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GLABRA2 Directly Suppresses Basic Helix-Loop-Helix Transcription Factor Genes with Diverse Functions in Root Hair Development

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Abstract

The *Arabidopsis thaliana GLABRA2* (*GL2*) gene encodes a transcription factor involved in the cell differentiation of various epidermal tissues. During root hair pattern formation, *GL2* suppresses root hair development in non-hair cells, acting as a node between the gene regulatory networks for cell fate determination and cell differentiation. Despite the importance of *GL2* function, its molecular basis remains obscure because the GL2 target genes leading to the network for cell differentiation are unknown. We identified five basic helix-loop-helix (bHLH) transcription factor genes—*ROOT HAIR DEFECTIVE6* (*RHD6*), *RHD6-LIKE1* (*RSL1*), *RSL2, LjRHL1-LIKE1* (*LRL1*), and *LRL2*—as GL2 direct targets using transcriptional and post-translational induction systems. Chromatin immunoprecipitation analysis confirmed GL2 binding to upstream regions of these genes *in planta*. Reporter gene analyses showed that these genes are expressed in various stages of root hair development and are suppressed by *GL2* in non-hair cells. *GL2* promoter-driven green fluorescent protein fusions of LRL1 and LRL2, but not those of the other bHLH proteins, conferred root hair development on non-hair cells. These results indicate that GL2 directly suppresses bHLH genes with diverse functions in root hair development.

INTRODUCTION

The morphological differentiation and deposition patterns of cells are crucial determinants of functional plant structures. Among the plant tissues composed of multiple types of cells, the root epidermis of *Arabidopsis thaliana* has served as an excellent model system for studying the morphological differentiation and pattern formation of plant cells (Dolan et al.,

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1993; Schiefelbein, 2000; Tominaga-Wada et al., 2011). In *Arabidopsis*, the root epidermis is composed of non-hair (N) and hair (H) cell files which are in contact with either one or two underlying cortical cell file(s), respectively, and only the cells in H cell files develop root hairs (Dolan et al., 1994; Galway et al., 1994; Clowes, 2000; Schiefelbein et al., 2009). A mechanism that converts a positional cue from the underlying structure to a signal directing the root hair pattern is assumed to involve a receptor-like kinase expressed in the root epidermis (Kwak et al., 2005; Kwak and Schiefelbein, 2007; Kwak and Schiefelbein, 2008). Downstream of this mechanism, numerous transcription factor genes, including *GLABRA2* (*GL2*), constitute regulatory networks responsible for cell pattern formation and subsequent cell differentiation (Schiefelbein et al., 2009; Tominaga-Wada et al., 2011; Grebe, 2012; Schiefelbein et al., 2014).

GL2 encodes a homeodomain-leucine-zipper transcription factor and is thought to be a negative regulator of root hair development because it is expressed preferentially in N cell files and because mutant defects in *GL2* result in ectopic root hair formation in N cell files (Rerie et al., 1994; Di Cristina et al., 1996; Masucci et al., 1996). *GL2* is activated in a cell file-specific manner by a transcription factor complex consisting of the WD40 protein TRANSPARENT TESTA GLABRA1 (TTG1), the basic helix-loop-helix (bHLH) protein GLABRA3 (GL3) or ENHANCER OF GLABRA3 (EGL3), and the R2R3-type MYB protein WEREWOLF (WER) or MYB23 (Galway et al., 1994; Masucci et al., 1996; Lee and Schiefelbein, 1999; Payne et al., 2000; Bernhardt et al., 2003, 2005; Zhang et al., 2003; Kang et al., 2009). The complex also transcriptionally activates the R3-type MYB protein gene *CAPRICE* (*CPC*) and its paralogs in N cell files (Wada et al., 1997; Schellmann et al., 2002; Esch et al., 2004; Kirik et al., 2004; Simon et al., 2007; Tominaga et al., 2008). However, these R3-type MYB proteins move to H cell files to negatively regulate the transcription factor complex by replacing WER and MYB23, thereby suppressing the expression of *GL2* with consequent root hair development in H cell files (Schellmann et al., 2002; Bernhardt et al., 2003; Esch et al., 2004; Kirik et al., 2004; Kurata et al., 2005; Tominaga et al., 2007, 2008). Consistent with this gene regulatory network, mutant defects in the transcription factor complex and its negative regulators confer ectopic root hair and scarce root hair phenotypes, respectively, and defects in *GL2* are epistatic to them with respect to root hair development (Wada et al., 1997; Hung et al., 1998; Lee and Schiefelbein, 1999; Bernhardt et al., 2005; Wang *et al.*, 2010). Furthermore, because the expression of a modified GL2 with a constitutive transactivating function (VP16-GL2 $\,$ N) resulted in the formation of root hair-like structures from various epidermal cells, including leaf pavement cells (Ohashi et al., 2003), GL2 is thought to recognizes a set of genes that are sufficient for root hair development. Unlike root hair development, the file-specific cell fate of the root epidermis is thought to be determined independently of *GL2* function because the characteristics of H and N cell files in cell length and vacuolation timing, which are affected in *ttg* mutants, are normal in *gl2* mutants (Galway et al., 1994; Masucci et al., 1996). This line of evidence clarifies the role of *GL2* as a genetic switch that suppresses root hair development between cell fate determination and cell differentiation during root hair pattern formation.

A regulatory network that acts downstream of *GL2* for root hair cell differentiation comprises many bHLH transcription factor genes (Bruex et al., 2012; Pires et al., 2013). Mutants of the *ROOT HAIR DEFECTIVE6* (*RHD6*)/*bHLH83* gene, which encodes a transcription factor belonging to the bHLH VIIIc subfamily (Heim et al., 2003), have fewer root hairs than the wild type (Masucci and Schiefelbein, 1994). *RHD6* expression is normally specific to H cell files (Masucci and Schiefelbein, 1994), but extends to include N cell files in *gl2* mutants (Menand et al., 2007) suggesting that GL2 suppresses *RHD6* expression directly or indirectly in N cell files. *RHD6 LIKE1* (*RSL1*)/*bHLH86*, the closest paralog of *RHD6*, has a function partly redundant with that of *RHD6* in the initiation of root hair development (Menand et al., 2007). Other VIIIc subfamily genes, *RSL2*/*bHLH85* and *RSL4*/*bHLH54*, are functionally redundant with each other in root hair elongation (Yi et al., 2010). Post-translational induction experiments using a modified RHD6 protein with the glucocorticoid receptor (GR) domain showed that RHD6 activated the *RSL4* and *RSL2* genes directly and indirectly, respectively (Yi et al., 2010). *Lotus japonicus* mutants of the transcription factor gene *ROOTHAIRLESS1* (*LjRHL1*) exhibited severe defects in root hair development (Karas et al., 2009). Cross-species complementation analysis showed that the *Arabidopsis* bHLH XI subfamily genes *LjRHL1-LIKE1* (*LRL1*)/*bHLH66, LRL2*/*bHLH69* and *LRL3*/*bHLH82* (Heim et al., 2003) are functional equivalents of *LjRHL1* (Karas et al., 2009). Multiple mutants of these genes have shorter root hairs than the wild type (Karas et al., 2009), while single mutants exhibited only moderate phenotypes in root hair length and morphology (Bruex et al., 2012) suggesting redundancy of their functions in root hair elongation. The transcript levels of *LRL3*, but not those of *LRL1* or *LRL2*, were affected markedly in the *rhd6-3 rsl1-1* double mutant (Karas et al., 2009) indicating that *LRL3* is regulated differently from *LRL1* and *LRL2*.

Despite the pivotal role of *GL2* in suppressing root hair development, the molecular basis of the suppression remains obscure because, to date, few GL2 target genes are known and none of them are connected to the gene regulatory network for root hair cell differentiation. The *PHOSPHOLIPASE D*ζ*1* (*PLD*ζ*1*) gene is suppressed directly by GL2 in N cell files and its ectopic expression causes root hair bulges in N cell files, suggesting its involvement in promoting root hair development (Ohashi et al., 2003). *CELLULOSE SYNTHASE5* (*CESA5*) and *XYLOGLUCAN ENDOTRANSGLYCOSYLASE17* (*XTH17*), both of which encode enzymes involved in polysaccharide synthesis, are also GL2 target genes, but their roles in root hair development are unclear (Tominaga-Wada et al., 2009). *GL2* and *MYB23* constitute a positive feedback loop in the shoot epidermis where GL2 directly recognizes *MYB23* (Khosla et al., 2014). Recently, GL2 was shown to negatively regulate anthocyanin biosynthesis through direct recognition of the bHLH IIIf subfamily gene *TRANSPARENT TEST8* (*TT8*) and the R2R3-type MYB genes *PRODUCTION OF ANTHOCYANIN PIGMENT2* (*PAP2*) and *MYB113* (Wang et al., 2015). Upstream regions of these genes typically contain the L1 box-like sequence 5'-TAAATGT-3' (Abe et al., 2001; Tominaga-Wada et al., 2009; Lin and Aoyama, 2012; Khosla et al., 2014; Wang et al., 2015). Although GL2 physically interacted with DNA regions containing this sequence (Ohashi et al., 2003; Khosla et al., 2014; Wang et al., 2015), whether the sequence is necessary and/or sufficient for recognition by GL2 *in planta* remains unclear.

In this study, we identified five bHLH transcription factor genes, *RHD6, RSL1, RSL2, LRL1*, and *LRL2*, as direct targets of GL2 using transcriptional and post-translational induction systems with the chimeric transcription factors VP16-GL2 N and GR-VP16-GL2 N, respectively. Chromatin immunoprecipitation (ChIP) analysis using a green fluorescence protein (GFP)-fused GL2 protein confirmed the physical interaction of GL2 with these genes *in planta*. Expression analyses showed that they are suppressed by *GL2* in N cell files and are expressed in various developmental stages of root hair development. Moreover, the phenotypes caused by the ectopic expression of their GFP-fusion proteins differed among them. These results indicate that GL2 acts as a negative factor for root hair development via multimodal pathways by targeting bHLH genes with diverse functions.

RESULTS

RHD6, RSL1, RSL2, LRL1, and LRL2 Are Candidate GL2 Target Genes

We performed a microarray screening to identify candidate GL2 target genes that were transcriptionally activated in response to the expression of the modified GL2 VP16-GL2 N, which has a constitutive transactivating function (Ohashi et al., 2003), in a glucocorticoidinducible gene expression system using the GVG transcription factor (Figure 1A). Triplicate microarray experiments using total RNA from seedlings harboring the GVG-inducible VP16-GL2 N gene detected 864 genes the steady-state transcript levels of which were consistently increased by more than two fold after dexamethasone (DEX) treatment (Supplemental Table 1). These candidate genes were categorized by gene product functions as defined using the agriGO web-based software tool (Du et al., 2010). The top five groups with most significant enrichment in the candidate target genes are associated with lipid metabolic processes, regulation of RNA metabolic processes, regulation of transcription, cell wall modification, and hormone-mediated signaling pathways, respectively (Figure 1B). Of these, the group involved in the regulation of transcription contained the largest number of genes, suggesting the presence of a large transcriptional network downstream of *GL2*. We subsequently focused on the transcription factor genes *RHD6* and *LRL1* because they are known to be involved in root hair development (Masucci and Schiefelbein, 1994; Menand et al., 2007; Karas et al., 2009; Bruex et al., 2012).

RHD6 and *LRL1* belong to the bHLH VIIIc and XI subfamilies, respectively (Heim et al., 2003). Other members of these subfamilies, *RSL4, LRL3, bHLH7*, and *bHLH59* were not induced significantly, and probes for *RSL1, RSL2, LRL2* and *RSL3*/*bHLH84* were absent from the microarray used in this study (Table 1). We examined the transcript levels of all the bHLH VIIIc and XI subfamily genes, together with that of *PLD*ζ*1* as a positive control, individually by quantitative real-time reverse transcription polymerase chain reaction (qRT-PCR) analysis using the same RNA preparations as used in the microarray experiments. The transcript levels of *RHD6, RSL1, RSL2, LRL1*, and *LRL2*, but none of the other VIIIc or XI subfamily genes, were induced significantly by the DEX treatment (Figure 1C). Therefore, we analyzed the five bHLH genes further as candidate GL2 target genes.

GL2 Recognizes RHD6, RSL1, RSL2, LRL1, and LRL2 Directly

To examine the direct recognition of the five bHLH genes by GL2 *in planta*, we used a posttranslational induction system with a GR domain-fused transcription factor. Transgenic seedlings harboring the 35S promoter-driven GR-VP16-GL2 $\,$ N gene (Figure 2A) were treated or not treated with DEX for 1–4 hours in the presence of cycloheximide (CHX), a protein synthesis inhibitor that prevents the secondary induction of transcription by the protein products of GL2 target genes. Following the treatment, qRT-PCR analysis using total RNA from the transgenic roots showed that the transcript levels of the five bHLH genes were increased by more than five fold, while the time required for the maximum induction differed for each bHLH gene (Figure 2B). In contrast, the non-candidate gene *LRL3* was not activated significantly by DEX treatment (Figure 2B). These results indicate that GR-VP16-GL2 N directly recognized the five bHLH genes through the GL2 DNAbinding domain, and that GL2 can also recognize these genes directly.

Based on previous findings suggesting that GL2 recognizes the L1 box-like sequence 5'- TAAATGT-3' (Lin and Aoyama, 2012), we searched the 3-kb regions upstream of the initiation codons of the five bHLH genes and identified at least one L1 box-like site in each (Table 2). To test the possibility that GL2 recognizes the five bHLH genes via the L1 boxlike sequence, we performed *Agrobacterium*-mediated transient expression analysis of GFP reporter genes containing the upstream regions of the five bHLH genes as promoters in tobacco leaves (Figure 3). GFP fluorescence levels were increased for all five of the upstream regions by co-introduction with the $35S$ promoter-driven VP16-GL2 N gene (Figure 3A to 3J and 3P). Changing the L1 box-like sequence to 5'-TACATCT-3' at all the sites in each upstream region reduced the increment in all instances (Figure 3F to 3P). These results suggest that GL2 recognized the five bHLH genes via the L1 box-like sequence.

GL2 Binds to Upstream Regions of the bHLH Transcription Factor Genes in planta

To examine the physical interaction of GL2 with the five bHLH genes *in planta*, we performed ChIP analysis using an anti-GFP antibody and a transgenic line harboring the *GL2* promoter-driven GFP-GL2 fusion gene *GL2pro-GFP-GL2* in the *gl2-5* mutant background (Figure 4A). To ensure that the fusion protein gene was functionally equivalent to the endogenous *GL2* gene, we confirmed that the transgene complemented *gl2-5* mutant phenotypes in the root and leaf epidermis (Figure 4B to 4E) and was expressed in the same manner as reported previously for *GL2* in the root epidermis (Figure 4F; Galway et al., 1994; Masucci et al., 1996).

A chromosome fraction was prepared from the roots of the transgenic line and fragmented DNA bound to GFP-GL2 was co-immunoprecipitated using the anti-GFP antibody. We examined the co-immunoprecipitates by quantitative real-time PCR to detect the short 100– 150-bp DNA regions containing the L1 box-like sequence upstream of the bHLH genes. The short DNA regions of all five bHLH genes and the *PLD*ζ*1* positive control gene were enriched significantly in the co-immunoprecipitates, whereas either a negative control region of *LRL1* or upstream regions of the negative control genes *ACT7* and *LRL3* were not (Figure 4G and Supplemental Figure S1). These results indicated that GL2 physically interacts with

the upstream regions of the five bHLH genes in roots, thereby confirming that they are GL2 target genes.

GL2 Suppresses the Expression of the bHLH Transcription Factor Genes in N Cells

To investigate how GL2 modulates the expressions of the five bHLH genes via binding to their upstream regions, we analyzed the promoter activity of each upstream region in wildtype and *gl2-5* mutant roots using the histochemical β-glucuronidase (GUS) reporter system. The upstream regions of *RHD6, RSL1, RSL2*, and *LRL2* exhibited promoter activity preferentially in the H cell files of the wild-type root epidermis (Figure 5A to 5C and 5E), but their activities extended to all cell files in the *gl2-5*mutant (Figure 5F to 5H and 5J) indicating that the promoter activities of these upstream regions were suppressed by *GL2* gene function in N cell files. The *LRL1* upstream region exhibited little promoter activity in the root epidermis based on the GUS reporter analysis (Figure 5D and 5I); however, a GFP reporter gene containing the *LRL1* upstream region and a genomic region encompassing the entire protein-coding sequence produced GFP florescence preferentially in H cell files in the wild type (Figure 5K). This preference disappeared in the *gl2-5*mutant (Figure 5L) indicating that *LRL1* is also suppressed by *GL2* in N cell files. Consistent with these results, qRT-PCR analysis showed that the transcript levels of the five bHLH genes were significantly higher in *gl2-5* mutant roots than in wild-type roots (Figure 5M). These results, together with the evidence for GL2 binding, indicate that the transcription of the five bHLH genes is suppressed directly by GL2 in N cell files.

The upstream regions of *RHD6* and *RSL1* exhibited promoter activities at earlier stages in root hair development than those of *RSL2* and *LRL2* (Figure 5A to 5C and 5E). The GFP reporter gene for *LRL1* was active in the root tip epidermis including the cell proliferation zone (Figure 5K). These results strongly suggest that the five bHLH genes are expressed differentially during the various stages of root hair development.

LRL1 and LRL2 redundantly function in root hair elongation downstream of GL2

RHD6 functions downstream of *GL2* in root hair initiation redundantly with *RSL1* (Masucci and Schiefelbein, 1994; Menand et al, 2007), and *RSL2* functions further downstream in root hair elongation (Yi et al, 2010). To examine the functional relationship of *LRL1* and *LRL2* with *GL2* in root hair development, the epistasis of their mutant effects was analyzed. Although the lines SALK_006430 (*Atlrl1-2* in Karas et al, 2009, and *bhlh66-1* in Bruex et al, 2012) and SALK_029317c (*bhlh69-1* in Bruex et al, 2012), which are designated as *lrl1-2* and *lrl2-2*, respectively, contain T-DNA insertions in introns (Supplemental Figure S2A), transcripts from both mutant genes were only scarcely detected by the RT-PCR analysis that produced intense signals for the wild-type genes (Supplemental Figure S2B). This indicates that the expressions of *LRL1*and *LRL2* are severely affected in these mutants.

Under the conditions used in this study, neither *lrl1-2* nor *lrl2-2* exhibited significant difference from the wild type in root hair length (Figure 6A to 6C and 6K). Whereas the homozygous double mutant plants could not be obtained, the partially homozygous double mutants *lrl1-2 lrl2-2/+* and *lrl-12/+ lrl2-2* exhibited short root hair phenotypes (Figure 6D, 6E, and 6K), as reported previously (Karas et al, 2009). Regarding root hair density, none of

the single or partially homozygous double mutants exhibited significant differences from the wild type (Figure 6L), as also reported previously (Karas et al, 2009). The *gl2-5* mutant exhibited significantly longer root hairs than the wild type (Figure 6F). However, root hairs of the *gl2-5 lrl1-2* and *gl2-5 lrl2-2* double mutants were significantly shorter than those of *gl2-5* and similar in length to those of the wild type (Figure 6F to 6H and 6K). Moreover, the partially homozygous triple mutants *gl2-5 lrl1-2 lrl2-2/+* and *gl2-5 lrl-12/+ lrl2-2* exhibited root hairs of similar lengths to those of *lrl1-2 lrl2-2/+* and *lrl-12/+ lrl2-2*, respectively (Fig 6D, 6E, and 6I to 6K). These results provide genetic evidence that *LRL1* and *LRL2* function redundantly as positive factors in root hair elongation downstream of *GL2*. In contrast, the *gl2-5* phenotype of higher root hair density due to ectopic root hair development in N-cell files was not affected by the *lrl* mutations (Figure 6L).

The bHLH Transcription Factors Have Diverse Functions in Cell Differentiation

To investigate the biological functions of the five bHLH transcription factors, we examined the effects of ectopic expression of their GFP-fusion proteins directed by the *GL2* promoter. Of transgenic plants that expressed the five fusion proteins at comparable levels (Supplemental Figure S3), those harboring *GL2pro-LRL1-GFP* and *GL2pro-LRL2-GFP* exhibited severe and moderate phenotypes, respectively, characterized by ectopic root hairs in N cell files and an increase in the number of root hairs (Figure 7F, 7G, 7J, and Supplemental Figure S4). In contrast, none of the other fusion genes exhibited significant effects on the root hair pattern (Figure 7C to 7E and 7J).

Because it was suggested that ectopic expression of a set of GL2 target genes caused severe phenotypes in cell differentiation including root hair-like structures produced from aerial epidermal cells (Ohashi et al., 2003), the cooperative effects of the fusion genes were examined in the F1 progeny of homozygous transgenic plants. Among the F1 plants, those that harbored both *GL2pro-RHD6-GFP* and *GL2pro-LRL1-GFP* exhibited a phenotype of ectopic and branching root hairs similar to the *GL2pro-VP16-GL2ΔN* plants (Figure 7H and 7I), while they did not develop aerial protuberant structures like those observed in the *GL2pro-VP16-GL2* N plants (Figure 7L). Their leaf epidermis exhibited various degrees of abnormalities in the initiation and branching pattern of trichomes. Some plants had no trichomes, similar to the *GL2pro-VP16-GL2* N plants (Figure 7M and 7N), and others had fewer trichomes with abnormal spatial patterns and divergent branch numbers (Supplemental Figure S5 and Supplemental Table S2). These results from single- and double-transgenic plants indicate that the five bHLH transcription factors have diverse functions in cell differentiation.

DISCUSSION

In this study, we showed that GL2 directly suppresses the five bHLH transcription factor genes *RHD6, RSL1, RSL2, LRL1*, and *LRL2* in N cell files. These genes have been identified as positive factors at various stages in root hair development. *RHD6* and *RSL1* promote root hair initiation in a partly redundant manner and *RSL2* subsequently promotes root hair elongation redundantly with *RSL4* (Menand et al., 2007; Yi et al., 2010; Pires et al., 2013). *LRL1* and *LRL2*, together with *LRL3*, are thought to function redundantly in root hair

elongation based on the root hair phenotypes of their single and multiple mutants (Karas et al., 2009; Bruex et al., 2012). This diversity in function is supported by the result that the upstream regions of *RHD6* and *RSL1* exhibited promoter activities at earlier stages in root hair development than those of *RSL2* and *LRL2*.

We confirmed that the partially homozygous double mutants of *LRL1* and *LRL2, lrl1-2 lrl2-2/+* and *lrl1-2/+ lrl2-2*, exhibited shorter root hairs than the wild type and found that their phenotypes in root hair lengths were epistatic to those of *gl2-5*. The possibility that they are also involved in processes other than root hair elongation cannot be excluded, however, because *lrl1 lrl2* double homozygous mutants could not be obtained. *LRL1* and *LRL2* are functional homologs of the *Lotus japonicas LRL1*, a mutant of which exhibited no indication of root hair initiation on almost all parts of the root surface (Karas et al., 2005). Moreover, the *GL2pro-LRL1-GFP* and *GL2pro-LRL2-GFP* transgenes caused ectopic root hairs in N cell files where *RHD6* and *RLS1* are normally inactive (Menand et al., 2007). These findings suggest that *LRL1* and *LRL2* are also involved in root hair initiation. This idea is supported, especially for *LRL1*, by the result that the GFP reporter gene for *LRL1* was expressed in root epidermal cells including those in the cell proliferation zone.

Different sets of transcription factor genes are known to be responsible for root hair development under various conditions. While *RHD6* is predominantly responsible for root hair initiation under normal growth conditions on agar media, *RSL1* contributes to an extent comparable to *RHD6* when roots are grown on cellophane disks (Menand et al., 2007). Moreover, in the presence of exogenous auxin, root hair initiation does not require either *RHD6* or *RSL1* (Yi et al., 2010) suggesting that other transcription factor genes are responsible for the root hair initiation. Despite such diversity of transcription factor genes for root hair initiation, the file-specific pattern of root hair formation is sustained robustly (Masucci and Schiefelbein, 1994). The mechanism ensuring this robustness might involve the targeting of multiple bHLH transcription factor genes by GL2. GL2 may also act as a negative factor for root hair elongation by suppressing *RSL2, LRL1*, and *LRL2*. Because *GL2* is expressed at a low level in H cells (Lee and Schiefelbein, 2002), GL2 might modulate root hair elongation in H cells via these bHLH genes. This idea is supported by the result that *gl2-5* exhibited significantly longer root hairs than the wild type. These findings suggest that for the fine and robust regulation of root hair development, GL2 suppresses multimodal pathways by targeting bHLH genes with diverse functions.

Transgenic plants harboring both the *GL2pro-RHD6-GFP* and *GL2pro-LRL1-GFP* transgenes exhibited abnormalities in root hair morphology and the initiation frequency and branching pattern of trichomes. Because these abnormalities were not observed in *GL2pro-RHD6-GFP* or *GL2pro-LRL1-GFP* homozygous transgenic plants, but were observed in the F1 progeny of crosses between them, the phenotype is thought to be due to a cooperative effect of the transgenes. The ectopic expression of GL2 target genes directed by VP16- GL2ΔN resulted in the development of root hair-like structures from aerial organs (Ohashi et al., 2003). In a related finding, ectopic coexpression of the *Physcomitrella patens* RHD6 like transcription factors PpRSL1 and PpRSL2 converted developing leafy shoot axes into rhizoids, which are protuberances similar in structure and function to root hairs (Jang et al., 2011). However, plants harboring both the *GL2pro-RHD6-GFP* and *GL2pro-LRL1-GFP*

genes did not develop aerial protuberant structures like those observed in the *GL2pro-VP16- GL2ΔN* plants while they exhibited root hair and trichome phenotypes similar to those of the *GL2pro-VP16-GL2 N* plants. GL2 target genes in addition to those encoding bHLH transcription factors may be required for such protuberant cell morphogenesis from aerial organs.

The GL2 DNA-binding domain has been shown to recognize DNA regions containing the L1 box-like sequence 5'-TAAATGT-3' in *in vitro* DNA foot-printing analysis (Ohashi et al., 2003) and yeast one-hybrid analysis (Tominaga-Wada et al., 2009; Khosla et al., 2014). The GL2 target genes identified in this study contain at least one copy of the L1 box-like sequence in each 3-kb upstream region and we confirmed the binding of GFP-GL2 in the close proximity of each L1 box-like sequence by ChIP analysis. To date, more than ten L1 box-like sites, including those from previous studies (Khosla et al., 2014; Wang et al., 2015), have been confirmed individually by ChIP analysis. Alignment of their sequences does not show strong conservation in the sequences surrounding the L1 box-like sites except for a relatively high A/T content (Supplemental Figure S6). Although this reflects a critical contribution of the L1 box-like sequence to the recognition by GL2, some imperfect L1 boxlike sequences might be recognized by GL2 because the mutated *RSL1* upstream region that lacked all of the L1 box-like sequence sites still functioned as a promoter activated by VP16-GL2 N (Figure 3L and 3P).

The gene regulatory networks for the cell fate determination in the root epidermis and for the morphological differentiation of root hair cells are the best-studied mechanisms of cell pattern formation and cell morphogenesis, respectively, in plants. This study integrates these two elements into a regulatory cascade via the direct interaction between GL2 and its target genes as illustrated in Figure 8. In N cells, a transcription factor complex composed of WER/MYB23, GL3/EGL3, and TTG1 activates *GL2* expression and GL2 negatively regulates the bHLH transcription factor genes *RHD6, RSL1, RSL2, LRL1*, and *LRL2* to suppress root hair development. In H cells, *GL2* is not activated due to the inhibitory function of CPC and its paralogs and the bHLH genes remain active to promote root hair development. In addition to the bHLH genes, GL2 may recognize a large number of target genes with various functions as shown in the list of candidate GL2 target genes in Supplemental Table S1. The enrichment for genes related to signal transduction (*i.e.*, those for lipid metabolic processes and hormone-mediated signaling pathways) and gene expression (*i.e.*, those for regulation of RNA metabolic processes and regulation of transcription) among the candidate genes is notable. Systematic approaches to identify GL2 target genes will help to elucidate the roles of *GL2* in the differentiation of various types of epidermal cells.

METHODS

Plant Materials and Growth Conditions

Arabidopsis thaliana, ecotype Columbia-0, was served as the wild type. The mutant line *gl2-5* (the Columbia genetic background; Ohashi et al., 2003) was gifted by G. Morelli (Food and Nutrition Research Centre, Agricultural Research Council, Rome, Italy). *lrl1-2* (SALK_006430) and *lrl2-2* (SALK_029317c) are ordered from ABRC (Alonso et al., 2003).

Sequences of the primers using in genotyping were presented in Supplemental Table S3. Plants were grown as described previously (Qin et al., 2005) unless otherwise noted.

Construction of Transgenes and Transformants

The GVG-inducible and *GL2* promoter-driven VP16-GL2 N genes were constructed previously (Ohashi et al., 2003). To construct the 35S promoter-driven GR-VP16-GL2 N gene, the promoter of the *GL2* promoter-driven VP16-GL2 N gene was replaced with the 35S promoter of pBI121 (Bevan et al., 1983), and a GR-coding fragment amplified by PCR from the GVG gene (Aoyama and Chua, 1997) was inserted at the 5'-end of the VP16 coding region in an in-frame manner. The *GL2* promoter-driven GFP-GL2 gene was constructed by assembling PCR products of the 2.1-kb *GL2* promoter fragment (Ohashi et al., 2003), and fragments encoding GFP and GL2 in the three-fragment recombination vector pK7m34GW (Invitrogen). To construct reporter genes for transient expression analysis, upstream fragments of bHLH genes were amplified by PCR from Arabidopsis genomic DNA, and assembled with the GFP-coding fragment in the vector pK7FWG0 (Invitrogen). To make mutant fragments, in which the L1 box-like sequence 5'- TAAATGT-3' was altered to 5'-TACATCT-3' at all the sites, the Q5 Site-Directed Mutagenesis Kit (New England Biolabs) was used. To construct the 35S promoter-driven VP16-GL2ΔN gene, the DNA region encoding VP16-GL2ΔN was cloned into pENTR-D-TOPO (Invitrogen) and transferred into the pK7WG2 expression vector (Invitrogen). To construct reporter genes for histochemical GUS analysis, upstream fragments of bHLH genes were amplified by PCR from *Arabidopsis* genomic DNA, and cloned into the vector pKGWFS7 (Invitrogen). To construct *GL2* promoter-driven bHLH-GFP fusion genes, bHLH-coding fragments were amplified by PCR from *Arabidopsis* cDNA, and assembled with the *GL2* promoter fragment and the GFP-coding fragment in the vector pK7m34GW. The DNA cloning into the vectors pK7m34GW, pK7FWG0, pK7WG2, and pKGWFS7was performed using Gateway system (Invitrogen) as described previously (Tao et al., 2013). Constructs were transformed into *Agrobacterium tumefaciens* GV3101/pMP90, using a freeze-thaw procedure. *Arabidopsis* transformation and transgenic plant screening were conducted as described previously (Qin et al., 2005). PCR primers used in DNA construction are shown in Supplemental Table S3.

Analyses for Transcript Levels

Total RNA was extracted using TRIzol reagent (Invitrogen) and treated with RNase-free DNase (TaKaRa). cDNA was synthesized from the total RNA using the Superscript II RT kit (Invitrogen). Microarray analysis was performed using GeneChip® ATH1 Arabidopsis (Affymetrix) according to Affymetrix protocols. qRT-PCR was performed as described previously (Wu et al., 2011) using SYBR Green Real-time PCR Master Mix (TOYOBO) on ABI7500 FAST (Applied Biosystems). Gene expression levels were standardized to the constitutive expression level of reference. At least three replications were performed for each sample in one experiment. The relative expression level of each gene was calculated based on the cycle threshold (CT) 2[−] ^{CT} method (Livak and Schmittgen, 2001). PCR primers used in qRT-PCR are shown in Supplemental Table S3.

Seedlings grown on agar medium (MS salts, 1% sucrose, 0.8% agar) around the four-leaf stage (about two weeks after germination) were transferred onto filter paper on agar medium (MS salts, 0% sucrose, 2% agar), and adapted to the open-air condition gradually for 3 days. Then, they were sprayed with induction or control solution, and kept for appropriate induction periods. For the microarray analysis, seedlings were treated with the induction solution containing 30 μM DEX and 0.01% Silwet L-77 for 5 hours. The induction solution for the DEX/CHX analysis contained 30 μM DEX, 30 μM CHX, and 0.01% Silwet L-77. For both analyses, the control solution was the same as the induction solution except for DEX.

Transient Expression Analysis in Tobacco Leaves

The Reporter and effector constructs were introduced into abaxial leaf epidermal cells of 2– 4-week-old *Nicotiana benthamiana* plants by *Agrobacterium* (GV3101/pMP90)-mediated transfection as described previously (Sparkes et al., 2006). Fluorescence images were captured using a confocal laser-scanning microscope TCS SPE (LEICA) 3 days after the transfection. The experiment was replicated three times for each effector-reporter combination. The intensity of GFP signal was quantified on the image with the assistance of ImageJ (<http://rsbweb.nih.gov/ij/>).

ChIP Analysis

ChIP analysis was performed using total root tissues of the wild-type and *GL2pro-GFP-GL2*/*gl2-5* plants as described previously (Gendrel et al., 2005). Immunoprecipitation was performed using Protein A beads (Millipore), anti-GFP antibody (Abcam), and IgG serum (Millipore). Co-immunoprecipitated DNA containing a particular region was quantified by real-time PCR with primers shown in Supplemental Table S3. The quantified value from the co-immunoprecipitated DNA was first normalized to that from the input DNA. The relative enrichment fold was then calculated by dividing the normalized value using the anti-GFP antibodywith that using the IgG sample.

GUS and GFP Reporter Analyses

The histochemical GUS reporter analysis was performed as described previously (Jefferson et al., 1987). Tissues were submerged in 90% acetone for 30 min at −20°C. After several washes with 0.1 M Na₂HPO₄ (pH 7.0), tissues were infiltrated under vacuum for 30 min, and then incubated at 37°C for 2–4 hours in the solution of 0.5 mg/mL 5-bromo-4-chloro-3 indolyl glucuronide, 0.1 M Na₂HPO₄ (pH 7.0), 10 mM Na₂EDTA, 0.5 mM potassium ferricyanide/ferrocyanide, and 0.06% Triton X-100. The staining solution was removed, and tissues were cleared in 70% ethanol. The observation and image capture of GFP fluorescence signal was performed using a confocal laser-scanning microscope TCS SPE (LEICA).

Observation and Evaluation of Root Hair and Trichome Phenotypes

To evaluate root hair phenotypes, plants were germinated and grown on the surface of vertically positioned agar medium containing MS salts, 1% sucrose, and 1% Phytagel™

(Sigma-Aldrich). Images of the primary root on the agar medium were captured with a CCD camera DFC425 C (LEICA) on stereomicroscope M205 FA (LEICA). The observation and image capture of root surface structures was performed using a microscope Axioplan 2 (Zeiss) equipped with a confocal laser-scanning unit CSU-X1 (Yokogawa) after staining with 5 μg/ml propidium iodide (PI). For root hair lengths, root hairs elongated horizontally to the agar surface in the region 5–7 mm from the root tip were measured on the image with the assistance of ImageJ. Immature root hairs or bulges with length less than 5 μm were omitted. For root hair numbers, both mature and immature root hairs including bulges in the visible side in the region 5–7 mm from the root tip were counted. Trichomes were observed by a variable-pressure scanning electron microscope S-3500 N (Hitachi) equipped with a cooling stage. The 10th–12th leaves were cut from the plants and directly observed invariable-pressure mode (100 Pa) at −20°C. To evaluate trichome and trichome branch numbers, 2-week-old seedlings germinated and grown on the surface of agar medium containing MS salts, 1% sucrose, and 1% Phytagel[™] (Sigma-Aldrich) were used.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Microarray and Individual Expression Analyses for Candidate GL2 Target Genes. **(A)** The structure of the modified GL2 gene (*GVGipro-VP16-GL2ΔN*) used in the expression analyses for candidate GL2 target genes is schematically illustrated. The promoter is glucocorticoid-inducible via the function of the GVG transcription factor. **(B)** The candidate target genes obtained from the microarray analysis were categorized by gene product functions as defined using the agriGO web-based software tool (Du et al., 2010), and the top five groups with most significant enrichment are shown with the ratios of

their populations in the candidates (white box) and the total *Arabidopsis* genes (gray box). *p*values are indicated on the right of boxes.

(C) Relative induction folds (DEX/Mock) of transcript levels in the expression analysis for individual bHLH VIIIc and XI subfamily genes, and the *PLD*ζ*1* gene as a positive control, are shown (mean \pm s.d., $n = 3$). Asterisks indicate significant differences between the levels of the DEX and Mock samples (* *P* < 0.05, ** *P* < 0.01, Student's *t*-test).

Lin et al. Page 18

Figure 2.

GL2 Direct Target Analysis of the Candidate GL2 Target Genes.

(A) The structure of the modified GL2 gene with a post-translationally inducible function (35Spro-GR-VP16-GL2 N) used in the GL2 directtarget analysis is schematically illustrated.

(B) Results of the inducible expression analysis using 35 *Spro-GR-VP16-GL2 N* are shown for the candidate GL2 target genes *RHD6, RSL1, RSL2, LRL1* and *LRL2*, and a negative control gene, *LRL3*. Relative induction folds (CHX DEX/CHX Mock) of transcript levels 1– 4 hours after induction are presented by different gray colors (mean ± s.d., *n* = 3). Asterisks indicate significant differences between the levels of the CHX-DEX and CHX-Mock samples (* *P* < 0.05, ** *P* < 0.01, Student's *t*-test).

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Figure 3.

Transient Expression Analysis for the Promoter Activity Directed by VP16-GL2 N. **(A)** to **(O)** Representative fluorescence images of the *N. benthamiana* leaf epidermis transfected with each bHLH promoter-driven GFP gene, *bHLHpro-GFP* **(A** to **E)**, each *bHLHpro-GFP* and the trans-factor gene *35Spro-VP16-GL2ΔN* **(F** to **J)**, or each mutant bHLH promoter-driven GFP gene, *bHLH-mL1pro-GFP*, and *35Spro-VP16-GL2ΔN* **(K** to **O)** are shown. The structures of the genes are schematically illustrated on the left of the images. The L1 box-like sequence 5'-TAAATGT-3' was altered to 5'-TACATCT-3' at all sites in each mutant bHLH promoter.

(P) Intensities of the GFP fluorescence signal were quantified and the relative intensity of the signal was calculated with the mean value for each *bHLHpro-GFP* arbitrarily set as 1 (mean \pm s.d., $n = 3$). Asterisks indicate that the values are significantly different from their corresponding *bHLHpro-GFP* values (* *P* < 0.05, ** *P* < 0.01, Student's *t*-test). Hashes indicate that the values are significantly different from their corresponding *bHLHpro-GFP*/ *35Spro-VP16-GL2ΔN* values (# *P*< 0.05; ## *P*< 0.01, Student's *t*-test).

Figure 4.

ChIP Analysis for the Binding of GL2 to the GL2 Target Genes. **(A)** The structure of the GL2 promoter-driven GFP-GL2 gene (*GL2pro-GFP-GL2*) used in the ChIP analysis is schematically illustrated. **(B)** to **(E)** Phenotypes in the root **(B** and **C)** and leaf **(D** and **E)** epidermis of the *gl2-5* mutant **(B** and **D)** and *GL2pro-GFP-GL2/gl2-5* transgenic **(C** and **E)** plants are shown. **(F)** The GFP fluorescence signal in a root tip region of the *GL2pro-GFP-GL2/gl2-5*

transgenic plant is shown.

(G) Results of the ChIP analysis are shown. The quantified DNA regions *PLD*ζ*1*-S, *RHD6*- S, *RSL1*-S1, *RSL1*-S2, *RSL2*-S1, *RSL2*-S2, *RSL2*-S3, *LRL1*-S1*, LRL1*-S2, and *LRL2*-S correspond to the sites of the L1 box-like sequence listed in Table II. The regions *RSL2*-S3 and *LRL2*-S1 contain two sites of the L1 box-like sequence for each. An upstream DNA region of *ACT7* was used as a negative control. Relative enrichment folds (anti-GFP/IgG) of *GL2pro-GFP-GL2*/*gl2-5* roots and the wild-type roots are shown in white and gray boxes, respectively (mean \pm s.d., $n = 3$). Asterisks indicate significant differences between the enrichment fols in *GL2-pro-GFP-GL2*/*gl2-5* and in the wild type (* *P*< 0.05, ** *P*< 0.01, Student's *t*-test). Bar = 100 μm **(B**, **C**, and **F)**, 500 μm **(D** and **E)**

Figure 5.

Expression Analyses of the GL2 Target Genes.

(A) to **(J)** Histochemical GUS expression patterns in 7-DAG root tips under the wild-type **(A** to **E)** and *gl2-5* **(F** to **J)** genetic backgrounds are shown for the upstream regions of *RHD6* **(A** and **F)**, *RSL1* **(B** and **G)**, *RSL2* **(C** and **H)**, *LRL1* **(D** and **I)**, and *LRL2* **(E** and **J)**. **(K)** and **(L)** The GFP fluorescence signal directed by the GFP reporter gene containing the upstream and total genomic regions of *LRL1* (*LRL1pro-GFP-LRL1g*) is shown. Root tips of the 7-DAG transgenic plants harboring *LRL1pro-GFP-LRL1g* under the wild-type **(K)** and

gl2-5 **(L)** genetic backgrounds were observed. Vertical bars indicate the root tip region containing proliferating cells.

(M) Relative transcript levels of *RHD6, RSL1, RSL2, LRL1*, and *LRL2* in 7-DAG wild-type (white box) and *gl2-5* (gray box) roots are shown. Transcript levels were determined by qRT-PCR, and their relative values were calculated by setting each wild-type mean value arbitrarily as 1 (mean \pm s.d., $n = 3$). Asterisks indicate significant differences between the transcript levels in $gl2-5$ and in the wild type (* $P < 0.05$, ** $P < 0.01$, Student's *t*-test). Bar = 100 μm

Figure 6.

Root Hair Phenotypes of *gl2-5, lrl1-2, lrl2-2*, and Their Multiple Mutants. **(A)** to **(J)** Main roots of the wild type **(A)**, *lrl1-2* **(B)**, *lrl2-2* **(C)**, *lrl1-2 lrl2-2/+* **(D)**, *lrl1-2/+ lrl2-2* **(E)**, *gl2-5* **(F)**, *gl2-5 lrl1-2* **(G)**, *gl2-5 lrl2-2* **(H)**, *gl2-5 lrl1-2 lrl2-2/+* **(I)**, and g*l2-5 lrl1-2/+ lrl2-2* **(J)** at 7-DAG are shown. Bar = 500 μm

(K) Root hair lengths of the wild type, *gl2-5, lrl1-2, lrl2-2*, and their multiple mutants at 7- DAG are shown. Root hairs in the region 5–7 mm apart from the root tip were measured (mean \pm s.d., $n = 400$). Asterisks indicate that the root hair lengths are significantly different from that of the wild type (* *P*< 0.05, ** *P*< 0.01, Student's *t*-test). Hashes indicate that the root hair lengths are significantly different from that of *gl2-5* (# *P*< 0.05, ## *P*< 0.01, Student's *t*-test).

(L) Root hair numbers of the wild type, *gl2-5, lrl1-2, lrl2-2*, and their multiple mutants at 7- DAG are shown. Root hairs and bulges in the visible side in the region 5–7 mm apart from the root tip were counted (mean \pm s.d., $n = 10$). Asterisks indicate that the root hair numbers are significantly different from that of the wild type (*P*< 0.01, Student's *t*-test). Hashes indicate that the root hair lengths are significantly different from that of $g/2-5$ ($P < 0.01$, Student's *t*-test).

Figure 7.

Phenotypes Caused by the *GL2* Promoter-driven bHLH-GFP Genes.

(A) to **(I)** Main roots of the wild-type **(A)** and *gl2-5* **(B)** plants, and the transgenic plants harboring *GL2pro-RHD6-GFP* **(C)**, *GL2pro-RSL1-GFP* **(D)**, *GL2pro-RSL2-GFP* **(E)**, *GL2pro-LRL1-GFP* **(F)**, *GL2pro-LRL2-GFP* **(G)**, *GL2pro-VP16-GL2* N **(H)**, and both *GL2pro-LRL1-GFP* and *GL2pro-RHD6-GFP* **(I)** at 7-DAG are shown. Typical branching root hair structures are indicated by arrows in **(H)** and **(I)**.

(J) Root hair numbers of the wild-type and *gl2-5* roots, and the transgenic roots harboring GL2pro-RHD6-GFP, GL2pro-RSL1-GFP, GL2pro-RSL2-GFP, GL2pro-LRL1-GFP, GL2pro-LRL2-GFP, GL2pro-VP16-GL2 N, and both *GL2pro-LRL1-GFP* and *GL2pro-*

RHD6-GFP at 7-DAG are shown (mean \pm s.d., $n = 10$). Asterisks indicate that the root hair numbers are significantly different from that of the wild type (*P*< 0.01, Student's *t*-test). **(K)** to **(N)** Phenotypes of the transgenic seedlings harboring *GL2pro-VP16-GL2 N* and both *GL2pro-LRL1-GFP* and *GL2pro-RHD6-GFP* in aerial organs. Seedlings of the wild type **(K)** and transgenic lines harboring *GL2pro-VP16-GL2ΔN* **(L** and **M)** and both *GL2pro-LRL1-GFP* and *GL2pro-RHD6-GFP* **(N)** are shown. The root hair-like structures in abaxial surface of the hypocotyl/cotyledon junction are indicated by an arrow in **(N)**. Bar = 250 μ m **(A** to **I)**, 1 cm **(K** to **N)**

Figure 8.

A Model of the Transcriptional Network Surrounding *GL2* in Root Epidermal Cells. In N cells (gray color), the WER/MYB23-GL3/EGL3-TTG1 complex activates *GL2.* GL2 directly suppresses the bHLH transcription factor genes *RHD6, RSL1, RSL2, LRL1*, and *LRL2*. In H cells (white color), WER and MYB23 in the complex are replaced by CPC and its paralogs. Consequently, the activation of *GL2* doesn't occur and the bHLH genes remain active to promote various stages of root hair development.

Table 1

Changes in Transcript Levels of *Arabidopsis* bHLH VIIIc and XI Subfamily Genes Caused by the Inducible Expression of VP16-GL2 N.

*** The value is the average fold-change of three independent experiments using GeneChip® ATH1 Arabidopsis microarray. False discovery rate (*q*value) is shown in parentheses.

****Data were not represented on the ATH1 microarray.

Table 2

Candidate Binding Sites of GL2 within the 3-kb Upstream DNA Regions of *PLD*ζ*1*, *RHD6*, *RSL1*, *RSL2*, *LRL1* and *LRL2*.

*** The sequence of the candidate GL2-binding site is presented. The L1 box-like sequence is indicated by boldface.

****Base-pair length from the initiation codon to the L1 box-like sequence is presented. c represents the complementary strand.