Genome-Wide Analysis of Spatiotemporal Expression Patterns During Rice Leaf Development

Masayuki Miya  
The University of Tokyo: Tokyo Daigaku

Takanori Yoshikawa  
Kyoto University: Kyoto Daigaku

Yutaka Sato  
National Agriculture and Food Research Organization

Jun-Ichi Itoh (ajunito@g.ecc.u-tokyo.ac.jp)  
The University of Tokyo: Tokyo Daigaku  
https://orcid.org/0000-0001-8952-2697

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Abstract

Background: Rice leaves consist of three distinct regions along a proximal-distal axis, namely the leaf blade, sheath, and blade-sheath boundary region. Each region has a unique morphology and function, but the genetic programs underlying the development of each region are poorly understood. To fully elucidate rice leaf development and discover genes with unique functions in rice and grasses, it is crucial to explore genome-wide transcriptional profiles during the development of the three regions.

Results: In this study, we performed microarray analysis to profile the spatial and temporal patterns of gene expression in the rice leaf using dissected parts of leaves sampled in broad developmental stages. The dynamics in each region revealed that the transcriptomes changed dramatically throughout the progress of tissue differentiation, and those of the leaf blade and sheath differed greatly at the mature stage. Cluster analysis of expression patterns among leaf parts revealed groups of genes that may be involved in specific biological processes related to rice leaf development. Moreover, we found novel genes potentially involved in rice leaf development using a combination of transcriptome data and in situ hybridization, and analyzed their spatial expression patterns at high resolution. We successfully identified multiple genes that exhibit localized expression in tissues characteristic of rice or grass leaves.

Conclusions: Although the genetic mechanisms of leaf development have been elucidated in several eudicots, direct application of that information to rice and grasses is not appropriate due to the morphological and developmental differences between them. Our analysis provides not only insights into the development of rice leaves but also expression profiles that serve as a valuable resource for gene discovery. The genes and gene clusters identified in this study may facilitate future research on the unique developmental mechanisms of rice leaves.

Background

Leaves, which are the main site of photosynthesis in higher plants, are usually polarized along three axes: proximal-distal, adaxial-abaxial, and medial-lateral. Tissues arranged along these axes have characteristic morphologies and functions. As leaves are derived from immature cell populations protruding from the shoot apical meristems, their morphology and functions must be acquired during the course of development. Leaf development is a tightly orchestrated process incorporating multiple events crucial to organogenesis: axis determination, pattern formation, and identity establishment. Additionally, the growth of leaf primordia, which relies on cell proliferation and differentiation, is precisely regulated both temporally and spatially to produce typically shaped leaves.

The morphology of leaves varies greatly among species and developmental phases and environments, and this variation is driven by differences in leaf genetic programs among species. Hence, the mechanisms regulating leaf morphogenesis should be studied in a wide variety of species. Most information currently available has been obtained from analyses of the model eudicot plant Arabidopsis. A number of genes regulating leaf development have been identified in Arabidopsis [1], and the molecular
mechanisms of leaf morphogenesis in various species have been elucidated based on information obtained from *Arabidopsis*.

Grasses belong to the monocot clade, and their leaf morphology is distinct from that of *Arabidopsis*. Although grass leaves are polarized along the same three axes as those of other plants, they are unique in that distinct regions with differing morphology and function are located along the proximal-distal axis (Fig. 1a). The leaf blade is the distal part of the leaf; it has a flat structure and is rich in mesophyll cells, in which photosynthesis occurs. The leaf sheath is located at the basal part of the leaf and has a thick structure that protects inner leaves and provides structural support to the blade. The boundary region between the blade and sheath comprises the lamina joint, ligule, and auricle. The lamina joint acts as a hinge that allows the blade to bend abaxially, thereby optimizing light capture by the blade. Each of the three regions undergoes different developmental processes. In addition, spatiotemporal coordination of tissue differentiation during development contributes to the final leaf morphology. As with *Arabidopsis*, tissue differentiation in grass leaves proceeds in the basipetal direction, suggesting that these processes are under precise spatial and temporal control by genetic mechanisms.

To date, several genes that are important for grass leaf morphology have been identified. Related to the organs and tissues that are differentiated along the proximal-distal axis, *LIGULELESS1*, a member of the *SQUAMOSA PROMOTER BINDING-LIKE* (*SPL*) gene family, is essential for the differentiation of organs in the blade-sheath boundary region [2, 3, 4]. Homologs of *Arabidopsis BLADE-ON-PETIOLE* genes in rice are important for sheath development [5]. Class I KNOX genes important for the maintenance of the shoot apical meristem (SAM) are believed to provide proximal cues for leaf primordia [6, 7]. Meanwhile, cell proliferation patterns have been reported to affect grass leaf morphology. Cell proliferation in immature tissues of leaf primordia is controlled by protein complexes encoded by two gene families, *GROWTH-REGULATING FACTORS* (*GRFs*) and *GRF-INTERACTING FACTORS* (*GIFs*) [8, 9, 10, 11]. Changes in their protein complex composition reportedly serve as a switch for the transition from cell division to cell expansion [12]. However, few of the genes that play important roles in grass leaf development have been identified.

For elaboration of leaf development and morphology, gene expression should be precisely regulated both temporally and spatially. Thus, revealing the expression patterns of genes during development would contribute significantly to understanding the genetic mechanisms behind the process of leaf establishment. Transcriptome analysis is a powerful method for exploring gene expression dynamics at both the genome-wide and single-gene levels. To date, transcriptome analysis of leaf development has been performed in species including *Arabidopsis* [13, 14], maize [15, 16, 17, 18, 19, 20, 21, 22], and rice [23, 24, 25]. In particular, transcriptome changes accompanied by tissue differentiation have been intensively studied in the developing leaf blade in maize. However, no study to date has reported the temporal transcriptomic changes occurring from leaf initiation to leaf maturation in rice. Furthermore, most transcriptome studies of grass leaves have been performed on tissues from only one of the regions along the longitudinal axis, making it difficult to draw direct comparisons among regions. Therefore, to
fully elucidate gene expression profiles and characterize gene function, it is necessary to investigate spatiotemporal changes in leaf transcriptomes from each region along the longitudinal axis of the leaf.

In this study, we performed transcriptome analysis of rice leaf development using the Agilent rice 44K microarray, which is compatible with the rice expression database RiceXpro [26, 27]. Our experimental design included a broad range of developmental stages and several distinct regions along the leaf longitudinal axis, which allowed us to capture overall transcriptome dynamics throughout leaf development. Our data analysis uncovered trends in the expression patterns of certain gene clusters during leaf development and revealed relationships between developmental events and those gene clusters. In addition, we performed in situ hybridization with 49 selected genes based on the data from our transcriptome analysis. As a result, we identified multiple genes with localized expression in tissues characteristic of grass leaves. The present work provides a foundation for future analyses of genes with novel functions in rice leaf development.

**Results**

**Experimental design for microarray dataset**

Rice leaf ontogeny, i.e., the developmental process from initiation to maturation, is described in Itoh et al. (2005) [28]. Briefly, according to the staging system based on plastochron numbers (Pn), the P1 leaf primordium protrudes from the SAM and then grows to surround the SAM at stage P2. During the P1 and P2 stages, the leaf primordium consists of undifferentiated cells with no morphological characters. During the P3 stage, the boundary between the blade and sheath is established, and the future blade and sheath parts can be distinguished. In addition, the ligule primordium is formed in the boundary region at this stage. Although most of the P3 leaf primordium is comprised of undifferentiated cells, the outermost cells on the distal side of the primordium begin to differentiate into epidermal cells. During stage P4, the leaf blade elongates rapidly, and the difference between the blade and sheath becomes more pronounced. The P4 leaf primordium exhibits a clear gradient of cell states along its longitudinal axis; cells in the proximal region remain undifferentiated, whereas those in the distal region are differentiated. During stage P5, the leaf sheath elongates rapidly, and the growth and maturation of the leaf are completed by the P6 stage, whereas bending of the lamina joint occurs between stages P5 and P6.

To obtain a comprehensive transcriptome of leaf development in rice, we sampled 12 leaf parts representing various stages and components along the longitudinal axis (Fig. 1b–d; Table 1). Rice seedlings at the four-leaf stage were dissected into 12 parts: shoot apex containing the SAM and P1 and P2 leaf primordia (Fig. 1d); entire P3 leaf primordium (Fig. 1c); apical, middle, and proximal parts of P4 leaf blade; P4 leaf sheath; and the leaf blade, sheath, and boundary region of P5 and P6 leaves. Three biological replicates were prepared for each part, and their RNA was hybridized to a 44K rice microarray (Agilent Technologies, Santa Clara, CA) [26, 29]. Out of 43,144 probes corresponding to 29,864 genes on the rice 44K microarray platform, 31,996 probes corresponding to 24,022 genes were expressed in at least one sample. Normalized expression levels of these 31,996 probes corresponding to 24,022 genes were
used in this study. Pearson correlation analysis showed strong correlations among the three replicates, indicating that our dataset was highly reproducible (Supplemental Fig. 1).

**Transcriptome dynamics during rice leaf development**

To elucidate the transcriptome dynamics that occur during leaf development, principal component analysis (PCA) was performed using all samples. The first, second, and third principal components (PC1, PC2, and PC3) explained 60.9%, 13.2%, and 8.1% of the total variance among samples, respectively (Supplemental Fig. 2). Plotting the samples within the three-dimensional space defined by PC1, PC2, and PC3 allowed the relationships among the samples to be visualized, reflecting the two properties of tissue differentiation state and tissue identity (Supplemental Fig. 2a). However, these properties were poorly represented by each principal component (Supplemental Fig. 2b and c) due to the fact that PC2 was excessively attracted toward P4Bm, which was located distant from the other samples. Generally, PCA is not robust to outliers, and principal components tend to be attracted toward them, interfering with the detection of the overall dataset structure. To avoid excessive attraction of PC2 toward the outlier P4Bm, we modified PC1, PC2, and PC3 while retaining positional relationships between the samples, as follows. PC1, PC2, and PC3 scores were treated as variables, and PCA was again applied using all samples except P4Bm, resulting in modified principal components (mPC1, mPC2, and mPC3) that were not excessively affected by P4Bm. Altogether, mPC1, mPC2, and mPC3 explained 60.9%, 11.9%, and 9.5% of the total variance among samples, respectively, and successfully captured the characteristic patterns of the dataset (Fig. 2).

Two groups were separated by mPC1, namely, immature tissues represented by SA, P3, P4S, and P4Bb, mature tissues represented by P4Ba, and samples derived from P5 and P6 stage leaves. This result suggests that mPC1 represented the differences between immature and mature tissues (Fig. 2b). Conversely, mPC2 characterized samples with intermediate tissue differentiation, most notably P4Bm, suggesting that mPC2 represented the transient state of the transcriptome during tissue differentiation (Fig. 2b). Thus, an arrow fitting the distribution of the samples in the space defined by mPC1 and mPC2 would represent the change in transcriptome dynamics associated with tissue differentiation from the immature state through the transient state to the mature state. Collectively, mPC1 and mPC2 explained 72.8% of the total variance among samples, suggesting that tissue differentiation state has profound effects on the leaf transcriptome. Moreover, samples derived from the P4 leaf exhibited large transcriptomic variations, whereas all P4-stage leaf samples including sheath samples were aligned along the arrow. This result indicates that the shift in the transcriptome associated with leaf maturation is found throughout the leaf during P4, coinciding with intensive basipetal tissue differentiation at stage P4 [30].

In addition, mPC3 separated samples of the leaf blade P4Bm, P4Ba, P5B, and P6B from those of the leaf sheath and blade-sheath boundary region P5S, P6S, P5BS, and P6BS, indicating that mPC3 represented differences between the leaf blade and sheath (Fig. 2c). On the other hand, only slight differences were observed among immature leaf samples such as P3, P4S, and P4Bb, suggesting that the transcriptomic difference between the leaf blade and sheath becomes more pronounced during maturation.
Overall, our results suggest that the transcriptome of each part of the leaf changes with the progression of tissue differentiation and the acquisition of tissue identity.

**Gene expression patterns during leaf development and their associations with gene function and transcriptional regulation**

To uncover the major gene expression patterns during rice leaf development, we conducted cluster analysis of genes based on their expression patterns. Prior to cluster analysis, analysis of variance (ANOVA) was applied to detect differentially expressed genes among different parts of the leaf. Of 31,996 probes corresponding to 24,022 genes, 31,043 probes corresponding to 23,350 genes were extracted (p-value = 0.001 when adjusted for the false discovery rate [FDR]). K-means clustering was performed on these probes, and 28 clusters with distinct expression patterns were obtained (Supplemental Fig. 3; Supplemental Table 2). Consistent with the results of PCA, the expression pattern of each cluster was affected by differences in tissue differentiation state among samples. For Clusters 1 to 8, higher expression levels were found in samples containing immature tissues (Supplemental Fig. 3a). For Clusters 9 to 14, transiently elevated expression was found in samples with intermediate tissue differentiation states (Supplemental Fig. 3b). Genes in Clusters 15 to 24 were most strongly expressed in samples containing mature tissues (Supplemental Fig. 3c). For some of the clusters, there were large differences in expression levels among samples and characteristic expression patterns, suggesting that some groups of genes undergo similar changes in gene expression, and that such changes are associated with events during leaf development.

To evaluate how the gene expression patterns and dynamics of these gene clusters are related to the functions of the genes, we conducted Gene Ontology (GO) enrichment analysis on each cluster (Supplemental Fig. 4). In addition, given the importance of transcriptional regulation to development, transcription factors and transcriptional regulators were extracted from each cluster (Supplemental Fig. 5). These analyses identified the characteristic functions of genes within each cluster, including several genes that may be involved in specific processes during rice leaf development (Fig. 3; Table 2).

Cluster 1, a group of genes that was specifically expressed in the shoot apex, was enriched in genes involved in transcriptional regulation (regulation of transcription, p = 4.8e−07). Within this cluster were class I KNOX genes, which are important for SAM maintenance [31], and OsNAM/OsCUC3 genes, which may be involved in organ boundary formation [32]. Thus, Cluster 1 was predicted to include genes related to SAM function and leaf initiation. This cluster also included genes in the BBM clade of the PLETHORA family [33], which are expressed in crown root primordia [34], and OsTB1, which is expressed in axillary buds [35]. Because the shoot apex tissue used in this study contained stem tissue as well as the SAM and leaf primordia, the presence of root- and axillary bud-related genes in this cluster was not surprising.

Cluster 2 contains genes that were highly expressed in tissues undergoing active cell proliferation. In this cluster, GO terms associated with cell division and cytokinesis (microtubule-based movement, p = 8.6e−05) were detected. Moreover, it contained ANT clade genes of the PLETHORA family [33], GRF family genes [12], and OsGIF1/MKB3 [11], which have been described as promoters of cell proliferation in
leaf primordia. Thus, Cluster 2 was expected to contain important genes related to cell proliferation in leaf primordia.

Cluster 9 genes were highly expressed in the middle parts of the P4 leaf blade and P5 leaf sheath. GO analysis revealed that this cluster contains class III peroxidases (response to oxidative stress, p = 4.2e−07). Some class III peroxidases regulate reactive oxygen species homeostasis in the apoplast, thereby affecting cell-wall stiffness [36, 37]. GO terms for cell-wall remodeling enzymes including XTHs (carbohydrate metabolism, p = 1.3e−3) [38] were also enriched in Cluster 9. Thus, Cluster 9 appears to be enriched in genes involved in the control of cell-wall extensibility and cell elongation.

Cluster 12 genes were mostly expressed at high levels in the mature leaf blade. GO terms for genes involved in photosynthesis (photosynthesis, p = 7.9e−21; carbon utilization by fixation of carbon dioxide, p = 8.5e−10; and electron transport, p = 3.3e−07) were enriched. This cluster included C2C2-CO-like family genes [39] and phytochrome-interacting bHLH factors (PIFs) [40]. Thus, Cluster 12 was expected to contain a high concentration of genes associated with photosynthesis and light-mediated signal transduction.

Cluster 7 genes were highly expressed in immature samples and samples from mature tissues from the sheath and blade-sheath boundary region. This cluster was enriched in genes involved in carbohydrate metabolism (main pathways of carbohydrate metabolism, p = 3.7e−05). The leaf sheath is believed to act as sink tissue for carbohydrates prior to heading [41]. In addition, this cluster includes OsBOP genes that are important for sheath development [5]. Thus, Cluster 7 was enriched in genes involved in the carbohydrate sink function and sheath development.

Cluster 15 consisted of genes that were preferentially expressed in the mature sheath and blade-sheath boundary region. This cluster was enriched in genes related to GO terms for protein kinases (protein amino acid phosphorylation, p = 6.2e−05). The blade-sheath boundary contains the lamina joint, which bends between stages P5 and P6. Various phytohormones and environmental stressors affect the bending process [42, 43], and many protein kinases exhibit temporal changes in expression during lamina-joint bending [25]. Thus, we assumed that Cluster 15 contains genes involved in lamina-joint bending.

Taken together, these analyses revealed genome-wide gene expression patterns during rice leaf development, and relationships between expression pattern and function were found in certain gene clusters.

Identification of genes with localized expression during leaf development

To identify novel genes that play important roles in early development and subsequent morphogenesis and tissue formation in the rice leaf, we selected genes from Clusters 1 to 8 using K-means clustering analysis. These clusters contain genes that tended to be highly expressed during the early stages and weakly expressed during the later stages, and thus were expected to include candidate genes. Forty-nine
genes, most of them involved in transcriptional regulation, were selected from the gene clusters, and their spatial expression patterns around the shoot apex were examined through in situ hybridization. Examples of these gene expression patterns are described below.

Cluster 2 included the PLATZ family transcription factor Os02g0172800, which is a co-ortholog of ORESARA15[44]. This gene was expressed in the basal part of immature leaves, and its expression was strongest in the abaxial side of the leaf primordium (Fig. 4a, b).

Tissue-specific expression patterns of three genes in Cluster 3 were detected. Expression of OsbHLH080, a member of the bHLH gene family, was observed in the abaxial base of leaf primordia and developing ligules (Fig. 4c). Another bHLH gene, OsbHLH166, was expressed mainly in the presumptive lysigenous aerenchyma areas of leaf primordia (Fig. 4d, e). Os05g0363500, which encodes a WD40 repeat-containing protein, was expressed in the developing diaphragms of leaf sheaths (Fig. 4f).

OsGH3-4, a member of the GH3 family involved in auxin conjugation, was placed in Cluster 7 and exhibited elevated expression levels mainly in the leaf sheath and blade-sheath boundary region. Expression of OsGH3-4 was detected in both the central domain of the SAM and tissues adaxially adjacent to vascular bundles in the leaf sheath and midrib, where bundle sheath extension cells differentiate (Fig. 4g, h).

Additionally, Cluster 7 contained four OsARF paralogs belonging to the ARF6/8 subfamily [45]. These genes, OsARF6/12/17/25, were expressed at the basal part of the leaf primordium around stage P3, with especially high expression levels at the margin (Fig. 5a–d, arrowheads). Moreover, these four OsARFs were also expressed in the developing ligule and marginal parts of the blade-sheath boundary region that were expected to differentiate into auricles (Fig. 5a–d). In addition to these common expression patterns among the four OsARFs, the characteristic expression of each gene was also observed. OsARF6 was expressed in the epidermis of the basal region of P4 leaf blades (Fig. 5a), whereas OsARF12 was expressed throughout the sheath and basal region of the blades (Fig. 5b). OsARF17 expression was detected in the adaxial epidermis at the blade-sheath boundary at stage P3 and in the lamina joint of P4 leaves (Fig. 5c), whereas OsARF25 was strongly expressed in the lamina joint of P4 leaves and at the base of sheaths in stages P4 and P5 (Fig. 5d).

In addition, ARF6/8 orthologs including the four OsARFs listed above are known targets of miR167, which is an evolutionarily conserved microRNA in seed plants [45]. To clarify the relationships between the expression patterns of the four OsARFs and miR167, the accumulation patterns of mature miR167 in the shoot apex were examined through in situ hybridization using a probe containing BNA\textsuperscript{NC}, which is a bridged nucleic acid derivative of a locked nucleic acid (Fig. 5e). Signals specific to mature miR167 were detected at higher levels in the distal part and lower levels in the basal part, indicating that the accumulation of miR167 and the expression of the four identified OsARFs were largely exclusive.

In addition to the genes described above, we explored a number of genes with localized expression during leaf development based on the list obtained through K-means clustering (Supplemental Fig. 6). Therefore,
our strategy of using K-means clustering facilitated the selection of genes that were differentially expressed among stages or tissues. In addition, the genes identified here should be further analyzed to uncover their functions in rice leaf development.

Discussion

Leaf primordia, which originate from the flank of the SAM, undergo various developmental processes from differentiation to maturation. Leaf primordia of grasses develop distinct regions along the longitudinal axis, and each region differentiates into a morphologically unique structure. Tissue differentiation in grass leaves during development proceeds in a basipetal direction, which has been extensively studied using the maize developing leaf blade as a model [16, 46, 47, 48, 49, 50, 51, 52, 53]. These studies revealed that basipetal tissue differentiation is accompanied by dynamic changes in mRNA, proteins, and various metabolites. However, the resolution of these studies is limited spatially and temporally, as only parts of a single developing leaf blade at a specific stage were used, whereas other stages and tissues were not examined. Thus, our study is unique in terms of providing genome-wide expression profiles both across developmental stages and distinct tissues along the longitudinal axis.

In this study, the PCA results suggest that the leaf transcriptome dynamically changes among stages of tissue differentiation, as shown in the space defined by mPC1 and mPC2 (Fig. 2b). On the other hand, the difference between the leaf blade and sheath, which was represented by mPC3, accounted for less transcriptomic variation among samples than did mPC1 or mPC2 (Fig. 2c), suggesting that organ identity has a smaller effect on the leaf transcriptome than tissue differentiation. In particular, marked transcriptomic changes were observed in the P4 leaf along its longitudinal axis (Fig. 2b). The transcriptomes of the proximal parts, P4S and P4Bb, resembled that of the leaf primordium in early stages, such as P3. On the other hand, the distal part (P4Ba) exhibited transcriptomic similarities to leaf parts at later stages, such as P5 and P6. Thus, our results suggest that P4 is the stage at which tissue differentiation proceeds in the basipetal direction along the whole leaf and dynamic transcriptome changes occur.

Along the longitudinal axis of developing grass leaves, it has been suggested that cell proliferation and elongation occur actively in the basal and middle regions [54] and that photosynthetic activity and related gene expression is high in the apical region [16, 52]. Our cluster and GO analyses revealed that the genes in Clusters 2, 9, and 12 may be involved in cell proliferation, cell elongation, and photosynthesis, respectively (Fig. 3; Table 2). The expression patterns of these clusters in various parts of the P4 leaf are consistent with previous findings; namely, different developmental events occur in a single leaf primordium along its longitudinal axis. Thus, P4 is the stage wherein drastic developmental reprogramming occurs in rice. Moreover, acquisition of photosynthetic competence is initiated between the P3 and P4 stages [24], which is in accordance with our finding that genes in Cluster 12 were weakly expressed at stage P3 and strongly expressed at stage P4 (Fig. 3; Table 2). In addition to these clusters, we identified Cluster 7 genes as being highly expressed in the sheath and blade-sheath boundary regions (Fig. 3; Table 2). OsBOP genes, which are regulators of leaf sheath identity, were included in this cluster and
highly expressed at stage P3, suggesting that a number of upstream or downstream genes exhibit polarized expression patterns along the longitudinal axis from an early stage of leaf development.

The rice leaf has distinct morphological features that are not present in Arabidopsis. The development of these features is likely controlled in part by genetic mechanisms unique to grasses. Previous studies have identified several genetic factors in the morphogenesis of characteristic organs in grass leaves. LIGULELESS1 (LG1) is required for the development of structures at the boundary between the blade and sheath [2, 3, 4]. DROOPING LEAF is important for the development of the midrib [55, 56]. However, most of the genetic factors and their networks that underpin grass leaf morphogenesis remain unknown.

In general, to explore novel genes that play roles in the developmental process of interest, an efficient screening method is required. Expression profiling is one practical method of gene discovery. Transcriptome analysis and in situ hybridization are representative methods of expression analysis with different advantages. Transcriptome analysis can provide global expression profiles at the genome-wide scale, whereas in situ hybridization can reveal the spatial expression pattern of a gene at the tissue and cellular levels. By combining these methods, many studies have attempted to identify novel genes that function in a developmental process of interest [15, 22, 57, 58, 59, 60, 61, 62, 63, 64].

In this study, we identified a number of genes with localized expression patterns that may be associated with morphological features unique to rice and grasses. For example, Os02g0172800 was expressed in immature leaf primordia at higher levels in the abaxial side of the basal part of the leaf primordia (Fig. 4a, b). This expression pattern is similar to that of MKB3, which is a positive regulator of cell proliferation in leaf primordia [11]. Furthermore, Os02g0172800 is a co-ortholog of ORESARA15, which reportedly promotes cell proliferation through a genetic pathway mediated by AN3, an ortholog of MKB3 [44]. It has been suggested that the formation of the leaf sheath requires a decreasing gradient of cell proliferation from the abaxial side of the leaf primordium to the adaxial side, which may be associated with the expression pattern of MKB3 [11]. Thus, Os02g0172800 might be also involved in cell proliferation in leaf primordia, and its expression pattern might reflect the unique cell proliferation pattern during the development of the leaf sheath.

Expression of OsbHLH80 was detected in the developing ligule (Fig. 4c). Ligule development was found to be disrupted by dysfunction of brassinosteroid (BR) signaling [65]. OsBC1, a paralog of OsbHLH80, is considered to regulate lamina-joint bending in response to BRs [66]. Thus, characterizing the roles of OsbHLH80 in ligule development in association with BR signaling is a worthwhile topic for future research.

OsGH3-4 was expressed in tissues adaxially adjacent to vascular bundles in the leaf sheath and midrib, where bundle sheath extension cell differentiation occurs (Fig. 4g, h). In addition, tissue-specific expression of OsbHLH166 and Os05g0363500 were observed in the presumptive region of lysigenous aerenchyma and developing diaphragms, respectively (Fig. 4d–f). During the development of the leaf sheath and midrib, a group of parenchyma cells collapse to form lysigenous aerenchyma, whereas parenchyma cells adaxially adjacent to the vascular bundles remain intact and differentiate into bundle
sheath extension cells. Additionally, some parenchyma cells transform into stellate parenchyma cells in diaphragms that vertically separate the lysigenous aerenchyma [67]. Although these structures are well-developed in the leaves of rice and some wetland plants, the genetic networks regulating the development of these structures remain unknown. It has been demonstrated that the formation of constitutive aerenchyma in rice roots is regulated by auxin signaling [68]. Thus, OsGH3-4 may play an important role in the patterning of bundle sheath extension cells through the modulation of the spatial patterns of auxin accumulation in leaf primordia. Functional analysis of OsbHLH166 and Os05g0363500 would provide insights into the developmental programs underlying the formation of aerenchyma and elaboration of the unique cell shape of stellate parenchyma cells forming diaphragms.

OsARF6/12/17/25 exhibited polarized expression along the longitudinal axis and was expressed in developing ligules as well as the marginal parts of the blade-sheath boundary, where auricle differentiation occurs (Fig. 5a–d). Moreover, OsARF17 exhibited a unique expression pattern in the adaxial epidermis of the blade-sheath boundary in leaf primordia (Fig. 5c). Similarly, OsLG1 exhibited localized expression in marginal parts of the blade-sheath boundary (Supplemental Fig. 7). Recent research suggested that a member of the ARF6/8 subfamily acts downstream of the OsLG1 ortholog in wheat [4]. Thus, these genes may be important for the development of organs in the blade-sheath boundary region. Furthermore, OsARF25 exhibited strong expression in the leaf-sheath pulvinus, which is a gravitropic tissue in the leaf base that is involved in shoot bending in grasses [69] (Fig. 5d). This expression pattern suggests that OsARF25 may play an important role in the bending capability of shoots at the leaf-sheath pulvinus in response to gravity. In addition to the leaf-sheath pulvinus, OsARF25 was strongly expressed in the lamina joint (Fig. 5d). Several genes controlling shoot bending were previously reported to be expressed in both the lamina joint and leaf-sheath pulvinus [70, 71, 72]. These gene expression patterns suggest that the lamina joint and leaf-sheath pulvinus use similar genetic programs to achieve bending.

Post-transcriptional regulation by miR167 has been reported to be required for attaining the correct spatial expression patterns of ARF6/8 during reproductive development in Arabidopsis [73]. Our analyses revealed mutually exclusive expression of miR167 and OsARF6/12/17/25 along the longitudinal axis of leaf primordia. This result indicates that miR167 post-transcriptionally downregulates OsARF6/12/17/25 expression in the distal part of the leaf primordium (Fig. 5). A similar accumulation pattern for mature miR167 was reported in Arabidopsis cotyledons [74]. Thus, post-transcriptional regulation of ARF6/8 orthologs by miR167 in the distal part of the leaf primordium may be evolutionally conserved between rice and Arabidopsis.

Our microarray analysis had some limitations in resolution despite the sampling of extensive leaf stages and regions. For example, the sample SA contained the SAM, P1 and P2 leaf primordia, and immature stem tissues below the shoot apex, which makes it impossible to distinguish differences in expression profiles among these tissues (Fig. 1d). To improve our transcriptomic resolution, the separation of small tissues through laser microdissection, as performed in maize transcriptome studies on different domains of the SAM [15, 22], may facilitate more detailed gene expression profiling at the early stages of rice leaf
development. In addition, the 44K microarray platform used in this study does not cover all transcripts in the rice genome. RNA sequencing analysis is required to obtain the expression profiles of all transcripts and identify a greater number of novel genes related to leaf development. However, our expression data are indispensable for elucidating the global transcriptome and true nature of individual gene expression levels in rice due to their compatibility with various datasets in the expression profile database Ricexpro (http://ricexpro.dna.affrc.go.jp) [26, 27].

Conclusions

We identified a large number of genes that exhibit localized and unique expression patterns during rice leaf development (Figs. 4 and 5; Supplemental Fig. 6). Due to the recent development of CRISPR/Cas9 technology, any genes in the rice genome can be easily knocked out [76]. Thus, reverse genetics strategies to reveal gene function can be applied to the genes identified in this study. Our findings will provide the foundation for future research on the development of grass leaves and contribute to the elucidation of genetic programs unique to grasses.

Methods

Plant materials and growth conditions

Rice (Oryza sativa L. ssp. japonica cv. Nipponbare) seeds were obtained from the National Agriculture and Food Research Organization. They were sown in germination boxes, and the seedlings were grown in a growth chamber (14-h light period at 30 °C and 10-h dark period at 25 °C). At 14–15 days after germination, seedlings in which the tip of the fifth leaf had just emerged from the fourth leaf were collected. The seedlings were dissected under a dissecting microscope to separate tissues at different developmental stages and locations along the longitudinal axis of the leaves. The dissected tissues were used for RNA extraction of microarray analysis.

RNA extraction and microarray analysis

Total RNA was extracted from the collected samples using a RNeasy Mini Kit (Qiagen, Hilden, Germany), and labeling was performed using Quick Amp Labeling Kit, One-Color (Agilent Technologies) in the presence of cyanine-3 (Cy3)-CTP according to the manufacturer’s protocol. The resulting Cy3-labeled cRNA was purified using a RNeasy Mini Kit (Qiagen, Hilden, Germany). A total of 1000 ng Cy3-labeled cRNA was fragmented and hybridized onto a slide of the rice 4×44K microarray RAP-DB (G2519F#15241; Agilent Technologies). After washing, the slides were scanned on an Agilent G2505B DNA microarray scanner, and background correction of the raw Cy3 signals was performed using Feature Extraction 10.5.1.1 software (Agilent Technologies).

Statistical analysis
The processed raw signal intensities of all probes were transformed to log$_2$ scale, and normalization procedures were performed, including 75$^{th}$-percentile normalization for inter-array comparison and baseline correlation to the median value within each dataset, using GeneSpring GX12 software (Agilent Technologies). In this study, we used 31,996 probes corresponding to 24,022 loci, which had raw signal intensities >50 in at least one of the 36 microarray datasets. Pearson correlation analysis was performed on all samples using the ggcorrplot package in R [77]. PCA based on the variance-covariance matrix was first performed on all samples using the prcomp function in R. Subsequently, PCA was applied again to all samples except for P4Bm, using the PC1, PC2, and PC3 scores as variables to compute modified principal components (mPC1, mPC2, and mPC3). The distribution of the samples in the space defined by mPC1 and mPC2 was approximated with a quadratic formula using the nls function in R. One-way ANOVA with Benjamini and Hochberg FDR correction was applied to detect the probes that were differentially expressed among samples (FDR-adjusted p-value = 0.001) using GeneSpring GX12. The extracted probes were classified into 28 clusters through K-means cluster analysis based on Euclidean distance using the cclust package in R [78]. GO enrichment analysis was performed on the genes in each cluster using a tool based on a rice gene coexpression database, RiceFREND [75]. GO terms corresponding to fewer than five genes were discarded. As GO terms are organized in a hierarchical structure, only the child terms were retained. Transcription factor and transcriptional regulator genes were extracted from each cluster using the list in the Plant Transcription Factor Database [79]. In addition, the enrichment of each gene family in each cluster was statistically tested by comparing the number of family members in each cluster with that obtained from the rice 44K microarray platform, using one-sided Fisher's exact tests with Benjamini and Hochberg FDR correction with the fisher.test and p.adjust functions in R. The expression profiles shown in Supplemental Fig. 3 and Fig. 3, and the heatmaps in Supplemental Figs. 4 and 5 were drawn using the ggplot2 package in R [80].

**Gene annotation used for analyses**

The gene annotation used in this study was obtained from RAP-DB [29]. The list of genes associated with the biosynthesis, catabolism, and signaling of six phytohormones was derived from Hirano et al. (2008) [81]. For the extraction of transcription factor and transcriptional regulator genes from the rice genome, the list from the Plant Transcription Factor Database was used.

**In situ hybridization**

Rice (*O. sativa* ssp. japonica cv. Nipponbare) seeds were obtained from the National Agriculture and Food Research Organization. They were sown in soil, and the seedlings were grown in a growth chamber (14-h light period at 30 °C and 10-h dark period at 25 °C). At 14–15 days after germination, the shoot apex of each four-leaf stage seedling was dissected and fixed in 4% paraformaldehyde in 0.1 M sodium phosphate buffer for 24 h at 4 °C, and then dehydrated in a graded ethanol series. The ethanol was replaced with Histo-Clear (National Diagnostics, Atlanta, GA), and the samples were embedded in Paraplast Plus (Leica, Wetzlar, Germany). Paraffin sections (thickness, 8 mm) were placed on microscope slides and coated with 3-aminopropyl triethoxysilane (Matsunami Glass, Osaka, Japan). To generate
probes for 30 genes (see Supplemental Table 3), the corresponding full-length cDNA clones were obtained from NIAS Genebank [82] and used as templates. For 20 genes, cDNA fragments were amplified using PCR and cloned into the pCR Blunt II TOPO vector (Invitrogen, Carlsbad, CA) using primers specific to each gene (Supplemental Table 3). The cDNA was amplified via PCR or digested with restriction enzymes and then transcribed using digoxigenin-labeled antisense riboprobes using a MAXIscript In Vitro Transcription kit (Life Technologies, Carlsbad, CA) with digoxigenin-11-UTP (Roche, Basel, Switzerland), or using T7, SP6 (Takara, Shiga, Japan), or T3 (Roche) RNA polymerase with DIG-RNA labeling mix (Roche). For detecting mature miR167, a 3'-end digoxigenin-labeled probe with the sequence 5' - AgAtCaTgCtGgCaGcTtCa-3' was used, where uppercase and lowercase letters represent BNA NC[NMe] and DNA, respectively. In situ hybridization and immunological detection of the hybridization signals were performed as described by Kouchi and Hata (1993) [83].

**Abbreviations**

SPL: *SQUAMOSA PROMOTER BINDING-LIKE*

SAM: Shoot apical meristem

GRF: *GROWTH-REGULATING FACTORS*

GIF: *GRF-INTERACTING FACTORS*

Pn: Plastchron number

PCA: Principal component analysis

ANOVA: Analysis of variance

FDR: False discovery rate

GO: Gene ontology

PIF: Phytochrome-interacting bHLH factors

BNA NC: Bridged nucleic acid

LG1: *LIGULELESS1*

BR: Brassinosteroid

**Declarations**

**Ethics approval and consent to participate**

Not applicable.
Consent for publication

Not applicable.

Availability of data and materials

The datasets generated and analyzed during the current study are available in the GEO repository through accession number GSE159047 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE159047].

Competing interests

No conflicts of interest declared.

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Author's contributions

MM and JI designed experiments; MM, TY, YS and JI performed experiments; MM and JI wrote the article. All authors have read and approved the manuscript.

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**Tables**
Table 1
Description of samples used for microarray analysis

| Sample               | Stage  | Tissue                                           | Abbreviation |
|----------------------|--------|--------------------------------------------------|--------------|
| Shoot Apex           | P0, P1, P2 | Shoot apex containing SAM and P1 and P2 leaf primordia | SA           |
| P3                   | P3     | Whole P3 leaf primordia                          | P3           |
| P4Sheath             | P4     | Leaf sheath                                      | P4S          |
| P4Blade_basal        | P4     | Basal part of leaf blade                         | P4Bb         |
| P4Blade_middle       | P4     | Middle part of leaf blade                        | P4Bm         |
| P4Blade_apical       | P4     | Apical part of leaf blade                        | P4Ba         |
| P5Sheath             | P5     | Leaf sheath                                      | P5S          |
| P5Boundary           | P5     | Boundary region between leaf blade and sheath    | P5BS         |
| P5Blade              | P5     | Leaf blade                                       | P5B          |
| P6Sheath             | P6     | Leaf sheath                                      | P6S          |
| P6Boundary           | P6     | Boundary region between leaf blade and sheath    | P6BS         |
| P6Blade              | P6     | Leaf blade                                       | P6B          |
Table 2

| Cluster   | Enriched GO terms                  | TF/TRs                                                                 |
|-----------|------------------------------------|------------------------------------------------------------------------|
| Cluster1  | regulation of transcription,       | OSH1/6/15/71 (Class I KNOX), OsNAM/OsCUC3, OsPLT2/3/4/5/6 (BBM clade PLETHORA), OsTB1 |
|           | p=4.8e-07                          |                                                                        |
| Cluster2  | microtubule-based movement,        | OsPLT1/7/8/9 (ANT clade PLETHORA), OsGRF1/6/7/9/10, OsGIF1/MKB3        |
|           | p=8.6e-05                          |                                                                        |
| Cluster7  | main pathways of carbohydrate      | OsBOP1/2/3                                                             |
|           | metabolism, p=3.7e-05               |                                                                        |
| Cluster9  | response to oxidative stress,      |                                                                        |
|           | p=4.2e-07                          |                                                                        |
| Cluster12 | photosynthesis, p=7.9e-21          | OsBBX8/10/12/17 (C2C2-CO-like), OsPIL12/13 (PIF)                        |
|           | carbon utilization by fixation of   |                                                                        |
|           | carbon dioxide, p=8.5e-10           |                                                                        |
|           | electron transport, p=3.3e-07       |                                                                        |
| Cluster15 | protein amino acid phosphorylation,|                                                                        |
|           | p=6.2e-05                          |                                                                        |

See Supplemental Figs. 4 and 5 for enriched GO terms and TF/TRs in all 28 clusters obtained through K-means clustering, respectively.