromosome segments from the Tropical Japonica Group varieties, as one of the traits for genetic improvements to increase rice yield production (Khush, 2000). Fujita et al. (2009) reported 334 introgressions lines (INLs) with the genetic background of an Indica Group variety IR 64 and harboring several chromosome segments from NPT varieties which were bred by the crosses between Indica Group and Tropical Japonica Group upland varieties. In the other words, these chromosome segments in INLs were mainly originated from the Tropical Japonica Group upland varieties. Uddin et al. (2016a) found 18 QTLs for yield component traits including a QTL for low panicle numbers (as an indicator of tiller number), qPN7, on chromosome 7, by association analyses with 35 INLs harboring chromosome segments from the NPT variety, IR 65600-87-2-2-3 among them (Figure 1). IR 65600-87-2-2-3 harbors the chromosome segments of the Tropical Japonica Group upland rice variety, Ketan Lumbu. The other QTLs for culm length, panicle length, panicle weight, harvest index (panicle weight/total weight), and fertility rate were also found in the same region detected qPN7 on chromosome 7. To confirm the detailed chromosome position of the QTLs detected,
advanced genetic studies were conducted using two hybrid populations. A total of 88 F₃ family lines derived from a cross between IR 64 and YTH34 which was one of 35 INLs harboring the segment on chromosome 7 from the NPT variety, IR 65600-87-2-2-3. The values of panicle numbers, spikelet numbers, and dry weight of whole plant in YTH34 were lower than those of IR 64 under the cultivation in irrigated lowland condition, and remarkable reductions of these values were observed under upland cultivation (Table 2) and an F₃ population consisting of 72 plants, which was self-pollinated from an F₂ plant, F₂-JII-IV-10, harboring the heterozygote chromosome region for qPN7 (Uddin et al., 2016b). These segregation analyses under upland conditions confirmed the single gene segregations. It was found a QTL for low tiller and designated as a recessive gene, ltn2, originated from the Tropical Japonica Group rice (Figure 2). Linkages were also found with some Simple Sequence Repeats (SSR) markers RM505, MRG5344, and RM21950. The genetic distances between RM505 and ltn2 were 3.4 cM, and between RM21950 and ltn2 they were 1.1 cM, and ltn2 was mapped most closely with RM21950. Advanced QTL analysis also confirmed other QTLs for culm length, culm weight, panicle length, panicle weight, seed fertility, and harvest index in the same chromosome region. These QTLs decreased the traits’ values with the YTH34 allele. The INLs, YTH34, with IR 64 genetic background, is

### Table 2 | Agronomic traits of IR 64 and YTH34 under irrigated lowland and upland.

| Conditions  | Variety | Panicle No./plant | Panicle length (cm) | Spikelet No./panicle | Seed fertility (%) | Panicle weight (g) | Dry weight (g) | Harvest index (%) | Days to heading |
|-------------|---------|-------------------|---------------------|---------------------|-------------------|-------------------|----------------|------------------|----------------|
| Irrigated   | IR 64   | 22.0              | 70.3                | 25.3                | 130.6             | 48.2              | 25.1           | 78.5             | 31.9           | 107.6          |
|             | YTH34   | 15.0 (68)         | 70.8 (100)          | 25.9                | 109.1             | 85.4              | 32.2           | 66.6 (84)        | 48.4           | 108.3          |
| Lowland     | IR 64   | 47.0              | 51.3                | 25.9                | 138.0             | 63.4              | 51.9           | 140.0            | 38.0           | 140.0          |
|             | YTH34   | 17.0              | 39.9                | 22.8                | 102.0             | 66.6              | 14.6           | 48.0 (34)        | 30.0           | 151.0          |

Rice varieties were cultivated at Tropical Agricultural Research Front, JIRCAS, Ishigaki, Okinawa Japan, in 2013 (Date were refereed from Uddin et al., 2016b). Mean values among 12 plants were calculated in each trait. ( ) = Relative value (%) in compared with that of IR 64.

![Figure 2](https://example.com/fig2.png)

FIGURE 2 | Chromosome location of low tiller number gene, ltn2 and QTLs detected on chromosome 7. (A) Uddin et al. (2016a); (B, C) Uddin et al. (2016b); DRO3: Uga et al. (2015); qSFR7: Tomita and Fukuta (2019); qBRT1: Li et al. (2011); qDR1: Luo et al. (2015); qS7: GWA7: Laffitte et al. (2004); qSOR1: Uga et al. (2012); qDTH: Monkada et al. (2001). Y: YTH34 allele decreased value in upland; Y: YTH34 allele decreased value in irrigated lowland. CW, culm weight; PW, panicle weight; PW/TW, ratio of panicle weight to total plant weight; CL, culm length; PL, panicle length; PN, panicle number; FS/TS, ratio of fertile spikelet per panicle; BRT, Basal root thickness; RDR, Ratio of deep rooting; SOR, Soil surface rooting; SF, Spikelet fertility; GW, Grain weight; DTH, Days to heading. *Irrigated lowland in 2011, **Upland in 2011, ***Irrigated lowland in 2012, and ****Upland in 2012. **Physical position of the Chromosome 7 was redrawn from the genome database in Oryzabase (https://shigen.nig.ac.jp/oryzabase/) using the Nipponbare reference genome. Black line: YTH34, Dashed line: YTH34, Dotted line: YTH34, Flanked region of QTL.
introduced only the genetic factor(s) for tiller/panicle numbers and those for the other traits locating the other chromosomes were not harbored. These results indicated that In2 originated from a tropical Japonica Group upland rice variety, Ketan Lumbu, had a strong genetic effect for occurring of tiller and panicle with negative effects for the other traits. The ecotype of upland landrace rice is characterized such as traits; low panicle/tiller numbers, long and thick culm, and large panicle. These traits are controlled by several genetic factors which were distributed on the rice genome chromosomes in each, and these were accumulated in the genetic background of upland ecotype rice. The tiller/panicle numbers have been known as the most difficult trait to detect genetically, because of large environmental error (Sasahara, 1997). In the other words, these were easy to be influenced and changed by environmental conditions. To control the tiller/panicle numbers, the genetic factor(s) with strong effects may be needed under the serious and unfavorable environmental conditions of upland. The low tiller gene and the other QTLs detected on chromosome 7, which were originated from the Tropical Japonica Group variety, were estimated to play the important role for developing shoot architecture of upland rice with strong effects.

**GENETIC FACTORS FOR ROOT ARCHITECTURE**

One major QTL for soil-surface roots, qSOR1, was found using 124 recombinant inbred lines derived from a cross between an Indonesian lowland rice cultivar, Gendjah Beton, with soil-surface roots and a Japanese lowland rice cultivar, Sasanishiki, was detected and localized on the long arm of chromosome 7 by Uga et al. (2012).

A QTL for deep root-growth-angle, DRO1, was found on chromosome 9, using 26 chromosome segment substitution lines (CSSLs) which were developed from the cross combination between an upland variety, Kinandang Patong (Tropical Japonica Group), and IR 64 as the recurrent parent (Uga et al., 2015). And another QTL, DRO3, was also found on the long arm of chromosome 7 using the progenies derived from a cross between IR 64 and a CSSL harboring DRO1. DRO3 contributed to the deep root growth angle trait with the allele of Kinandang Patong. The nearest marker RM5508 for DRO3 locates in the same chromosome regions of In2 with the distance of 0.7 cM. Another QTLs for deep rooting, qRDR7, was also reported in the same region of chromosome 7 and the allele of an upland variety IRAT109 contributed to deep rooting in the genetic background of Zhenshan 97B which was a shallow rooting parent by Luo et al. (2015).

Tomita et al. (2017) tried to clarify the genetic variation for the root angle distributions using 97 accessions, and these were classified into two cluster groups, A and B. The accessions of cluster group A showed shallow rooting including the soil-surface root type, but and the numbers of accession were few. An INL, YTH16, by Fujita et al. (2009) harboring chromosome segments from a common NPT variety, IR65600-87-2-2-3, was also included. The accessions in cluster group B showed a wide variation from shallow to deep rooting types in both Indica and Japonica Groups, lowland and upland ecotypes, and landraces and improved types. There were no relationships between the root vertical angle and total root numbers among them. These findings indicated that the root angle distributions were not related with the differentiations between Japonica Groups and Indica Groups, among ecotypes for lowland and upland, and among degrees of genetic improvement, and the accessions with soil-surface root were rare and unique among natural variations. Tomita and Fukuta (2019) clarified that the soil-surface-rooting in YTH16 was controlled by three QTLs on chromosomes 2, 5, and 7, and one QTL, qSFR7, on chromosome 7 had the biggest effect and played a main role. qSFR7 was detected in the same region as In2 by Uddin et al. (2016b). The QTL for soil surface-rooting on chromosome 7 is also one of the unique characters originated from the Tropical Japonica Group variety in rice. The genetic factor(s) for root architecture may also be relevant to upland adaptation together with other traits.

**ECOTYPE DIFFERENTIATED GENES**

With the basis of the rice genome sequence and annotation databases, NCBI: http://www.ncbi.nlm.nih.gov/gene/, TIGR: http://rice.plantbiology.msu.edu/, we found the physical locations of MRG5433 on chromosome 7 and three putative and expressed genes close to the target region of the Japonica Group rice genome (cultivar: Nipponbare), such as Os07g0607500, Os07g0607700, and Os07g0607800, were found that were related to abiotic stress tolerance (Sottosanto et al., 2007; Zhang et al., 2008; Huang et al., 2009; Kera et al., 2012).

Using the entire genome resequencing data from a large panel of 84 upland and 82 irrigated lowland rice accessions, Lyu et al. (2014) reported some selective deviations in rice chromosomes, which were called as the ecotype differentiation genomic regions (EDRs). In the EDRs, several individual ecotype’s differentiated genes (EDGs) that are critical in upland adaptation or the phenotypical differentiation between the two ecotypes, lowland and upland, were also detected. A type-A response regulator (ARR) gene, Os07g0449700, on chromosome 7, is of special interest as it belongs to the ARR gene family induced by cytokinin, plays important roles in cytokinin signaling, and has impacts on root and shoot development. This ecotype differentiation of this ARR gene, Os07g0449700, may explain some differences of root and plant height between upland and irrigated rice varieties. Further reciprocal transgenic experiments between upland and irrigated rice will clarify the effects of Os07g0449700 in the adaptation of upland rice.

Xia et al. (2014) reported seven selective loci that are ecotype preferable alleles expressed under drought stress by the analyses for 47 drought-responding expressed sequenced tags-simple sequence repeats (EST-SSRs) markers, using 377 rice landraces collected from China. These EST-SSRs markers are selected from the DNA transcribe regions and are closely related to the functional gene expressed under drought stress. Among the seven,
two loci, E3735 and E4208, were related to the differentiation between upland and lowland varieties, and these corresponded to Os07g0260000 and Os07g0546500, which were detected on the rice chromosome 7, which may have an important effect on drought resistance or drought tolerance in rice.

CONCLUSION

Crops adapted to different agro-ecosystems always promote the variation of agriculturally important genes (Xia et al., 2014). The knowledge and information regarding the genetic differentiation for eco-typical variations of traits and adaptations will greatly contribute to harnessing the genetic resources for the breeding and cultivation of rice under upland as well as unfavorable environments. We reviewed the involvement of a region on chromosome 7 in the genetic differentiations for rice ecotypes to lowland and upland ecosystems.

The ltn2 region on chromosome 7 showed the unique reactions: low tiller, low dry matter production, short panicle length, and short culm lengths (Uddin et al., 2016b), and soil surface rooting (Tomita et al., 2017; Tomita and Fukuta, 2019). Usually, the upland landrace varieties are shown the unique characters: low tiller, heavy and big panicle, and short and early maturation, and these agricultural traits might contribute to avoid the risk of short water supply and to keep the stable yield production based on the minimalizing reproductive tillers under rainfed conditions. This unique shoot architecture of low tiller and panicle may be an important trait for adaptation to serious upland environmental conditions (Uddin et al., 2016b). YTH34 had a lower plant height, spikelet numbers, dry matter productions, and longer time to heading, and did not show good performances compared with that of IR 64 under the upland condition (Table 2). These results suggest that the genetic background of ltn2 is efficient in the upland variety. This ltn2 related to one of the genetic factors for the adaptation of rice cultivar to upland conditions might be very useful for the genetic improvement of rice cultivars under upland or unfavorable conditions with the other genetic factors.

YTH34 harboring ltn2 and the genetic information of the low tiller gene including linked SSR markers as well as the proposed candidate genes will be useful for the genetic modification of plant architecture in the rice variety, and for understanding the genetic mechanism of differentiation between lowland and upland ecotypes. The model plant type for adaptation to upland conditions will be able to be reconstructed by ltn2. The information of gene pyramiding based on ltn2 will help us to understand the architecture of upland rice cultivars.

The ecotype differentiated genes such as Os07g0449700 for root and shoot development (Lyu et al., 2014), EST-SSRs based selective loci E3735 and E4208 and their corresponding genes Os07g0260000 and Os07g0546500 for drought resistance (Xia et al., 2014), and other detected QTLs (Uga et al., 2015; Uddin et al., 2016a; Uddin et al., 2016b) on the rice chromosome 7 also play key roles in upland adaptation and result in the phenotypical differentiation. Clarification of the relationships among these candidate genes, ltn2, and QTL for the soil surface root on chromosome 7, will be investigated in future studies.

This review will not only help the geneticists to understand the underlying molecular basis of adaptive divergence, but also provide to breeder and agronomist valuable information for rice domestication and adaptation, especially on the upland/water stress and drought tolerant in rice.

AUTHOR CONTRIBUTIONS

MNU prepared the manuscript basically, and YF made the plan for the conformance of the manuscript. The paper was prepared based on the PhD course study for MNU under the supervision of YF in University of Tsukuba.

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