bHLH heterodimer complex variations regulate cell proliferation activity in the meristems of Arabidopsis thaliana

Control of cell proliferation

SAM
TMO5 LHW
TMO5 LL2

LM
TMO5 LHW
T5L1 LHW
T5L2 LHW

RAM
TMO5 LHW
T5L1 LHW
TMO5 LL1
TMO5 LL2

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Highlights
Expression of TMO5 and LHW bHLH clade members varies in distinct meristems
Single mutant analyses reveal functional specificity in meristems
Variations in TMO5/LHW heterodimer complexes affect target gene regulation
TMO5/LHW complexes are regulators of cell proliferation in all plant meristems
bHLH heterodimer complex variations regulate cell proliferation activity in the meristems of Arabidopsis thaliana

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SUMMARY

Root, shoot, and lateral meristems are the main regions of cell proliferation in plants. It has been proposed that meristems might have evolved dedicated transcriptional networks to balance cell proliferation. Here, we show that basic helix-loop-helix (bHLH) transcription factor heterodimers formed by members of the TARGET OF MONOPTEROS5 (TM05) and LONESOME HIGHWAY (LHW) subclades are general regulators of cell proliferation in all meristems. Yet, genetics and expression analyses suggest specific functions of these transcription factors in distinct meristems, possibly due to their expression domains determining heterodimer complex variations within meristems, and to a certain extent to the absence of some of them in a given meristem. Target gene specificity analysis for heterodimer complexes focusing on the LONELY GUY gene targets further suggests differences in transcriptional responses through heterodimer diversification that could allow a common bHLH heterodimer complex module to contribute to cell proliferation control in multiple meristems.

INTRODUCTION

Post-embryonic plant growth and development is driven by the activity of three main pools of pluripotent stem cells contained in zones called meristems. These are tightly regulated to divide and differentiate into specific cell types and form new organs. The root apical meristem (RAM) is located at the growing root tip, laid down during embryogenesis and responsible for the formation and primary growth of below-ground organs. At the other extremity, the activity of stem cells in the shoot apical meristem (SAM) is responsible for aerial organ development (Wang et al., 2018). While the apical meristems (RAM and SAM) give rise to the primary plant body, plants use a third pool of proliferating cells located in lateral meristems (LM) to support secondary growth leading to an increase in root and stem girth or thickness (Ragni and Greb, 2018). These meristems represent vascular and cork cambia (Etchells and Turner, 2010; Serra et al., 2022). Meristem activity is essential for growth and development and thus needs to be tightly controlled to ensure optimal growth depending on the environmental conditions and to avoid excessive cell proliferation (Motte et al., 2019).

Several key regulators, including transcription factors (TFs), and ligand-receptor complexes have been identified, which contribute to this intricate regulation of each of these meristem regions (Shimotohno and Scheres, 2019). For example, the CLAVATA3 (CLV3)-CLV1-WUSCHEL (WUS) negative feedback loop is the central genetic mechanism that coordinates stem cell proliferation with differentiation in the SAM. Perturbation of this regulatory network leads to phenotypes ranging from a loss of the meristem to a meristem-like structure (Shimotohno and Scheres, 2019). For example, the CLAVATA3 (CLV3)-CLV1-WUSCHEL (WUS) negative feedback loop is the central genetic mechanism that coordinates stem cell proliferation with differentiation in the SAM. Perturbation of this regulatory network leads to phenotypes ranging from a loss of the meristem to a meristem-like structure (Shimotohno and Scheres, 2019).
The DNA binding with One Finger (DOF)-type TFs have also been shown to control cell division orientation and proliferation in the vascular cells in the RAM in a cytokinin-dependent manner (Miyashima et al., 2019; Smet et al., 2019). These act downstream a complex formed by two phylogenetically distant basic helix-loop-helix (bHLH) TFs, TARGET OF MONOPTEROS (TMOS) and LONESOME HIGHWAY (LHW) (De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b, 2014).

So far, most of the studies have thus focused on factors that seem to be almost exclusively specific to one of the three meristem regions. While dedicated regulatory networks are likely required in different meristems, the alternative possibility remains that we are simply yet to uncover common factors required to regulating proliferation in all meristems. The TMOS/LHW bHLH heterodimer complex is a good candidate to function in multiple meristems as individual members have been shown to be broadly expressed in vascular tissues throughout plant development (De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b, 2014). Moreover, bHLH dimers are well suited to allow diversification in functions by using three main parameters: spatio-temporal expression patterns, DNA binding specificity, and dimerization properties (Grove et al., 2009). Indeed, bHLH TFs display a variety of expression patterns, where the overlap can define their sites of action in space and time (Qian et al., 2021; Hao et al., 2021). DNA binding specificity is dictated by a highly conserved signature of amino acid motif that forms the basic DNA-binding regions and shows significant variations in the bHLH family (Massari and Murre, 2000). Finally, specificity in dimerization properties was highlighted as a determining factor for the majority of bHLH proteins (Grove et al., 2009).

TMOS acts downstream the auxin-dependent transcription factor MONOPTEROS/AUXIN RESPONSE FACTORS (MP/ARFs) (Schlereth et al., 2010). TMOS has three homologs, TMOS-LIKE1-3 (TSL1-3), all showing a similar expression pattern restricted to the xylem in the RAM (De Rybel et al., 2013). Loss-of-function of TMOS and its closest homolog TSL1 sharing 48% protein similarity (De Rybel et al., 2013) leads to a reduced vascular cell number compared to wild type (WT) and a monarch patterning defect with only one pole of phloem and xylem, compared to the diarch WT phenotype (De Rybel et al., 2013). Higher order mutants enhance the severity of these phenotypes, suggesting they are redundant family members (De Rybel et al., 2013). Similarly, LHW also has three homologs, LHW-LIKE1-3 (LL1-3), none of which have significant homology with TMOS clade members (De Rybel et al., 2013). Although LHW and its homologs have a broader expression pattern in the RAM (Ohashi-Ito and Bergmann, 2007; De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b), defects in LHW lead to identical phenotypes as the tmo5 t5l1 double mutant (De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b). Moreover, higher order mutants increase the severity of the phenotypes, indicating that their function is dose dependent (Ohashi-Ito and Bergmann, 2007; De Rybel et al., 2013). Combined misexpression of both TMOS and LHW factors triggers ectopic periclinal and radial cell divisions throughout the RAM, suggesting that they function as part of an obligate heterodimer complex (Ohashi-Ito and Bergmann, 2007; De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b; Smet et al., 2019). Indeed, members of the TMOS and LHW subclades, which overlap in expression in the young xylem cells of the primary RAM, interact and form heterodimers (De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b, 2014). The TMOS/LHW complex directly activates expression of LONELY GUY3 (LOG3), LOG4, and BETA-GLUCOSIDASE44 in the xylem cells, leading to higher levels of active cytokinins by increasing biosynthesis (LOG3/4) and deconjugation (BGLU44) (De Rybel et al., 2014; Ohashi-Ito et al., 2014; Yang et al., 2021). Cytokinins are thought to diffuse to the neighboring procambium and phloem where they trigger cell divisions. This activity is balanced by CYTOKININ OXIDASE3, which is induced by SHORT ROOT, itself a direct TMOS/LHW target gene (Yang et al., 2021).

Here, we show that the role of TMOS/LHW complexes in regulating cell proliferation activity is not restricted only to the primary root meristem region but is more broadly required for normal development of RAM, SAM, and also of one of the LM, the root vascular cambium. Our results suggest a scenario where different TMOS/LHW complexes could regulate these differently organized meristems due to differences in expression domains and likely due to heterodimer complex variations between members of the TMOS and LHW subclades, leading to target gene specificity.

**RESULTS**

**TMOS/LHW function is not restricted to primary root development**

To establish if the function of TMOS and LHW clade members is restricted to the RAM or whether they play a broader role during plant development, we first explored the effect of altered heterodimer levels in the SAM and in the vascular cambium during root secondary growth, using a constitutive misexpression line...
that uses the well-characterized meristematic promoter RPS5A (Weijers et al., 2001; Figure S1A). The RPS5A promoter is expressed in dividing cells and particularly active in meristematic regions, thus allowing to target primarily the issues analyzed in this work. We also used existing higher order mutants (tmo5 t5l1 double, tmo5 t5l1 t5l3 triple, and lhw ll1 double mutants) (De Rybel et al., 2013; Ohashi-Ito et al., 2013a). Given that vascular cell numbers are not easily quantified in the SAM (vascular cell differentiation occurs at a significant distance from the SAM surface) and that the capacity of TMO5 and LHW to induce cell division is not limited to vascular cells (De Rybel et al., 2013), we measured the SAM area as a readout for a possible effect on cell proliferation activity. We found significant changes in the SAM area in all the lines compared to control wild-type Col-0 (Figures 1A–1E and 1K) (see example of SAM surface analysis in Figure S1B). The misexpression line, the tmo5 t5l1 t5l3, and lhw ll1 mutants had a smaller SAM while tmo5 t5l1 had a slightly but significantly bigger SAM area. Changes in SAM size were only partial due to changes in cell size (Figure 1L) and cell number (Figure S1C), indeed suggesting a role of these genes in the regulation of cell proliferation throughout the SAM which is more complex compared to the RAM. Similar to the effects observed in the primary RAM, the number of vascular cell files was reduced during root secondary growth in a dose-dependent manner in the loss-of-function mutant lines and increased in the misexpression line (Figures 1F–1J and 1M). In summary, these results
suggest that the activity of TMO5/LHW and some of their homologs might not be restricted only to control cell proliferation in the primary RAM but in other meristems as well.

**TMO5/LHW is required and sufficient for root secondary growth**

Our results suggest that the role of the TMO5/LHW pathway in the regulation of cell proliferation could be conserved in primary meristems and also during root secondary growth. However, during secondary growth, it remains possible that the observed effects are a consequence of the persistent lack or overexpression of these factors during primary growth. Thus, to investigate if TMO5 is sufficient to trigger vascular proliferation during secondary growth specifically, TMO5 was exclusively expressed during this developmental stage by introducing a dexamethasone (DEX)-inducible ProRPS5A::TMO5-GR rescue construct (using the RPS5A promoter to target expression in meristematic regions) into the tmo5 t5l1 t5l3 triple mutant.

When grown on medium supplemented with 10 μM DEX, the ProRPS5A::TMO5-GR construct introduced in the triple mutant can rescue the proliferation defect in the tmo5 t5l1 t5l3 triple mutant to an almost non-phenotypical (t5l1 t5l3 double mutant) situation (Figures 2A, 2B, and 2I) (De Rybel et al., 2013). This inducible rescue system was next used as a tool to investigate if TMO5 expression during root secondary growth is sufficient to trigger vascular cell proliferation. tmo5 t5l1 t5l3 mutants with and without the inducible ProRPS5A::TMO5-GR rescue construct were grown on mock medium for 5 days and then transferred onto inducing medium supplemented with 10 μM DEX for another 5 days. Again, the number of vascular cell files in the root undergoing secondary growth was quantified. tmo5 t5l1 t5l3 mutants carrying the ProRPS5A::TMO5-GR rescue construct showed a significant increase in vascular cells numbers compared with the triple mutant without the rescue construct (Figures 2C, 2D, and 2I). To further confirm that the observed difference was specifically due to induction during secondary growth, this experiment was repeated by applying local DEX treatment only to the investigated root area. This was achieved by growing the seedlings on mock medium for 5 days and then transferring the plants for 5 additional days to a plate where mock and DEX media were physically separated (Figure S2). Quantification of the number of vascular cell files confirms that induction of TMO5 specifically during secondary growth is sufficient to trigger...
periclinal division leading to radial expansion. Next, we made use of the same inducible rescue system to investigate if the TMO5/LHW pathway is required for secondary growth. Seedlings were first grown on inducible DEX medium for 5 days and then transferred onto mock medium for an additional 10 days. This timing was chosen as the effect of the initial 5 days DEX treatment persists for several days after transfer to mock medium (Figures 2E, 2F, and 2I). After the transfer to mock medium and an additional 10 days of growth, no significant difference could be observed between the number of vascular cell files of the triple mutant compared to the triple mutant carrying the ProRPS5A::TMO5-GR rescue construct (Figures 2G–2I), indicating that TMO5 presence during primary growth is sufficient to initiate secondary growth but not to maintain it. These results therefore suggest that the TMO5/LHW pathway is both required and sufficient to allow vascular proliferation during root secondary growth.

TMO5 and LHW clade members show overlapping expression in distinct meristems

Although the redundant role of both TMO5 and LHW subclade members in vascular proliferation of the primary RAM has been described in detail, some of the observed phenotypes associated with higher order mutants of these factors are not restricted to the primary root meristem (De Rybel et al., 2013; Ohashi-Ito et al., 2013a, 2013b). Moreover, as mentioned previously, there are some indications that TMO5 and LHW bHLH subclade members are expressed in aerial tissues as well (Ohashi-Ito et al., 2013a), consistently with the effect on SAM size observed in mutants and in the TMO5/LHW misexpression line (Figures 1A–1E and 1K).

In order to first provide a detailed overview of the localization of these factors in Arabidopsis thaliana, we generated promoter-nuclear triple GFP-GUS fusion constructs for all members and used confocal microscopy to analyze their expression domains in the RAM, in the vascular cambium during root secondary growth, and in the SAM. As previously reported (De Rybel et al., 2013), TMO5 clade members show overlapping expression in the young xylem cells of the RAM (Figures 3A–3D). Similarly, during secondary growth in the root, TMO5 clade members showed expression in developing and differentiating xylem cells. TSL1 and TSL3 were also detected in some cells of the vascular cambium (Figures 3E–3H). In the SAM region, only TMO5 showed a specific provasculature-associated expression (Figure 3I). TSL1 was not detected in the SAM (Figure 3J) but did show expression in a few cell files in the vascular tissue below the
SAM (Figure S3A). T5L2 was found to be highly expressed in the L1 layer and at much lower levels in other cells of the SAM (Figure 3K). T5L2 was also largely excluded from the center of the meristem. T5L3 was expressed broadly in the SAM both in the L1 layer and in the internal tissues, except for the central part of the SAM where it was completely absent (Figure 3L).

Compared to TMO5 clade members in the RAM, members of the LHW clade showed a broader expression domain in vascular tissues (Figures 4A–4D), and in the case of LHW and LL1, this pattern was similar to previously published lines (Figures 4A and 4B) (De Rybel et al., 2013). Also, during secondary growth, LHW, LL1, and LL3 clade members showed expression in xylem and cambium tissues, while LL2 was only detected in the xylem (Figures 4E–4H). A broad expression domain was observed in the SAM for both LHW and LL1 (Figures 4I and 4J), but LHW was absent specifically from the L2 layer while LL1 was more specifically expressed in organ primordia in the peripheral zone. No expression was detected in the SAM for LL2 and LL3 (Figure 4K and 4L) and, similarly to T5L1, LL3 showed expression within the vasculature below the SAM (Figure S3B). Taken together, our results show that the TMO5 and LHW clade members are expressed in distinct meristematic regions, consistent with a general meristematic function for these factors throughout development that, at least in the SAM, is not restricted only to the regulation of vascular tissue development. Additionally, some of these TFs show prominent expression only in one of the meristems, while being absent in others. These results thus argue for a general function of the TMO5-T5Ls/LHW-LLs factors in all the meristems, but with some expression specificity that could result in alternative TMO5-T5Ls/LHW-LLs combinations in the different meristems.

**Single mutant analysis reveals functional specificity in TMO5 and LHW clade members**

Despite the fact that TMO5 and LHW homologs have a clear redundant role in primary root vascular proliferation, the observed diversity in expression patterns in other meristems suggests that there might be some functional specificity among the clade members. Indeed, even in the primary root meristem, TMO5, T5L1, and LHW are reported to be the most prominent factors driving vascular proliferation, while the other members might be less important for this specific developmental process (De Rybel et al., 2013; Ohashi-Ito et al., 2014). To get a global understanding of the functional specificity among the TMO5 and LHW clade members throughout plant development, we next analyzed single mutants for discernible
phenotypes in the three meristems in comparison to WT plants. In the RAM, all single mutants with the exception of t5l2 showed a significant reduction in cell files number (Figures 5A and S4A–S4I). Still, a clear distinction within the subclades was observed in the relative contributions to this phenotype, with tmo5, t5l1, t5l3, and lhw showing the strongest reduction in the number of vascular cell files. It is worth noting that lhw is the only single mutant with a monarch instead of the normal diarch vascular architecture (Ohashi-Ito and Bergmann, 2007), explaining the more pronounced phenotype. Similar observations were made in roots initiating secondary growth, with tmo5, t5l1, t5l2, lhw, and l1l-analyzed mutants showing a significant reduction in cell file numbers, but the relative contributions of the factors were different. Indeed, t5l1, t5l2, and lhw seem to be the major players in the establishment of secondary growth (Figures 5B and S4J–S4O). Additionally, the SAM area was significantly larger in t5l1 and l1l mutants as well as the number of meristem cells (Figures 5C and S4P–S4Y). Cell size was not significantly affected in single mutants, confirming an effect on cell proliferation in the SAM (Figure 5D). Additional phenotypes were found in the inflorescence of mutants. For example, tmo5 and t5l1 started initiating siliques before the last inflorescence branch (Figure S5A) suggesting a deviation in lateral organ/structure identity determination in the SAM, whereas t5l3 produced inflorescence branches incapable of upright growth (Figure S5B). Finally, inflorescence growth of the lhw mutant was slower, and much more affected in l1l double mutants (Figure S5A). In summary, these results show that, while a general effect on cell proliferation is observed, the effect of single mutations in the members of TMO5 and LHW clades differs depending on the meristem considered, suggesting a level of functional and tissue specificity with an opposite trend in root and shoot.

Figure 5. Single mutant analysis reveals functional specificity in TMO5 and LHW clade members
Quantification of Col-0, tmo5, t5l1, t5l2, t5l3, lhw, l1l, and l1l for vascular cell files number of cross sections (A) of root apical meristems, (B) of root undergoing secondary growth, (C) measurement of shoot apical meristem area, and (D) cell size in L1 layer. Boxplots show the median and interquartile range (25%–75%), and whiskers represent the 1.5 × interquartile range. Lowercase letters in charts indicate significantly different groups as determined using a one-way ANOVA with post-hoc Tukey HSD testing (p < 0.05).
Variations in bHLH heterodimer complexes show distinct phenotypic outputs

Although the observed tissue- and organ-specific expression of the TMO5 and LHW clade members could account for the phenotypic differences in single mutants, there was no perfect correlation, suggesting that there might be other layers of functional regulation. Given that TMO5 and LHW clade members form obligate heterodimer complexes (De Rybel et al., 2013; Ohashi-Ito et al., 2014), we next questioned if such functional specificity could be caused by the particular heterodimer complex that is formed. Although TMO5 and LHW interaction partners can likely all interact with each other (De Rybel et al., 2013; Ohashi-Ito et al., 2014), this indeed does not mean that these combinations would lead to a functional bHLH complex. We thus combined several individual constitutive misexpression lines (using the RPS5A promoter to drive expression in meristems) by crossing lines misexpressing TMO5 clade members with lines misexpressing LHW clade members and analyzing the effect in the RAM and the SAM (Figure 6). Compared to the wild type, different combinations resulted in a quantitative difference in the number of root vascular cell files with TMO5/LHW as the most potent combination and T5L2/LHW not showing any significant difference (Figures 6A–6E and 6K). Strong phenotypical effects were observed in the shoot as previously reported for TMO5/LHW (Vera-Sirera et al., 2015), including reduced stem height, curly, hyponastic, or jagged leaves (Figure S6). In the SAM, our analysis suggests that the resulting phenotypes and their severity show a tendency to be dependent on the combination of TMO5 and LHW clade members as in the root but the high variability of the parameters allows firm conclusions only for TMO5/L2 with a reduced SAM area (Figures 6F–6J and 6L) and SAM cell number (Figure S7), and TMO5/LHW with a reduced cell size (Figures 1 Ma n d 6M).

In summary, these experiments show that even when TMO5 and LHW clade members are ectopically expressed together, they do not always lead to the same phenotype. Thus, TMO5- and LHW-clade
heterodimer complex activity is not solely determined via the observed differential expression domains but also likely because of variations in the complexes or in the activity of the complexes which are being formed.

Variations in bHLH heterodimer complexes affect target gene specificity

TMO5/LHW complexes induce vascular cell proliferation in the root apical meristem via induction of cytokinins biosynthesis through direct binding to the promoter regions of LOG3 and LOG4 (De Rybel et al., 2014; Ohashi-Ito et al., 2014; Smet et al., 2019). These enzymes catalyze the final step of cytokinin biosynthesis (Kurakawa et al., 2007; Kuroha et al., 2009). We therefore set to explore quantitatively and functionally the differences in gene regulatory potential between the different heterodimers that can be formed by analyzing their potential to activate not only LOG3 and LOG4 but also several other LOG genes. To achieve this, we implemented a protoplast-based quantitative gene expression system that enables covering all possible combinations, obtaining quantitative data, and reducing interferences from other factors and tissue context. Protoplasts derived from Arabidopsis shoot tissues (see STAR Methods for details) were transiently transformed with constructs comprising each combination of the TFs under the control of a 35S constitutive promoter. The promoters of the tested genes, namely LOG1, LOG3, LOG4, LOG5, and LOG7, were cloned upstream the firefly luciferase gene, that served as a readout. A construct coding for constitutively expressed Renilla luciferase was included as a normalization element. We first assayed the effect of overexpression of single TFs on the expression of each of the LOG genes (Figure S8 A and Table S1). LHW and LL1 were able to induce LOG3 and LOG4 expression only, although to moderate levels, whereas all other TFs had no inducive effects on any LOG expression. These results are overall consistent with the idea that TMO5 and LHW clade members act as obligate heterodimers (De Rybel et al., 2013; Ohashi-Ito et al., 2014). It also suggests that LHW and LL1 can activate a basal level of transcription by themselves. Overexpression of any combination of two transcription factors from the TMO5 and LHW clades was able to induce expression of the direct target genes LOG3 and LOG4, but there was a clear quantitative difference with the highest induction values for all TMO5-clade combinations with LL1, and the lowest in combinations with LL2 (Figure 7 and Table S1). None of the other LOG genes analyzed were induced by any of the TF complexes (Figure 7 and Table S1), suggesting a specific regulation of LOG3 and LOG4 by the TMO5 and LHW clade members or that other regulatory factors not present in protoplasts might be needed for their induction. Taken together, these experiments show that LHW clade members play a key role in defining the strength of the transcriptional activation of LOG3 and LOG4 in the simplified protoplast system, and not the TMO5 clade.

To further validate these results in planta, we inspected the ProLOG3::n3GFP and ProLOG4::n3GFP reporter lines in lhw and tmo5 t5l1 mutant backgrounds (Figures S8B–S8G). In the wild-type background, ProLOG3::n3GFP was expressed in a diarch configuration in flower primordia in the SAM (Figure S8B; white arrows), similarly to the diarch expression in root protoxylem cells (De Rybel et al., 2014). The signal of

| ProLOG1 | 2.30 | 5.70 | 0.64 | 1.12 | 1.76 | 6.20 | 0.89 | 0.96 | 1.00 | 2.60 | 0.79 | 1.18 | 1.44 | 6.48 | 0.80 | 1.20 |
| ProLOG3 | 72.00 | 265.00 | 35.00 | 44.90 | 43.00 | 413.00 | 29.00 | 32.00 | 33.00 | 296.00 | 25.00 | 46.00 | 76.00 | 493.00 | 35.00 | 62.00 |
| ProLOG4 | 233.00 | 576.00 | 87.00 | 208.00 | 144.00 | 534.00 | 65.00 | 100.00 | 207.00 | 718.00 | 52.00 | 134.00 | 149.00 | 586.00 | 75.00 | 88.00 |
| ProLOG5 | 0.11 | 0.16 | 0.33 | 0.21 | 0.18 | 0.11 | 0.32 | 0.37 | 0.39 | 0.25 | 0.49 | 0.48 | 0.23 | 0.20 | 0.37 | 0.33 |
| ProLOG7 | 0.35 | 1.06 | 0.21 | 0.24 | 0.46 | 0.82 | 0.30 | 0.36 | 0.48 | 1.09 | 0.52 | 0.51 | 0.42 | 1.88 | 0.39 | 0.45 |

**Figure 7. Different combinations in bHLH heterodimer complexes affect target gene expression**

Heatmap shows relative changes of gene expression from LOG promoters after overexpression of combinations of TMO5 and LHW clade members’ pairs in quantitative gene expression assays in Arabidopsis protoplasts. Values represent a ratio between promoter expression with transcription factors and basal promoter expression without adding transcription factors.
**ProLOG3::n3GFP** was still present in the **tmo5 t5l1** double mutant (Figure S8C) despite a loss of activity of two TMO5 clade members affecting SAM phenotype (Figures 1C, 1L, and S4). However, the expression of ProLOG3::n3GFP in **tmo5 t5l1** was detected only in one cell axis compared to the control in an inspected SAM area (Figure S8C; yellow arrows) suggesting impaired vascular formation similar to the effects in the RAM (Ohashi-Ito and Bergmann, 2007; De Rybel et al., 2013). The expression of ProLOG4::n3GFP in the L1 layer (Figure S8E) decreased in a **tmo5 t5l1** double mutant but the overall pattern remained similar (Figure S8F). In contrast, the signal of both ProLOG3::n3GFP and ProLOG4::n3GFP was completely missing in a **lhw** single mutant (Figures S8D and S8G). Overall, these findings support the results of the protoplast assays showing that LHW activates LOG3 and LOG4 transcription together with TMO5 clade members, and that the LHW clade members are essential for LOG3 and LOG4 expression.

**DISCUSSION**

Current knowledge on the regulatory complexes governing cell proliferation suggests that plants have evolved specific networks for each of the meristem regions. Although most known examples are comparable since they are based on a peptide-receptor interaction pair, these are unique for one specific meristem context: CLV3-CLV1 in the SAM (Gaillochet et al., 2015), CLE41-PXY in the vascular cambium (Fisher and Turner, 2007; Suer et al., 2011; Etchells et al., 2013), and CLE40-ACR4 in the RAM (Stahl et al., 2013; Berckmans et al., 2020). One can however question if this is accurately reflecting an evolutionary reality where each meristem region has independently developed a regulated regulatory network, or whether we are simply yet to uncover common regulators in these distinct regions governing cell proliferation. Our results suggest that the TMO5/LHW and T5Ls/LLs bHLH heterodimer complexes could act as general regulators of cell proliferation expressed and active in different plant meristems, besides the previously described role in the RAM. Although all homologs show overlapping expression domains in the RAM, there is more variation in expression domains in the SAM and vascular cambium areas (Figures 3 and 4). Our work thus highlights the fact that overlapping expression patterns of TF regulating together development in a given meristem are not necessarily copied to other meristems, further complicating extrapolation of functional studies performed in one organ to another. The obligate heterodimer nature of this interaction is likely to be of key importance to explain how different combinations of TMO5 and LHW subclade members could regulate proliferation in different meristems. Indeed, homodimers present in e.g. single misexpression lines (De Rybel et al., 2013) do not give a strong phenotypical effect as when both partners are overexpressed. Moreover, heterodimers are required for an efficient regulation of the well-characterized target genes LOG3 and LOG4 in both protoplasts and in planta (Figures 7 and S8), although our results in protoplasts suggest that LHW and LL1 alone have a basal capacity to activate target genes that could be functionally significant. Additionally, higher order mutants of each subclade, such as **tmo5 t5l1 t5l2** and **lhw ll1** mutants, yield the same phenotype (De Rybel et al., 2013). Our results and previously published work thus suggest that heterodimer variations by combinations of TMO5 and LHW subclade members resulting primarily from expression in specific domains within different meristems (and only secondarily from differential expression between meristems, as e.g. for TSL1, LL2, and LL3 that are absent in the SAM) could provide the required specificity to adapt responses to a given developmental context, such as different meristems. The activity of TMO5/LHW heterodimers would then be determined by specificity of the promoter regions of TMO5/LHW genes, first restricting expression of TMO5/LHW subclade genes to specific regions in the different meristems. Identifying the factors that controls this specificity in expression will thus be required to understand how TMO5/LHW heterodimers fulfill their functions in different meristems. Expression of **TSL2**, **TSL3**, **LHW**, and **LL1** genes in the SAM proper, and not only in vascular tissues as in other meristems, exemplifies how divergent expression patterns could possibly lead to the formation of heterodimers in specific domains of meristems to fulfill their function in the regulation of cell proliferation in the different meristems. Note however that the phenotype we observed in the mutants was not always correlated to the expression patterns, notably in the SAM, which could be a compensation mechanism to cope with the loss of activity of a given gene by other members of the same subclade. Further work will thus be required to fully understand how expression patterns contribute to the function of a given heterodimer in different meristems.

Although our results suggest that TMO5/LHW heterodimers could act as a common regulator controlling cell proliferation, the exact downstream mechanism might not be conserved in each of the meristem contexts. LOG3 and LOG4 genes are induced by all tested heterodimer complexes in a protoplast system and complexes containing LHW, and LL1 is required for their expression both in the RAM and SAM (De Rybel et al., 2014) (Figure S8). This suggests that regulation of cytokinin biosynthesis by heterodimer
combinations between TMOS and LHW clade members could be a conserved mechanism in different meristems, even outside of vascular tissues as in the SAM. However, opposite responses on cell proliferation are observed upon misexpression in the SAM and RAM regions. Our results using protoplast indicates that this could be due to quantitative differences between heterodimers in their capacity to activate common target genes such as LOG3 and LOG4. However, this is also most likely be due to the differences in DNA binding specificity of the different heterodimers in each developmental context or participation of additional regulatory factors. It is indeed very likely that a different set of target genes will be activated by specific heterodimer complexes leading to further functional diversification. Some indications can be found in literature as TSL1/LHW misexpression was shown to have only partly overlapping target genes compared to TMOS/LHW misexpression (De Rybel et al., 2014; Ohashi-Ito et al., 2014). However, these gene lists have been obtained using a very different experimental set-up, precluding a direct comparison. As such, additional work using a genome-wide analysis of target genes for the different heterodimer complexes would be required to evaluate the precise downstream target gene sets activated by these complexes. In summary, our work suggests a scenario where a common bHLH heterodimer complex module controls cell proliferation in distinct plant meristems in Arabidopsis thaliana through heterodimer diversification leading to either quantitative differences in common target genes or to target gene specification.

An intriguing question emerging from our results is whether the bHLH heterodimer complexes are unique in their capacity to act as more general regulators that can be used in different developmental contexts; or whether this is a more general theme for most TFs which has simply not been uncovered so far. On one hand, one could argue for bHLH factors being unique as there are other examples of a same set of bHLH factors acting in multiple contexts. For example, the formation of trichomes and root hairs respectively depends on the bHLH proteins GL3 or EGL3, which interact with the MYB proteins WER or GL1 thus forming active transcriptional complexes used in these two developmental contexts (Bernhardt et al., 2003; Zhang et al., 2003; Zhao et al., 2008). On the other hand, as a general property, several TF families can form within-family heterodimers and TFs from a given family can interact with many other TFs from other families (Trigg et al., 2017). Thus, variations of heterodimer formation of TFs from other families than bHLH could occur in different meristems through differential expression and could be regulating meristems similarly to the TMOS/LHW heterodimer. Although it might thus be evolutionary efficient for organisms to use TFs dedicated to a given tissue, combinatorial TF interactions among different families are an alternative to achieve the same level of specificity needed in each developmental context through regulation of expression patterns within and not between tissues. It is thus likely that we are yet to uncover additional functions of known TFs in other developmental contexts which could emerge by interactions with other partners.

Limitations of the study

Our work provides data on the transcriptional domains of TMOS and LHW subclades members in Arabidopsis meristems but not of the corresponding proteins, which would be required to fully identify where heterodimers can form. Further studies are also needed to directly visualize the formation of various heterodimers in different meristems. In addition, we analyzed the regulation of target gene expression by these bHLH complexes using only some members of the LOG gene family. A wider gene target spectrum will be needed to understand the impact of heterodimer variations on transcriptional responses in a broader way. Finally, functional redundancy likely explains why the observed phenotypes are relatively weak in single mutants used in this study. Thus, further genetic analysis is needed to resolve these questions.

STAR+METHODS

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Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.105364.

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AUTHOR CONTRIBUTIONS
E.M., M.P., B.D.R., and T.V. designed the research. E.M. performed the experiments related to RAM and vascular cambia with the help of M.M. and J.N. L.R. and D.R. analyzed root secondary growth expression pattern. M.P. designed and performed the experiments related to SAM, shoot phenotype, and LUC assays with the help of G.C., C.G., J.A., and M.D.Z. E.M. and M.P. prepared figures and manuscript draft. E.M., M.P., B.D.R., and T.V. wrote the manuscript with input from all authors.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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# STAR METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| 10-beta Competent E. coli | New England Biolabs | C3019H |
| One Shot™ TOP10Chemically Competent E. coli (Invitrogen) | Thermo Fisher Scientific | C404006 |
| Agrobacterium tumefaciens strain CS8C1<sup>1</sup> (pMP90) LBA4404 | N/A | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| Dexamethasone | Merck | D4902 |
| MES | Duchefa | M1503.0100 |
| M5 | Duchefa | M0221.0050 |
| Plant Agar | Neogen | NCM0250A |
| Propidium Iodide | Merck | P4170 |
| Propidium iodide solution | Merck | P4864 |
| Kanamycin | Duchefa | CAT 25389-94-0 |
| Hygromycin | Duchefa | CAT 31282-04-9 |
| Sulfadiazine | Sigma-Aldrich | CAT 68-35-9 |
| D-luciferin | Biosynth AG | FL08608 |
| Q5<sup>®</sup> High-Fidelity DNA Polymerase | New England Biolabs | M0491L |
| **Critical commercial assays** | | |
| Technovit 7100 Kit | Heraeus Kulzer | CAT 64709003 |
| NucleoSpin RNA Plant Kit | Macherey-Nagel | 740949.50 |
| NucleoSpin Plasmid, Mini kit for plasmid DNA | Macherey-Nagel | 740588.50 |
| NucleoBond Xtra Midi kit for transfection-grade plasmid DNA | Macherey-Nagel | 740410.50 |
| Wizard® Plus Midipreps DNA Purification System | Promega | A7640 |
| SuperScript™ III Reverse Transcriptase (Invitrogen) | Thermo Fisher Scientific | 18080044 |
| NucleoSpin Gel and PCR Clean-up | Macherey-Nagel | 740609.50 |
| **Experimental models: Organisms/strains** | | |
| Arabidopsis: Col-0 | NASC | N1093 |
| Arabidopsis: tmo5 | De Rybel et al., 2013 | GK-143E03 |
| Arabidopsis: tmo5 like 1 | De Rybel et al., 2013 | RIKEN_12-4602-1 |
| Arabidopsis: tmo5 like 2 | De Rybel et al., 2013 | GK-824H07 |
| Arabidopsis: tmo5 like 3 | De Rybel et al., 2013 | SALK_109295 |
| Arabidopsis: liw | De Rybel et al., 2013 | SALK_023629 |
| Arabidopsis: liw like1 | Ohashi-Ito et al., 2013a | SALK_126132 |
| Arabidopsis: liw like2 | NASC | GK-523B12 |
| Arabidopsis: liw like3 | NASC | GK-262H03 |
| Arabidopsis: tmo5 ts11 | De Rybel et al., 2013 | N/A |
| Arabidopsis: tmo5 ts11 ts13 | De Rybel et al., 2013 | N/A |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Teva Vernoux (teva.vernoux@ens-lyon.fr).

Materials availability
Plasmids and plant lines generated in this study are available from the lead contact upon request.

Data and code availability
- Microscopy data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Arabidopsis: lhw II1 | Ohashi-Ito et al., 2013a | N/A |
| Arabidopsis: tmo5 t5l1 t5l3 x ProRPSSA::TMO5-GR | De Rybel et al., 2013 | N/A |
| ProRPSSA::TMO5-GR x ProRPSSA::LHW-GR | Smet et al., 2019 | N/A |
| ProRPSSA::TMO5 x ProRPSSA::LHW-LIKE1 | This Study | N/A |
| ProRPSSA::TMO5 x ProRPSSA::LHW-LIKE2 | This Study | N/A |
| ProRPSSA::LHW x ProRPSSA::TMO5-LIKE1 | This Study | N/A |
| ProRPSSA::LHW x ProRPSSA::TMO5-LIKE2 | This Study | N/A |
| ProRPSSA::3nGFP-GUS | This Study | N/A |
| ProTMO5:3nGFP-GUS | This Study | N/A |
| ProTMO5-LIKE1::3nGFP-GUS | This Study | N/A |
| ProTMO5-LIKE2::3nGFP-GUS | This Study | N/A |
| ProTMO5-LIKE3::3nGFP-GUS | This Study | N/A |
| ProLHW::3nGFP-GUS | This Study | N/A |
| ProLHW-LIKE1::3nGFP-GUS | This Study | N/A |
| ProLHW-LIKE2::3nGFP-GUS | This Study | N/A |
| ProLHW-LIKE3::3nGFP-GUS | This Study | N/A |
| ProLOG3::n3GFP | De Rybel et al., 2014 | N/A |
| ProLOG4::n3GFP | De Rybel et al., 2014 | N/A |

Oligonucleotides
Provided in Table S2 | N/A | N/A |

Recombinant DNA
Provided in Table S3 | N/A | N/A |

Software and algorithms
PlotsOfData webtool | UVA | https://huygens.science.uva.nl/PlotsOfData/ |
Fiji | NIH | https://fiji.sc/ |
ANOVA | Astatsa | https://astatsa.com/OneWay_Anova_with_TukeyHSD |
Auto seeded 3D watershed algorithm (MARS) | Fernandez et al. (2010) | https://gitlab.inria.fr/mosaic/publications/sam_layer_height |
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Arabidopsis thaliana

Unless otherwise mentioned, all plant material used was Arabidopsis thaliana, ecotype Columbia-0. Some transgenic and mutant lines have been described previously: ProLOG3::n3GFP (De Rybel et al., 2014), ProLOG4::n3GFP (De Rybel et al., 2014), ProRPS5A::TMO5-GR (De Rybel et al., 2013), ProRPS5A::TMO5-GR x ProRPS5A::LHW-GR (Smet et al., 2019). ProRPS5A overexpression lines (De Rybel et al., 2013) were used to generate the crosses: F1 seeds were used for RAM analysis, each seedling was genotyped to confirm the presence of the constructs. F2 seeds from genotyped plants were used for the SAM analysis. The n3GFP-GUS reporter lines were generated by MultiSite Gateway cloning (Karimi et al., 2007) in the pMK75*NfM14GW,0 destination vector. All constructs were transformed into the Arabidopsis thaliana Col-0 background.

For phenotype analyses, we used the following mutant lines: tmo5 (GK-143E03) (De Rybel et al., 2013), t5l1 (RIKEN_12-4602-1) (De Rybel et al., 2013), t5l2 (SALK_023629) (De Rybel et al., 2013), lhw (SALK_126132) (Ohashi-Ito et al., 2013a), ll1 (GK-262H03). Gene specific primers for genotyping were designed and are listed in Table S2, as are insertion-specific primers. For expression analysis in planta, ProLOG3::n3GFP and ProLOG4::n3GFP lines were crossed into lhw or tmo5 t5l1 mutant backgrounds. Homozygous plants were selected by PCR or antibiotic resistance as follows: final concentrations in cultivation medium of 25 mg/L kanamycin (Duchefa), 20 mg/L hygromycin (Duchefa), 10 mg/L sulfadiazine (Sigma-Aldrich).

Bacterial strains

Escherichia coli DH5α competent cells and Agrobacterium tumefaciens C58C1Rif(pMP90) LBA4404 were cultivated in LB medium supplemented with appropriate antibiotic in a shaker at 37 and 28°C, respectively.

METHOD DETAILS

Cultivation conditions

For root analysis, seeds were sterilized using a solution of 25% bleach and 75% ethanol. After 4 days of stratification at 4°C, plants were grown in half strength Murashige and Skoog medium (Duchefa) (Murashige and Skoog, 1962) without sugar and 0.8% Plant agar under continuous light conditions at 22°C. 10 mM dexamethasone (DEX) was used for induction of expression. For lateral meristem root analysis, plants were grown in half strength Murashige and Skoog medium (Duchefa) and 0.8% plant agar under continuous light conditions for 19–20 days at 22°C. For shoot analysis, plants were cultivated in soil under long-day conditions (16 h light/8 h dark) in growth chambers maintained at 21–22°C, with a light intensity of approximately 150 μmol m⁻¹ s⁻¹ and 40–60% relative humidity.

Shoot apical meristem dissection

Shoot apical meristems from inflorescence stems between 0.5 and 1.5 cm long were dissected and cultured in vitro for 3 h in a cultivation chamber as described previously (Brunoud et al., 2020). The meristems were stained with a water solution of 100 μg/mL propidium iodide (Sigma-Aldrich) for 5 min, then washed with water and used for microscopy.

Histochemical and histological procedures

For anatomical sections, 10-days-old roots were fixed overnight in 1% glutaraldehyde and 4% paraformaldehyde in 50 mM phosphate buffer, pH 7. Samples were dehydrated and embedded in Technovit 7100 resin (Heraeus Kulzer) according to the manufacturer’s protocol. For proper orientation of the samples, we used a two-step embedding methodology, with a pre-embedding step to facilitate orientation in 0.5 mL Eppendorf tubes (De Smet et al., 2004). Sections of 4 μm of root, taken 0.5 cm below junction between the root and the hypocotyl were cut with a Richert Jung microtome 2040, dried on Superfrost® plus microscopic slides (Menzel-Gläser), counterstained for cell walls with 0.05% ruthenium red for 5 min and rinsed in water. After drying, the sections were mounted in DPX mounting medium (Sigma-Aldrich) and covered with cover slips. Images were taken with an Olympus BX53 DIC microscope. mPS-PI staining was performed as described previously (Truernit and Haseloff, 2008). Briefly, the seedlings were fixed in 50% methanol and 10% acetic acid at 4°C for at least 12 h. Samples were then rinsed with water and incubated in 1% periodic acid (Sigma-Aldrich) for 40 min at room temperature (22°C). After another water rinse,
seedlings were incubated with Schiff’s reagent (100 nM sodium metabisulphite, 0.15N 37% HCl) with fresh propidium iodide (100 μg/mL) until visibly stained. To visualize, seedlings were transferred onto microscope slides in chloral hydrate solution. Quantification of vascular cell file numbers (cells within but excluding the pericycle) were performed using ImageJ software (https://imagej.nih.gov/ij/). Root apical meristems of n3GFP-GUS seedlings were stained with 0.1% Calcofluor White in ClearSee solution to visualize the cell wall (Ursache et al., 2018). To visualize GFP during secondary growth, a modified ClearSee protocol (Ursache et al., 2018; Ben-Targem et al., 2021) was employed. The most upper part of the root (0.5 cm below the hypocotyl root junction) was fixed with 4% PFA (Paraformaldehyde: Sigma, P6148) and 0.01% Triton in 1X PBS for 1 h under vacuum and embedded in 5% agarose blocks, then sections of 70–80 μm were obtained using a Vibratome (Leica VT-1000) and collected in water. Water was quickly replaced with ClearSee solution (10% xylitol 15% sodium deoxycholate, 25% urea) (Kurihara et al., 2015) and sections were kept in ClearSee for 24 h at room temperature and then stored at 4°C. Prior imaging, sections were stained with 0.05% Direct Red 23 (Sigma 212490) in ClearSee for 30 min, washed 3 times in ClearSee and mounted in ClearSee on a slide. Direct Red 23 stained the cell wall, and it was used to visualized cell outlay.

Microscopy
Confocal microscopy of shoot apical meristems was carried out using an upright Zeiss Axio Imager 2 equipped with a LSM700 confocal unit and 40×/1.0 DIC M27 water-dip objective. GFP was excited at 488 nm and detected at 490–530 nm; PI was excited at 555 nm and detected at 570–630 nm. Confocal microscopy of n3GFP-GUS root apical meristems was performed on a Leica SP8 using a 63× water-immersion objective. Calcofluor White and GFP were excited at 405 and 488 nm and visualized at 425–475 nm and 500–550 nm, respectively. mPS-Pi-stained roots were imaged at an excitation of 514 nm and emission of 600–650 nm. Confocal microscopy of root sections undergoing secondary growth was performed on a Zeiss LSM880 using a 20× dry objective and digital zoom. GFP and Direct Red 23 were excited at 488 and 561 nm and visualized at 490–544 nm and 580–642 nm, respectively. DIC microscopy of embedded samples was done using an Olympus BX53 microscope equipped with 10×, 20× and 40× air objectives.

cDNA synthesis
Total RNA was prepared from 100 mg of 11-day-old seedlings with the NucleoSpin RNA Plant Kit (Macherey-Nagel) according to the manufacturer’s instructions. cDNA was synthesized from 500 ng of total RNA using the SuperScript™ III Reverse Transcriptase Kit (Invitrogen).

Plasmid construction
DNA fragments were released by restriction from existing plasmids or amplified by PCR using primers synthesized by Sigma-Aldrich or Eurofins. The PCR reactions were performed using Q5 High-Fidelity DNA Polymerase (New England Biolabs). Gel extractions were performed using NucleoSpin Gel and PCR Clean-up Kits (Macherey-Nagel). Vectors were assembled via AQUA cloning technology (Beyer et al., 2015) and transformed into chemically competent E. coli strain 10-beta (New England Biolabs) or TOP10 (Invitrogen). Plasmid purifications were performed utilising Wizard Plus SV Minipreps DNA Purification Systems (Promega). New plasmids were tested by restriction enzyme digests and sequencing (Eurofins/GATC or Microsynth). All restriction enzymes were purchased from New England Biolabs. For testing promoters, the plasmid pMP010 was constructed as follows: the firefly luciferase gene (FLuc) was amplified by PCR from the pM2836 plasmid (Müller et al., 2014) using the oligonucleotides oMP025 and oMP029. The product was assembled via AQUA cloning into pGEN16 (Samodelov et al., 2016) digested with SacII/Xhol. Promoter sequences of LOG genes upstream from the ATG were amplified from genomic DNA extracted from 7-day-old seedlings using primers as follows: ProLOG1 (oMP036 and oMP037, 3138 bp), ProLOG3 (oMP040 and oMP041, 3564 bp), ProLOG4 (oMP022 and oMP023, 3999 bp), ProLOG5 (oMP042 and oMP043, 3024 bp), ProLOG7 (oMP026 and oMP027, 3187 bp). The products were inserted via AQUA cloning into pMP010 digested with SacII/Agel. For preparing vectors harboring cDNA of transcription factors, the plasmid pMP011 was constructed as follows: the nucleotide sequence of the HA tag (YPYDVPDYA) was amplified by PCR from the pMZ836 plasmid (Müller et al., 2014) using the oligonucleotides oMP025 and oMP029. The product was assembled via AQUA cloning into pGEN16 digested with AgeI/XhoI. Nucleotide sequences of transcription factors were amplified by PCR using the oligonucleotides oMP088 and oMP089. The product was assembled via AQUA cloning into pGEN16 digested with AgeI/Xhol. Nucleotide sequences of transcription factors were amplified from cDNA prepared previously using primers as follows: cTMOS (oMP107 and oMP108), cTSL1 (oMP121 and oMP122), cTSL2 (oMP125 and oMP126), cTSL3 (oMP123 and oMP124), cLHW (oMP109 and oMP110), cL1 (oMP131 and oMP132), cL2 (oMP129 and oMP130), cL3 (oMP127 and oMP128). PCR products were fused via AQUA cloning into pMP011 digested with AgeI/BstZ17I. All primers and plasmids used in this study are listed in Tables S2 and S3, respectively.
Luciferase protoplast assay
Protoplasts were isolated from shoots of 2- to 3-week-old Arabidopsis thaliana plants. Floatation was employed for isolation, and plasmids were transformed using a polyethylene-glycol-mediated approach as described previously (Ochoa-Fernandez et al., 2020). Plasmids were prepared with a Wizard® Plus Midi-preps DNA Purification System (Macherey-Nagel). Protoplasts were co-transformed with mixtures of the appropriate plasmids, 30 μg DNA in total. The transformed protoplasts were cultivated for 18–20 h at 19–22°C in the dark. After incubation, protoplasts were divided into aliquots of sufficient volume to measure six technical replicates for each sample. Firefly (FLuc) and Renilla luciferase (RLuc, in GB0109, (Sarrion-Perdigones et al., 2013)) activities were simultaneously quantified in intact protoplasts as described (Ochoa-Fernandez et al., 2016). Substrates for both luciferases were added directly before measurement: D-luciferin (Biosynth AG) for FLuc, Coelenterazine (Carl Roth) for RLuc. Chemiluminescence measurements were performed using a Berthold Centro XS3 LB 960 microplate luminometer (Berthold Technologies, Bad Wildbad, Germany) and a BertholdTriStar2 S LB 942 multimode plate reader (Berthold Technologies, Bad Wildbad, Germany). The FLuc/RLuc ratio was determined (n = 4–6) and showed in tables. Constitutively expressed RLuc served as an internal normalization element to obtain ratiometric data.

Quantitative analysis of shoot apical meristem
Images of shoot apical meristems stained with propidium iodide were segmented using an auto seeded 3D watershed algorithm derived from the MARS pipeline (Fernandez et al., 2010) in which the parameters were manually tuned for each sample. In the resulting segmented images, cells belonging to the L1, L2 and L3 layers were automatically identified. To do so, a triangle mesh representing the tissue surface was computed using the segmented image, and L1 cells were selected as those adjacent to the background region and closest to the vertices of the surface mesh. L2 and L3 cells were selected recursively by adjacency to cells belonging to the previous layer.

Finally, “meristematic cells” (cells belonging to the central zone, the peripheral zone and to organ initials) were distinguished from cells of organ primordia and boundaries using the surface curvature. Principal curvatures were estimated on the surface mesh based on the vertex normal vectors (Theisel et al., 2004), and a central meristematic region was identified by thresholding the minimum principal curvature value and performing morphological operations. The retained threshold value was −0.005 μm−1. The resulting binary property was projected on the closest L1 cells to identify L1 meristematic cells, and then propagated to L2 and L3 cells by adjacency with a triangle of meristematic cells at the previous layer. The results were obtained by filtering out non-meristematic cells and pooling the cell measures by cell layer.

Accession numbers
The sequence data of genes described this article can be found in The Arabidopsis Information Resource (https://www.arabidopsis.org/) or GenBank (http://www.ncbi.nlm.nih.gov/genbank/) databases under the following accession numbers: AT3G25710 for TMO5/bHLH32, AT1G68810 for T5L1/bHLH30/ABS5, AT3G56770 for T5L2/bHLH107, AT2G41130 for T5L3/bHLH106/STC8, AT2G27230 for LHW/bHLH156, AT1G64625 for LL1/LHL3/bHLH157, AT2G31280 for LL2/LHL2/bHLH155, AT1G06150 for LL3/LHL1/EMB1444, AT2G28305 for LOG1, AT2G37210 for LOG3, AT3G53450 for LOG4, AT4G35190 for LOG5, and AT5G06300 for LOG7.

QUANTIFICATION AND STATISTICAL ANALYSIS
All statistical analysis plots were generated using the PlotsOfData webtool at standard settings (https://huygens.science.uva.nl/PlotsOfData/). In all boxplots, boxes represent the 1st and 3rd quartiles, and the centre line represents the median. The lowercase letters associated with the boxplots indicate significantly different groups as determined by one-way analysis of variance (ANOVA) with post-hoc Tukey HSD testing (p < 0.001).