



Non-cell autonomous and spatiotemporal signalling from a tissue organizer orchestrates root vascular development

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During plant development, a precise balance of cytokinin is crucial for correct growth and patterning, but it remains unclear how this is achieved across different cell types and in the context of a growing organ. Here we show that in the root apical meristem, the TMO5/LHW complex increases active cytokinin levels via two cooperatively acting enzymes. By profiling the transcriptomic changes of increased cytokinin at single-cell level, we further show that this effect is counteracted by a tissue-specific increase in CYTOKININ OXIDASE 3 expression via direct activation of the mobile transcription factor SHORTROOT. In summary, we show that within the root meristem, xylem cells act as a local organizer of vascular development by non-autonomously regulating cytokinin levels in neighbouring procambium cells via sequential induction and repression modules.

The plant vasculature is a complex tissue composed of multiple cell types, each with a specific function¹. In the *Arabidopsis* root apical meristem during primary growth, vascular tissues are organized according to a bilateral symmetry, with a central xylem axis flanked by two phloem poles with intervening procambium cells^{1,2}. Previous work has shown that this patterning is established and maintained by a domain of high-auxin signalling in the xylem cells and a domain of high-cytokinin signalling in the procambium and phloem cell lineages^{3–5}, making this an excellent model system to study coordinated development involving intercellular communication, hormonal signalling and crosstalk. On a molecular level, growth and patterning of vascular tissues is in part driven by the heterodimer formed by the basic helix loop helix transcription factors TARGET OF MONOPTEROS 5 and LONESOME HIGHWAY (TMO5/LHW)^{6–11}. This complex triggers local biosynthesis of cytokinin via direct activation of LONELY GUY 3 and 4 (LOG3/4)^{4,5}. This xylem-derived cytokinin is thought to diffuse to neighbouring procambium cells where it drives vascular proliferation by activating downstream target genes, including members of the DNA-binding-with-one-finger (DOF)-type transcription factor family^{12,13}. Although it is clear that TMO5/LHW plays an important role in controlling vascular growth and patterning^{4,5}, it remains unclear how appropriate cytokinin levels are maintained in each cell type in the context of a growing tissue¹⁴.

BGLU44 and LOG4 cooperatively produce active cytokinin

TMO5/LHW activity is dependent on the phytohormone cytokinin as this dimer is inactive when cytokinin biosynthesis (for example,

in a *log1234578* mutant^{15,16}) or signalling (for example, in a *wol* mutant¹⁷) are perturbed⁴. Although *LOG3* and *LOG4* were identified as main target genes of the TMO5/LHW dimer^{4,5}, misexpression of *LOG* genes does not result in the strong cytokinin-related vascular phenotypes observed upon exogenous cytokinin treatment or TMO5/LHW induction⁴, suggesting that additional factors are involved in releasing active cytokinin. To identify such factors, we overlapped genes co-expressed with *LOG4* in a high spatiotemporal resolution single-cell dataset (Fig. 1a, Extended Data Fig. 1 and Supplementary Data 1)¹⁸ with a list of putative TMO5/LHW target genes¹³. The overlap contained the closely related *LOG3*^{4,5}, the negative regulator of TMO5/LHW activity *SACL3*^{7,11} and an uncharacterized beta-glucosidase family member *BGLU44/AT3G18080* (Fig. 1a, Extended Data Fig. 2a–c and Supplementary Data 1). By quantitative reverse transcription–polymerase chain reaction (RT-qPCR) analysis, we found that relative expression levels of *BGLU44* were increased upon TMO5/LHW induction and reduced in *tmo5* single, double and triple mutant backgrounds (Fig. 1b), similar to *LOG4*⁴. We next constructed a pBGLU44-nYFP/GUS reporter line and found *BGLU44* expressed in the root apical meristem along the xylem axis and in xylem pole-associated pericycle and endodermis cells as predicted by single-cell RNA-seq (scRNA-seq) atlas data¹⁸ (Fig. 1c,d and Extended Data Fig. 2a–c), and its expression patterns identical to those of *LOG4* in this tissue⁴. Induction of the TMO5/LHW heterodimer throughout the root meristem (using a dexamethasone (DEX) double inducible pRPS5A::TMO5:GR x pRPS5A::LHW:GR or dGR line¹³) triggered both increased and

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ectopic *pBGLU44::nYFP/GUS* expression (Fig. 1e,f and Extended Data Fig. 2d–g), suggesting that BGLU44 acts downstream of TMO5/LHW. By chromatin immunoprecipitation (ChIP)-RT-qPCR analysis, we found a significant binding of TMO5-GR/LHW-GR to the *BGLU44* promoter region, indicating that *BGLU44* is a direct target of TMO5/LHW (Extended Data Fig. 3a). Taken together, these results confirm the TMO5/LHW-dependent co-expression of *BGLU44* and *LOG4*.

BGLU44 encodes a member of the glycoside hydrolase family 1, comprising over 40 members in both *Arabidopsis* and rice¹⁹. Although *BGLU* proteins have been implicated in various developmental processes including mobilization of storage compounds²⁰ and reconstruction of cell walls²¹, the beta-glucosidase Zm-p60.1 in maize was shown to cleave biologically inactive cytokinin conjugates to release active cytokinin²². To investigate the possibility that *BGLU44* would have a similar role in *Arabidopsis*, we misexpressed *BGLU44* from the strong meristematic *RPS5A* promoter²³ (Extended Data Fig. 2h–j) and analysed xylem differentiation in the root as proxy for active cytokinin levels⁴. As positive control, TMO5/LHW misexpression (dGR line) resulted in a loss of protoxylem differentiation compared to the wild type control situation (Fig. 1k,l). We did not observe any differences in the *pRPS5A::BGLU44* line compared to wild type plants (Fig. 1g,i,l and Extended Data Fig. 2h). Considering that the *pRPS5A::LOG4* misexpression line shows mild vascular defects in the root meristem⁴ (Fig. 1g,h,l), we hypothesized that both enzymes might work in a cooperative manner. We thus combined both misexpression lines via crossing (*pRPS5A::LOG4* x *pRPS5A::BGLU44*) and observed a defect in siliques positioning on the stem (Extended Data Fig. 2h,i) and an increase in root hairs (Extended Data Fig. 2t–w) as also seen in TMO5/LHW-misexpressing¹⁸ and cytokinin-overproducing plants²⁴. When analysing the root meristem vascular tissues, we found an almost complete loss of protoxylem differentiation in the root (Fig. 1j–l), phenocopying the higher cytokinin levels found in TMO5/LHW misexpression lines⁴. Fitting with this observation, combined misexpression of *LOG4* and *BGLU44* resulted in a small but significant increase in the number of vascular cell files (Extended Data Fig. 2k–o). Similarly, a newly generated *bglu44* loss-of-function CRISPR line, which led to a large fragment deletion, did not result in a strong phenotype, but enhanced the reduction of vascular cell numbers when combined with *log4* or *log3 log4* mutants (Extended Data Fig. 4), suggesting that *LOG4* and *BGLU44* play a role in vascular proliferation. To further understand whether increased *LOG4* and *BGLU44* expression are related to increased levels of active cytokinin, we next analysed the two-component signalling sensor nuclear (TCSn) reporter for cytokinin signalling^{13,25} fused to the nuclear tdTomato fluorescent protein (*pTCSn::ntdT*) in the combined misexpression background. Confocal imaging confirmed that plants overexpressing both *LOG4* and *BGLU44*, but not the individual factors, caused increased expression of TCSn, which

was most prominent in the ground tissues that typically show very low TCSn expression in wild-type plants (Fig. 1m–q and Extended Data Fig. 2p–s). Finally, we assayed the enzymatic activity of the *BGLU44* protein (Extended Data Fig. 4d). First, we tested specificity of *BGLU44* activity in vitro for several glucose-conjugated cytokinin substrates and found that it is specific to O-glucoside cytokinin species (Fig. 1r). Moreover, *BGLU44* is able to cleave the inactive conjugated tZOG and tZROG species into the bio-active tZ and tZR (Fig. 1s and Source Data Fig. 1). Measuring the endogenous cytokinin profiles of 7-day-old root tips in *LOG4*, *BGLU44*, *LOG4/BGLU44* and dGR misexpression lines revealed that the combined misexpression of *LOG4* and *BGLU44* resulted in a similar increase in cytokinin levels as previously shown for TMO5/LHW⁴ (Fig. 1t and Source Data Fig. 1). Taken together, our results show that *BGLU44* and *LOG4* cooperatively act downstream of TMO5/LHW to release active cytokinin in the vascular bundle of the root meristem, thereby controlling primary vascular development.

CKX3 balances cytokinin levels downstream of TMO5/LHW

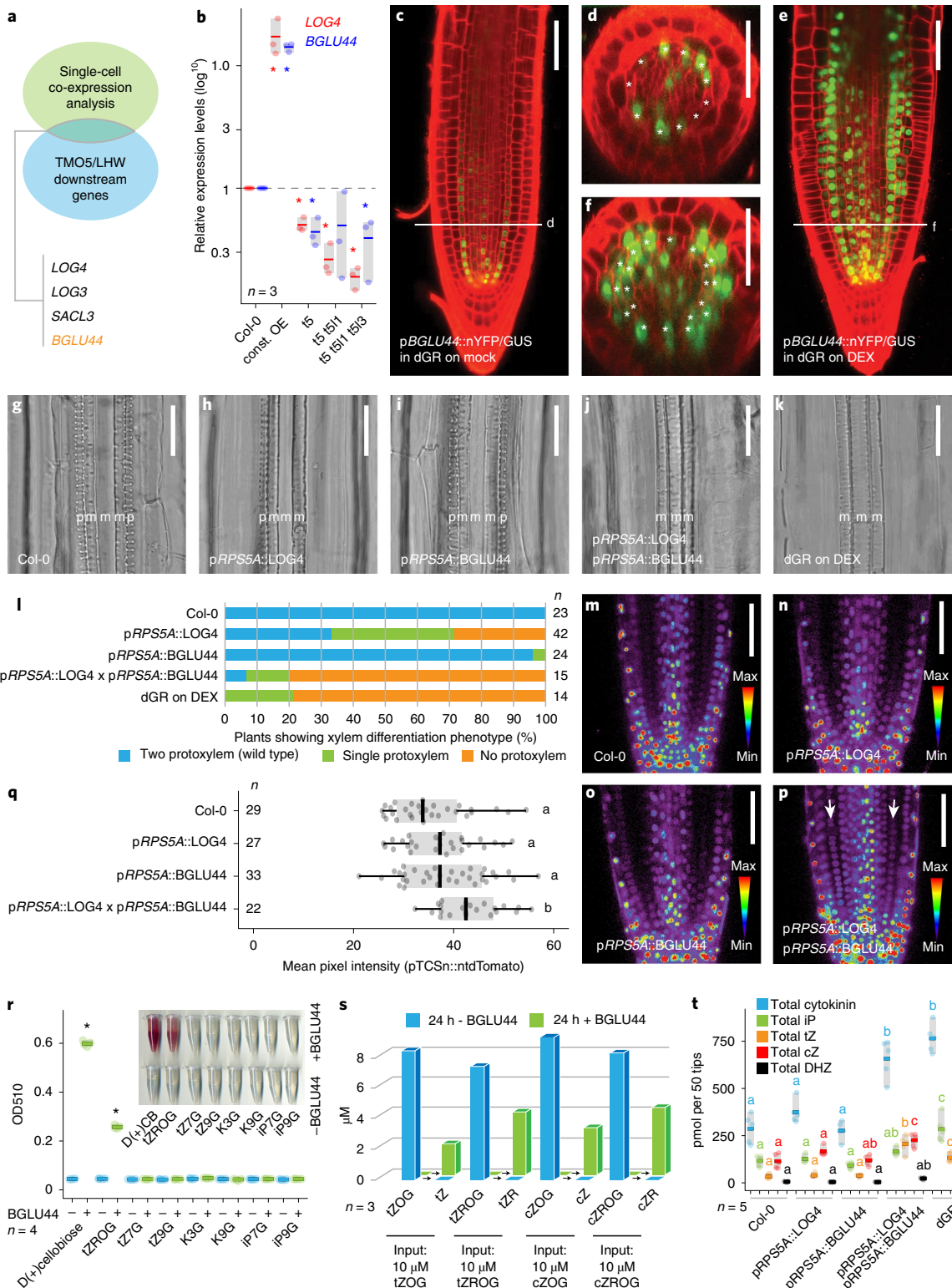
Cytokinin levels need to be well balanced in space and time to allow normal development. Indeed, high levels of active cytokinin disturb normal vascular cell proliferation, patterning and differentiation^{15,16,26,27} (Fig. 3n). The active cytokinin produced in the central xylem axis, where TMO5/LHW activity overlaps with *LOG4* and *BGLU44* expression, is thought to diffuse to neighbouring procambium and phloem cells^{3–5}, which show high levels of cytokinin signalling. To understand the tissue-specific responses of increased cytokinin levels, we profiled the transcriptional effect of cytokinin treatment on root meristem cells at single-cell resolution (Fig. 2a,b). After filtering, we retained about 10,000 high quality cells for each sample with a minimal unique molecular identifier (UMI) count of more than 1,000 (Fig. 2a). Clear transcriptional changes were observed for most cell types, while some subpopulations remained largely unaffected, such as the protoxylem and columella cells (Extended Data Fig. 5a–c), suggesting tissue-specific responses. As expected, primary response genes of the cytokinin signalling pathway, such as A-type ARABIDOPSIS RESPONSE REGULATORS (ARR), were recovered as cytokinin inducible in all cell clusters (Supplementary Data 2). We next created transcriptional reporter lines for 12 genes previously uncharacterized for cytokinin response in the root meristem and showing tissue-specific cytokinin responses. These lines show both the predicted tissue-specific expression pattern in the mock situation and the tissue-specific induction kinetics after cytokinin treatment in the root (Supplementary Fig. 1), thus validating the predictive power of our dataset.

Given that procambium cells are those responding to cytokinin levels with respect to cell proliferation in our system, we next searched our dataset for those genes specifically responding to the

Fig. 1 | *BGLU44* and *LOG4* cooperatively produce active cytokinin downstream of TMO5/LHW. **a**, Schematic representation of the strategy to identify *BGLU44* as putative target gene of TMO5/LHW. **b**, Relative expression levels of *BGLU44* and *LOG4* in wild type (Col-0), TMO5/LHW misexpression, and *tmo5*, *tmo5 tmo5like1* (*t5 t5l1*) and *tmo5 tmo5like1 tmo5like3* (*t5 t5l1 t5l3*) mutant backgrounds. The coloured lines indicate the median value of three biological replicates ($n=3$), while the grey area shows the data range (* $P < 0.05$). The dashed black line indicates the expression level of the control sample, which was set to 1. **c–f**, Expression of *pBGLU44::nYFP/GUS* in the dGR root meristem grown on mock medium and transferred to mock or 10 μ M DEX for 24 h. Asterisks indicate the endodermis cell layer. **g–k**, Microscopic images of xylem differentiation in the mentioned genotypes. **p**, protoxylem; **m**, metaxylem. **l**, Quantification of the different classes of xylem phenotypes shown in panels **g–k**. **m–p**, Confocal images of root meristems expressing *pTCSn::ntdT* reporter in the mentioned genotypes. Arrows in **p** indicate cortex cell layer. **q**, Quantification of the *pTCSn::ntdT* mean pixel intensity in the mentioned genotypes. Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. **r**, In vitro *BGLU44* enzymatic activity on a range of cytokinin glycoside substrates. **s**, In vitro cleavage of O-glucosylated cytokinins by *BGLU44*. **t**, Overview of total endogenous cytokinin levels in root tips of the indicated lines. Lowercase letters in graphs indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($P < 0.001$). Asterisks in graphs indicate significance values as determined by standard two-sided *t*-tests. Coloured lines indicate mean values and grey boxes indicate data ranges. Scale bars in **c–k** and **m–p**, 50 μ m. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Source Data Fig. 1. The experiments in **c–f** were repeated three times independently, with similar results.

increase in cytokinin levels in the procambium cell cluster only. We found that among the top candidates, *CYTOKININ OXIDASE 3* (*CKX3*/AT5G56970) was recovered as specifically induced by cytokinin in procambium cells (Fig. 2c,d and Supplementary Data 2). As CKX proteins have been shown to reduce levels of active cytokinin^{24,26,28}, *CKX3* could be the factor that counteracts the flow of active cytokinin from the xylem cells and ensure well-balanced cytokinin levels. Indeed, previous reports suggested that cytokinin

could balance itself by promoting cytokinin oxidase expression^{29,30}. We confirmed the cytokinin inducibility of *CKX3* in procambium cells by generating a p*CKX3::nYFP-GUS* transcriptional reporter using a 4.2 kb promoter fragment (Fig. 2e,f). Upon transfer to 10 μM 6-benzylaminopurine (BAP) for 3 h, a significant increase in *CKX3* expression levels in procambium cells was observed (Fig. 2g–i) as predicted by the scRNA-seq dataset (Fig. 2c,d and Extended Data Fig. 5e).



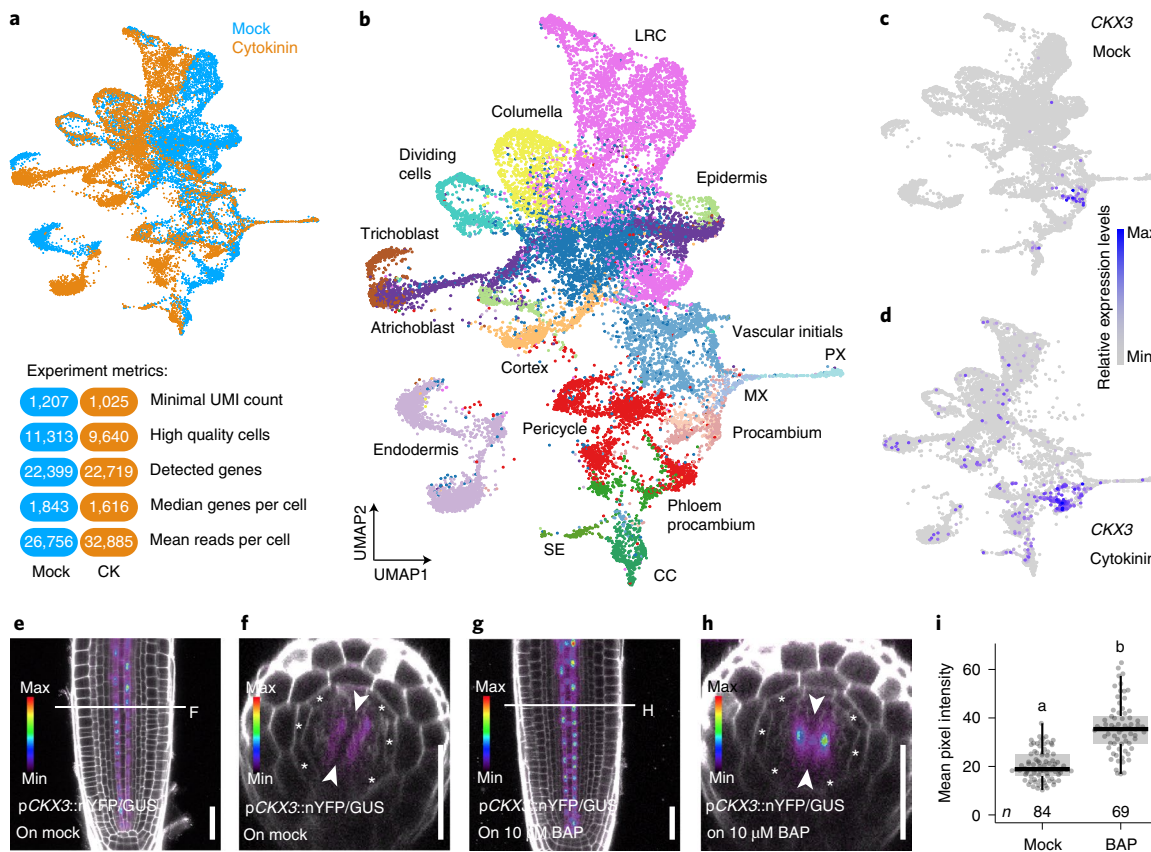


Fig. 2 | Single-cell transcriptional changes in root meristem cells upon cytokinin treatment. **a**, Uniform Manifold Approximation and Projection (UMAP) plot showing the merge of the mock and cytokinin samples, and an overview of the experimental metrics for both samples. **b**, UMAP plot of the merged data with indications of the different cell identities. LRC, lateral root cap; PX, protoxylem; MX, metaxylem; SE, sieve element; CC, companion cell. The most central dark blue cell cluster is the initial cell cluster. **c,d**, Feature plots of CKX3 expression in the mock (**c**) and cytokinin (**d**) datasets showing a tissue-specific induction in the procambium cells. **e–h**, Expression of pCKX3::nYFP/GUS in the root meristem grown on mock medium and transferred to mock or 10 µM BAP for 3 h. Asterisks indicate the endodermis cell layer. Arrowheads indicate xylem axis. Scale bars, 50 µm. **i**, Quantification of the experiment described in **e–h**. Lowercase letters on top of the boxplots indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($P < 0.001$). Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Source Data Fig. 2.

To further understand whether the CKX3 regulation by cytokinin is related to TMO5/LHW, we first confirmed by RT-qPCR that relative expression levels are increased upon induction of the TMO5/LHW heterodimer and downregulated in *tmo5* double and triple mutants (Fig. 3a), providing support that CKX3 indeed acts downstream of TMO5/LHW. We next introduced the transcriptional pCKX3::nYFP/GUS line (Fig. 3b–e) and a newly generated translational pCKX3::CKX3:GFP reporter line (Fig. 3f–i) in the dGR background. Upon misexpression of TMO5/LHW, expression of CKX3 increased and now marked the entire vascular cylinder (Fig. 3d,e,h,i). To further assess a possible role for CKX3 during vascular development, we analysed phenotypes upon loss of function of CKX3 and its close homologue CKX5, which shows a similar expression pattern as predicted by our scRNA-seq dataset (Extended Data Figs. 5f and 6a,b) and validated by a newly generated 3.5 kb pCKX5::nYFP/GUS reporter line (Extended Data Fig. 6c,d). The *ckx3 ckx5* double mutant²⁴, but not the *ckx3* or *ckx5* single mutants, showed additional metaxylem cell files in over 60% of plants analysed (Extended Data Fig. 6e–h,l) and vascular cell file number was increased (Extended Data Fig. 6m–q). This is opposite to the effect of reducing cytokinin biosynthesis in *log* higher-order mutants (Extended Data Fig. 6i–l)^{4,16}. These results suggest that CKX3 (in collaboration with CKX5) is involved in modulating vascular cytokinin levels. To further investigate the importance of CKX3 function

downstream of TMO5/LHW, we generated pRPS5A::CKX3:YFP misexpression lines in the dGR background. Unlike the dGR control situation where DEX treatment inhibited protoxylem differentiation (Fig. 3j,k,o) and increased vascular cell file number (Fig. 3p,q,t)⁶, CKX3 misexpression completely repressed TMO5/LHW function (Fig. 3l,m,o,r–t). These results show that TMO5/LHW not only promotes release of active cytokinin via LOG4 and BGLU44, but at the same time represses active cytokinin in procambium cells via CKX3 to maintain optimal levels of cytokinin for normal vascular development. To further examine the importance of TMO5/LHW in CKX3 regulation, we treated wild type and *tmo5* triple mutants with exogenous cytokinin. While cytokinin increased relative expression levels of CKX3 in a wild-type Col-0 background, this response was repressed in the absence of TMO5 activity (Extended Data Fig. 6r), suggesting that TMO5/LHW is an important regulator of CKX3 expression. Considering that TMO5 and CKX3 expression domains are in neighbouring cell types, we hypothesize that there must be a mobile intermediate connecting these two factors.

SHR bridges TMO5/LHW-dependent regulation of CKX3 expression

The mobile transcription factor SHORT-ROOT (SHR) has been shown to directly bind to the CKX3 promoter region and regulate its expression levels³¹. Although cytokinin levels³¹ and signalling

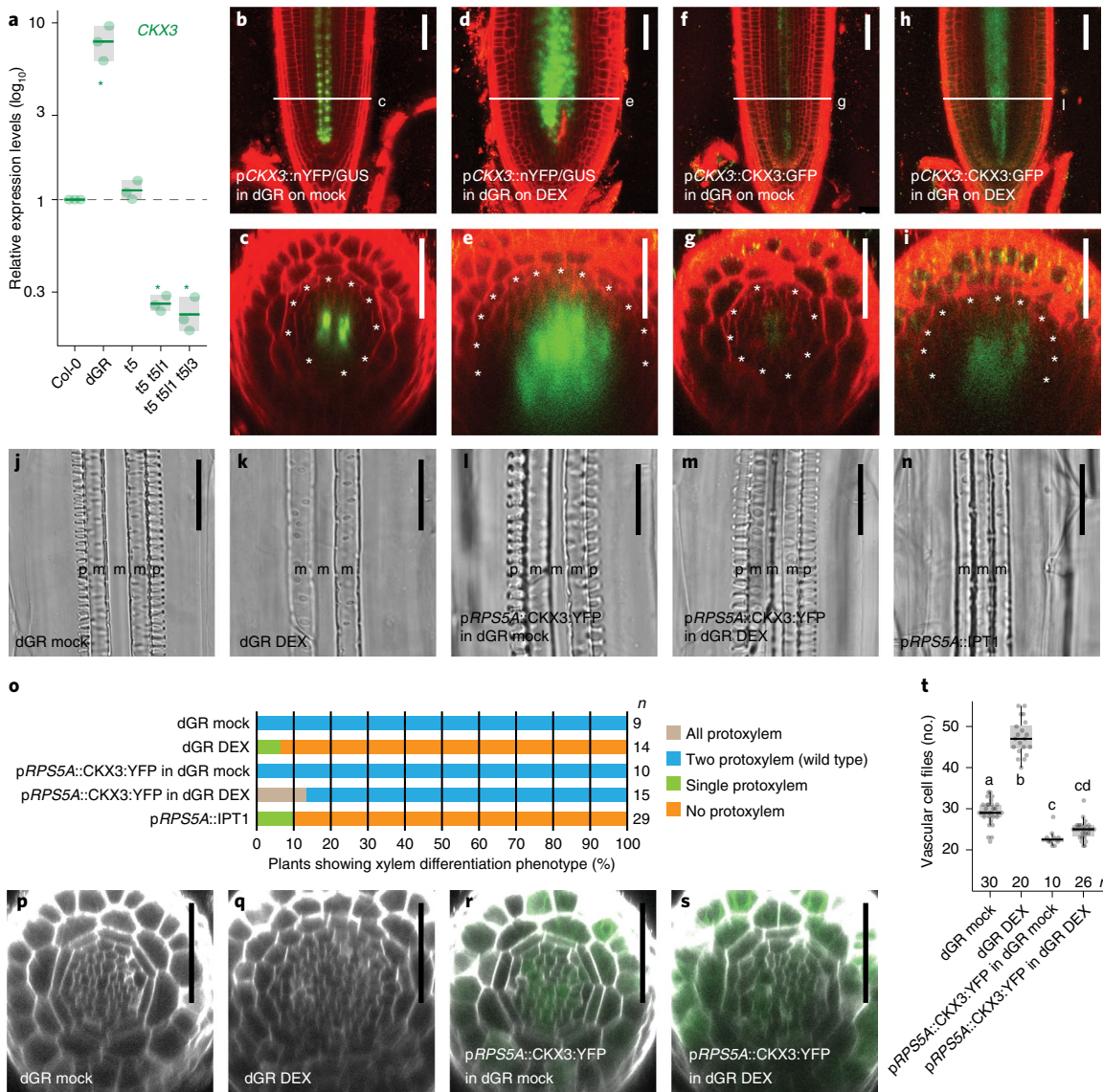


Fig. 3 | CKX3 balances cytokinin levels downstream of TMO5/LHW. **a**, Relative expression levels of CKX3 in wild type (Col-0), TMO5/LHW misexpression, and *tmo5*, *tmo5 tmo5like1* (*t5 t5*l1**) and *tmo5 tmo5like1 tmo5like3* (*t5 t5*l1* t5*l3**) mutant backgrounds. Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. $n=3$ biological replicates (* $P < 0.05$). The dashed black line indicates the expression level of the control sample, which was set to 1. **b–i**, Expression of pCKX3::nYFP/GUS and pCKX3::CKX3:GFP in the dGR root meristem grown on mock medium and transferred to mock or 10 μ M DEX for 24 h. Asterisks indicate the endodermis cell layer. **j–n**, Microscopy images of xylem differentiation in the mentioned genotypes. p, protoxylem; m, metaxylem. **o**, Quantification of the different classes of xylem phenotypes shown in **j–n**. **p–s**, Confocal cross-section images through the root meristem of the mentioned genotypes grown on mock medium and transferred to mock or 10 μ M DEX for 24 h. **t**, Quantification of the number of vascular cell files shown in **p–s**. Lowercase letters on top of the boxplots indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($P < 0.001$). Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. Asterisks in graphs indicate significance values as determined by standard two-sided t-tests. Black lines indicate mean values and grey boxes indicate data ranges. Scale bars in **b–i** and **p–s**, 50 μ m. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Source Data Fig. 3. The experiments in **b–i** were repeated three times independently, with similar results.

(Extended Data Fig. 7a–c) are high in a *shr-2* mutant background, CKX3 expression levels are reduced³¹. This suggests that although CKX3 expression levels can be controlled by cytokinin and SHR, both of which are mobile in the vascular tissues, the dominant form of regulation acts via SHR. To explore the importance of SHR in the connection between TMO5/LHW and CKX3 regulation during vascular development, we first introduced a pSHR::SHR:GFP translational reporter line in dGR plants. Upon induction of TMO5/LHW by DEX treatment, the SHR:GFP fusion protein was ectopically present throughout the root meristem (Extended

Data Fig. 7d–g). To understand if this was caused by regulation at the transcriptional level, we next introduced a pSHR::nYFP/GUS transcriptional reporter line in dGR. Also in this case, induction of TMO5/LHW caused ectopic expression of SHR throughout the root meristem (Fig. 4a–d), suggesting that TMO5/LHW might control SHR expression. These results were further supported by RT-qPCR data showing that relative expression levels of SHR were increased in a TMO5/LHW heterodimer misexpression line and repressed in *tmo5* higher order mutant lines (Extended Data Fig. 7h). To evaluate whether TMO5/LHW directly activates SHR

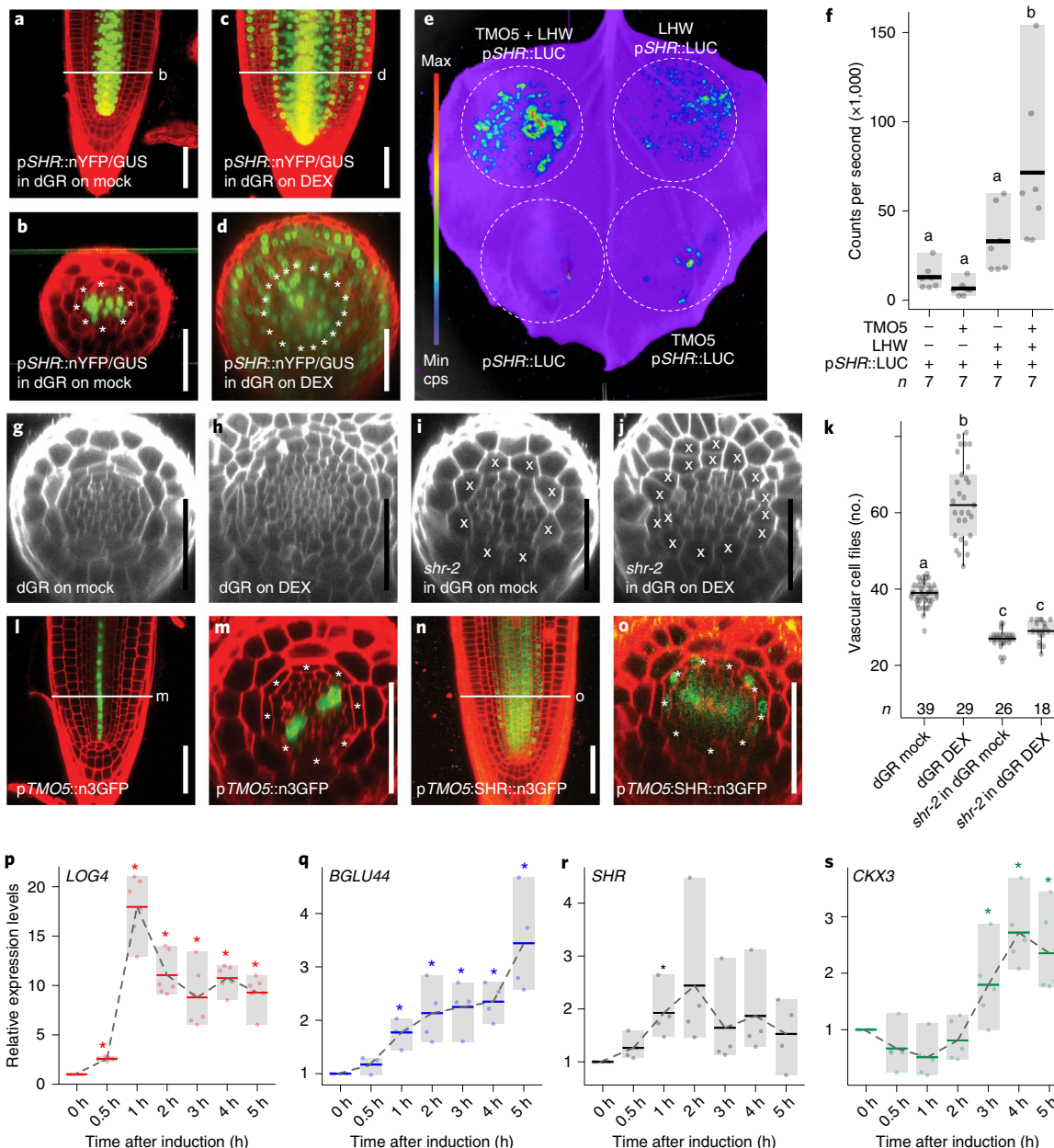


Fig. 4 | SHR bridges TMO5/LHW-dependent regulation of CKX3 expression. **a-d**, Expression of pSHR::nYFP/GUS in the dGR root meristem grown on mock medium and transferred to mock or 10 µM DEX for 24 h. Asterisks indicate the endodermis cell layer. **e,f**, Transient luciferase assay in tobacco leaves showing pSHR::LUC expression in the mentioned combination of introduced constructs. Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. $n=7$ independent tobacco leaves. **g-j**, Confocal cross-section images through the root meristem of the mentioned genotypes grown on mock medium and transferred to mock or 10 µM DEX for 24 h. The x in **i** and **j** indicates cells with mixed cortex-endodermis identity in the *shr-2* mutant. **k**, Quantification of the number of vascular and endodermis cell files shown in panels **g-j**. The x in **i** and **j** indicates cells with mixed cortex-endodermis identity in the *shr-2* mutant. Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. **l-o**, Expression of pTMO5::n3GFP (**l,m**) and pTMO5::SHR::GFP (**n,o**) in the root meristem grown on mock medium. **p-s**, Relative expression levels of LOG4, BGLU44, SHR and CKX3 in dGR grown on mock and transferred to 10 µM DEX for the indicated time before sampling. Lowercase letters on top of the graphs indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($P<0.001$). Boxes represent the 1st and 3rd quartiles, and the centre line represents the median. $n=4-6$ biological replicates. Asterisks in graphs indicate significance values as determined by standard two-sided *t*-tests (* $P<0.05$). Coloured lines indicate mean values and grey boxes indicate data ranges. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Source Data Fig. 4. The experiments in **a-d** and **i-o** were repeated three times independently, with similar results.

expression, we fused a 2.5 kb promoter fragment of *SHR*³² to luciferase and introduced this construct in tobacco leaves in the presence of TMO5/LHW. pSHR::LUC was induced by TMO5/LHW, while this was not the case in the negative controls and not significantly different in the presence of the individual members of

the dimer (Fig. 4e,f), fitting with the previous findings that TMO5 and LHW act as an obligate heterodimer^{6,9}. These results were confirmed by ChIP-RT-qPCR (Extended Data Fig. 3b). Taken together, these results suggest that TMO5/LHW directly binds to the *SHR* promoter region to activate its expression.

To further study the genetic relationship between SHR and the TMO5/LHW heterodimer complex, we introduced the *shr-2* mutation³³ into a dGR background by crossing. Although some pericinal divisions were still observed, vascular cell file numbers were not significantly increased in the presence of the *shr-2* mutation (Fig. 4g–k), suggesting that SHR is required for normal TMO5/LHW function. We observed a loss of protoxylem differentiation in *shr-2*, both with and without induction of dGR (Extended Data Fig. 7i–m), in line with the high cytokinin levels in this mutant background³¹. Moreover, we found that xylem-expressed SHR (pTMO5::SHR:GFP) is capable of moving throughout the vascular bundle (Fig. 4l–o), and is capable of rescuing the *shr-2* root length and ground tissue cell identity phenotypes in a dose-dependent manner (Extended Data Fig. 8). This indicates that TMO5/LHW-driven SHR protein can move from xylem cells into adjacent procambium cells to regulate CKX3 expression, thereby balancing overall cytokinin levels. The relevance of TMO5/LHW regulation on the extended SHR transcriptional pathway was further highlighted by the fact that also its interactor SCARECROW and downstream target miRNA165 are controlled by TMO5/LHW activity (Extended Data Fig. 7n–s). As such, the xylem axis works as a central organizer for vascular development and patterning via SHR.

Spatiotemporal regulation of vascular cytokinin levels

Our results suggest that downstream of TMO5/LHW, the LOG4 and BGLU44 enzymes cooperatively work to increase levels of active cytokinin in the xylem axis, while CKX3 reduces cytokinin levels in the neighbouring procambium cells via SHR as a mobile intermediate, and independent of cytokinin signalling. These results are in line with the hypothesis that the xylem axis acts as an insensitive source of active cytokinin, which is thought to diffuse to the neighbouring procambium cells^{4,5}. In the procambium domain, cells are responsive to cytokinin^{3,4,17,27} and thus require a repressive mechanism to cope with the influx of active cytokinin and obtain optimal levels for normal development. We next questioned whether these different factors responsible for the increase and decrease of active cytokinin levels are activated simultaneously or in sequence. By analysing the temporal changes in expression levels via RT-qPCR, we show that *LOG4* is increased in expression levels around 30 min to 1 h after TMO5/LHW induction (Fig. 4p), quickly followed by *BGLU44* induction starting around 1 h after TMO5/LHW induction (Fig. 4q). *SHR* expression levels are also induced from 1 h onwards (Fig. 4r), leading to the induction of CKX3 as a direct SHR target at 3 h after induction (Fig. 4s). A similar trend was observed when analysing the transcriptional reporter lines for *LOG4*, *BGLU44* and *SHR*, and a protein fusion reporter for CKX3 genes (Supplementary Figs. 2a–c and 3), further corroborating that induction of active cytokinin levels via LOG4 and BGLU44, and repression via SHR and CKX3 are sequential events upon TMO5/LHW induction. Previous mathematical models^{4,34} had predicted a network that was conceptually sufficient to determine the correct position of auxin response around an initial asymmetry in cellular pattern. However, these models considered only a fixed basal rate of auxin synthesis, and did not consider how robust this output would be in regards to fluctuations in auxin input. To better understand the spatiotemporal wiring of this network, we generated a mathematical model comprising these molecular players in their respective cells (for a detailed description, see Supplementary Methods: Mathematical Modelling), with auxin signalling as input to TMO5/LHW and cytokinin signalling as output. This model was based around a simplified tissue geometry, but considered fluctuations in the auxin signalling input over time. In the model, the wiring of activation and repression modules is capable of dampening fast oscillating auxin inputs into a continuous cytokinin response, while remaining responsive to slower changes in auxin (Model Description). This response was not dependent on the amplitude of the auxin

signalling input, meaning that both small and large changes have an impact on the cytokinin signalling output as long as the frequency of the modulation is low (Model Description).

In conclusion, our findings point to a tight spatiotemporal regulation of cytokinin levels via sequential induction and repression modules, all originating from the same bHLH heterodimer complex (Extended Data Fig. 9). As such, we found that a single transcription factor complex controls multiple biosynthesis and degradation steps of a phytohormone to regulate tissue development in space and time. The fact that these modules with opposite function are initiated as direct targets of the TMO5/LHW complex suggests that cytokinin levels might be balanced without the need for intermediate sensing via canonical CRE1/AHK4/WOL receptor signalling^{17,35}. Rather, differences in spatiotemporal activity of these modules might be sufficient to control levels of active cytokinin in the respective tissues and drive the observed self-organizing capacities of vascular patterning and growth^{4,34,36}. Intriguingly, our modelling efforts suggest that these cytokinin signalling-controlled processes would not be influenced by fast fluctuations in auxin signalling in the root apical meristem. Rather, vascular patterning and growth controlled by the TMO5/LHW dimer would be sensitive to slow and gradual changes in the auxin input. Although additional experimental work is required to support this hypothesis, this emerging property of the model makes sense in the context of a growing tissue where growth and patterning respond to gradual modulation of hormone levels. Our work also identifies the central xylem cells as a tissue organizer for vascular development and highlights TMO5/LHW as upstream regulator of SHR, a central transcriptional hub controlling several distinct aspects of growth and development in vascular and other tissues^{33,37–39}. Given that MONOPTEROS was previously shown to control SHR expression⁴⁰, it is conceivable that within the context of vascular development this regulation is indirect and requires TMO5 as an intermediate factor. It remains to be determined if and how TMO5/LHW would be connected to other SHR-controlled processes during plant development.

Methods

Plant materials and growth condition. *Arabidopsis thaliana* ecotype Columbia-0 (Col-0) was used in all transformation and phenotype analysis. All seeds were germinated on ½MS (Murashige and Skoog, Duchefa) medium after 3 d stratification at 4°C. Seedlings and plants were grown at 22°C in continuous light conditions. Glucocorticoid receptor (GR)-related inducible plants (7 days old) were transferred from 1/2 MS to 10 µM DEX-supplemented medium and allowed to grow for the indicated time. Plant lines and mutants used in this study are dGR¹³, pSHR::SHR:GFP³⁸, pTCSn::tdTomato¹³, pRPSA::LOG4¹, cck3³⁴, cck5³⁴, cck3 cck5³⁴, log4¹⁶, log3 log4¹⁶, log3 log4 log7¹⁶, shr-2³³, tmo5⁶, tmo5t 5l¹⁶ and tmo5 t5l1 t5l³³. The Arabidopsis Genome Initiative identifiers for the genes used in this study are as follows: *TMO5*, AT3G25710; *LHW*, AT2G27230; *LOG4*, AT3G3450; *BGLU44*, AT3G18080; *CKX3*, AT5G56970; *CKX5*, AT1G75450; and *SHR*, AT4G37650.

LOG4-based single-cell co-expression analysis. We identified potential co-expressed genes of *LOG4* by computing the Pearson correlation coefficient on the log-normalized UMI counts from a previously published scRNA-seq dataset¹⁸. Results were next sorted from high to low correlation and the top 50 genes were extracted for further analysis.

Constructions and transformation. All the constructs were generated on the basis of the Multisite Gateway system (Invitrogen)³¹. To generate transcriptional reporters, up to 4 kb promoters amplified using Col-0 genomic DNA as template were cloned into pDONR4P1R entry vectors by BP reaction (see detailed information in Supplementary Data 3). The entry vectors containing the promoter regions were next recombined into the binary vector pH7m34GW along with pDONR221-nYFP and pDONR2P3-GUS. Protein fusion reporters were obtained by cloning the respective promoter fragments into pDONR4P1R, the respective coding sequence without stop codon into the entry clone pDONR221, and a fluorescence tag (GFP or YFP) into pH7m34GW destination vector using Gateway cloning. To generate the luciferase (LUC) expression vector, coding sequences of the transcription factors TMO5 and LHW were cloned into the pDONR221 entry vector using the BP clonase system. The CaMV35S constitutive promoter (35S) fused to these transcription factors were generated via an LR clonase reaction with the pH7FWG2.0 vector. Promoter fragments of *LOG4* were

cloned into pDONR P4P1R. A promoter fragment fused to the luciferase reporter gene was developed by the two-fragment gateway system with pDONR221-LUC and pH7m24GW3. All constructs were verified by Sanger sequencing and were transformed into wild-type Col-0 or the dGR¹³ *Arabidopsis* plants using floral dipping. All primer sequences used for cloning and sequencing are listed in Supplementary Data 3. The resulting constructs were transformed into *Agrobacterium tumefaciens* strain GV3101.

BGLU44 CRISPR mutant generation. Four guide RNAs (gRNAs) were designed for targeting BGLU44 using the CRISPR-P tool⁴². Cloning of gRNA vectors was performed essentially as previously described¹³. Oligos were annealed and ligated into six Golden Gate gRNA entry modules using standard DNA ligation and the sequences verified. The gRNA entry plasmids were cloned into the pFASTRK_AtCas9_AG destination vector (<https://gateway.psb.ugent.be>) using Golden Gate assembly. Plants were transformed with *Agrobacterium* cultures using floral dip. A positive transgene was selected by red seed fluorescence. Deletion mutations in the target gene were confirmed in the T1 generation by PCR amplification and sequencing. T3 Cas9-free, homozygous knockout plants were selected and used for phenotype analysis. To generate *bglu44 log4* or *bglu44 log3 log4* triple mutants, homozygous *bglu44* mutants were crossed with *log4* or *log3 log4* mutants and homozygous seeds were identified by PCR. Primers used for PCR and sequencing can be found in Supplementary Data 3, *log3* and *log4* mutants identification primers were used as previously described¹³.

Imaging and processing. Xylem differentiation was observed using differential interference contrast (DIC) microscopy by mounting samples in a solution of 20% glycerol and 60% lactic acid, and imaged using an Olympus BX53 microscope equipped with DIC optics⁴. GUS staining of the *pBGLU44::nYFP-GUS* was performed using GUS staining methods⁴³. Cell wall staining for optical confocal cross sections was done using the ClearSee protocol including cell wall staining with 0.1% Calcofluor White⁴⁴. Confocal laser scanning microscopy was performed using a Leica SP8X device with 40X water-corrected objectives (NA 1.2) using an argon laser excitation wavelength of 405 nm for Calcofluor White, 488 nm for GFP, 514 nm for YFP and 561 nm for nuclear tdTomato (tdT)- and propidium iodide (PI)-stained samples. Calcofluor White, GFP, YFP, tdT and PI were visualized at an emission of 425–475 nm, 500–535 nm, 520–550 nm, 580–630 nm and 600–700 nm, respectively. Images were then processed using Leica Application Suit X. All optical cross sections were made in the middle of the meristem, determined by the quiescent center on one end and the first elongating cortex cell on the other end. The vascular cell file numbers were also counted in this position.

RNA extraction and qRT-PCR analysis. Total RNAs were extracted from 7-day-old seedling roots using Trizol reagent (Invitrogen) with RNeasy Kit (QIAGEN) as previously described⁴³ and reversely transcribed to first-strand complementary DNA by qScript cDNA SuperMix (QUANTABIO). qRT-PCR analysis was performed with SYBR GREEN I Master kit (Roche) and data were collected on a LightCycler480 apparatus (Roche) detection system in accordance with the manufacturer's instructions. Data analysis was performed with Q-BASE+ and transcript levels were normalized to EEF1a4 and CDKA1;1. All qRT-PCR primers used in this study are listed in Supplementary Data 3.

Transient expression in *Nicotiana benthamiana* and LUC assay. *A. tumefaciens* strains GV3101 carrying expression constructs (p35S::TMO5, p35S::LHW and pSHR::LUC) were grown in YEP media overnight, centrifuged and resuspended in infiltration buffer (10 mM MES, 10 mM MgCl₂ and 200 μM acetosyringone) at OD₆₀₀ 1.0. Equal volumes of cultures of different constructs were mixed for infiltration. To prevent gene silencing, p19 agrobacterial culture was mixed with the combinations. Cultures were then spot-infiltrated into 4-week-old *N. benthamiana* (tobacco) leaves. After 3 d, leaves were cut and sprayed with 1 mM luciferin, which were then imaged for an 11 min exposure using a NightSHADE LB 985 system (Berthold Technologies).

Chromatin immunoprecipitation (ChIP). ChIP was performed using universal plant ChIP kit (C01010152, Diagenode). One gram roots of 7-day-old dGR and Col-0 seedlings transferred to MS medium with DEX for 24 h were used. Anti-GR antibody (PA1-516, Invitrogen) was bound to Protein A Dynabeads (Diagenode) and incubated with chromatin sample overnight at 4 °C. Quantitative PCR reactions of the samples were performed using SYBR GREEN I Master kit (Roche), with the following PCR conditions: 3 min at 95 °C, followed by 45 cycles of 30 s at 95 °C, 30 s at 58 °C and 30 s at 72 °C. Primers used in this study are listed in Supplementary Data 3, SHR ChIP primers were used as previously described⁴⁶. Data were analysed by fold enrichment method (fold enrichment = 2^{-(Ct(Sample)-Ct(Control))}). Two independent biological replicates were performed.

Protein expression and purification. Mature BGLU44 without signal peptide (AA 24–512) was picked up from a cDNA library, with primers listed in Supplementary Data 3. The PCR fragment was cloned between the SphI and XhoI sites in the linearized ppSA-Ht plasmid (a derivative of the pPICZalphaB vector; provided by the VIB Protein Core) using the Gibson assembly protocol

from New England Biolabs. The gene of interest was cloned in the frame of the alpha mating factor for secreted expression. The obtained plasmid was then Pmel-linearized and transformed into NRRL Y-11430 wild-type *Komagataella phaffii* yeast expression strain. Transformed cells where plated on increasing zeocin concentrations (100 μg ml⁻¹, 250 μg ml⁻¹, 500 μg ml⁻¹ and 1,000 μg ml⁻¹) to screen for high-expressing colonies.

Purification was performed according to the method described in ref. ¹⁹. Briefly, biomass was grown by inoculating 250 ml buffered glycerol-complex medium with NRRL Y-11430(ppSA-HT_BGLU44) and incubating for 48 h at 30 °C and 200 r.p.m. Cells where collected by centrifuging for 10 min at 1,500 g and resuspended in buffered methanol-complex medium containing 1% MeOH to induce protein expression. Resuspended cultures were grown for 48 h at 30 °C and 200 r.p.m. and spiked with 1% MeOH every 12 h. Cells where pelleted by centrifuging the culture for 10 min at 1,500 g. Supernatant was collected and further processed. Diafiltration was performed to transfer the supernatant to 20 mM sodium acetate buffer (pH 5.5). The desalting supernatant was applied to a 5 ml Source 15S cation exchange column (Cytiva) pre-equilibrated with 20 mM sodium acetate buffer (pH 5.5). The column was washed with 20 column volumes of 20 mM sodium acetate buffer (pH 5.5) containing 200 mM NaCl. Protein was eluted with a linear gradient of 20–60% over 10 column volumes with 20 mM sodium acetate buffer supplemented with 1 M NaCl (pH 5.5). Fractions with proteins running on the correct molecular weight on an SDS-page gel were then collected and pooled together, and briefly concentrated to 4 ml with an Amicon ultra concentrator with 30 kDa cut-off (Millipore). The concentrate was then loaded and run on a Superdex 200 prep grade column (Cytiva) pre-equilibrated with 20 mM MES (pH 6.0). Fractions containing protein of the correct size were pooled and the concentration was determined using a Trinean dropsense 16 device (Unchained Labs).

BGLU44 enzyme activity (D-glucose method). To determine hydrolase activity towards all cytokinin glycoside substrates, 10 μg BGLU44 was incubated for 2 h at 30 °C in 50 μl assays that contained 200 μg substrate and 100 mM Na₂HPO₄/citric acid buffer (pH 4.5). After terminating the reaction by boiling for 2 min, liberated glucose was determined by the D-glucose assay kit (GOPOD Format, Megazyme). D(+)-cellobioside was used as a positive control. All assays were performed in quadruplicate. All cytokinin glycoside substrates used (trans-zeatin-O-glucoside riboside, tZROG; trans-zeatin-7-glucoside, tZ7G; trans-zeatin-9-glucoside, tZ9G; kinetin-3-glucoside, K3G; kinetin-9-glucoside, K9G; N6-isopentenyladenine-7-glucoside, iP7G; and N6-isopentenyladenine-9-glucoside, iP9G) were obtained from OlChemIm (Olomouc). Stock solutions were prepared as 50 μg μl⁻¹ in dimethylsulfoxide (DMSO; Sigma-Aldrich).

BGLU44 enzymatic assay (MS/MS method). Enzymatic reaction was carried out by incubating 5 μM purified BGLU44 with 10 μM tZOG, tZROG, cZOG or cZROG in Na₂HPO₄/citric acid buffer (pH 4.5) at 30 °C for 30 min, 1 h, 6 h, 12 h and 24 h. Enzymatic reactions were repeated in triplicate and stopped with equal volume of 100% MeOH. The supernatant fractions (2 μl per sample) were then transferred in insert-equipped vials and diluted in 5% (v/v) methanol up to 40 μl. Samples (1 μl) were further injected onto a reversed-phase column (Acquity UPLC BEH Shield RP18 column; 150 × 2.1 mm, 1.7 μm; Waters). Separation was performed on an Acquity UPLC System (Waters). The effluent was introduced into the electrospray ion source of a triple quadrupole mass spectrometer Xevo TQ-S MS (Waters) and analysed according to the UHPLC-ESI-MS/MS method for cytokinin metabolic profiling as described in ref. ⁴⁷.

Measurements of endogenous cytokinins. Quantification of cytokinin metabolites was performed according to the method described by ref. ⁴⁷. Samples (50 pieces of 1 cm root tips) were homogenized and extracted in 0.5 ml modified Bielecki buffer (60% MeOH, 10% HCOOH and 30% H₂O) together with a cocktail of stable isotope-labelled internal standards (0.25 pmol of cytokinin bases, ribosides, N⁶-glucosides and 0.5 pmol of cytokinin O-glucoside nucleotides per sample added). The extracts were purified using the in-tip microSPE on the basis of the StageTips technology⁴⁸. Briefly, combined multi-StageTips (containing C18/SDB-RPSS/cation-SR layers) were activated sequentially with 50 μl each of acetone, methanol, water, 50% (v/v) nitric acid and water (by centrifugation at 434 g, 15 min, 4 °C). After application of the sample (500 μl, 678 g, 30 min, 4 °C), the microcolumns were washed sequentially with 50 μl of water and methanol (525 g, 20 min, 4 °C), and elution of samples was performed with 50 μl of 0.5 M NH₄OH in 60% (v/v) methanol (525 g, 20 min, 4 °C). The eluates were then evaporated to dryness in vacuo and stored at -20 °C. Cytokinin levels were determined using ultra high performance liquid chromatography-electrospray tandem mass spectrometry (UHPLC-MS/MS). Quantification was performed by Masslynx software using a standard isotope dilution method⁴⁹. Five independent biological replicates were performed.

scRNA-seq: protoplasting conditions and FACS. Briefly, 6-day-old seedlings were transferred to 10 μM BAP-supplemented medium or an equal volume of DMSO as mock treatment and allowed to grow for 2 h. The root tips were then cut and incubated in protoplasting Solution B (1.5% (wt/vol) cellulysin and 0.1% (wt/vol) pectolyase in Solution A (600 mM mannitol, 2 mM MgCl₂, 0.1% (wt/vol) bovine

serum albumin, 2 mM CaCl₂, 2 mM MES, 10 mM KCl, pH 5.5)) for approximately 1 h at room temperature. Cells were filtered through a 70 µm cell strainer and spun down at 200 g for 6 min, resuspended in Solution A, filtered through a 40 µm cell strainer and stained for live/dead using 4',6-diamidino-2-phenylindole (DAPI) at 14 µM final concentration. Cells were sorted on a BD Aria II and protoplasts without the DAPI signal were selected for further analysis.

scRNA-seq: 10X genomics sample preparation, library construction and sequencing. Sorted cells were centrifuged at 400 g at 4 °C and resuspended in Solution A to yield an estimated concentration of 1,000 cells per µl. Cellular suspensions were loaded on a Chromium Single Cell 3' GEM, Library & Gel Bead Kit (V3 chemistry, 10X Genomics) according to the manufacturer's instructions. Sequencing libraries were loaded on an Illumina HiSeq4000 and sequenced following recommendations of 10X genomics at the VIB Nucleomics Core (VIB, Leuven).

scRNA-seq: raw data processing and generation of gene expression matrix. Demultiplexing of the raw sequencing data was done by the 10x CellRanger (version 3.1.0) software 'cellranger mkfastq', which wraps Illumina's bcl2fastq. The fastq files obtained after demultiplexing were used as input for 'cellranger count', which aligns the reads to the *Arabidopsis thaliana* reference genome (Ensemble TAIR10.40) using STAR and collapses them to UMI counts. The result is a large digital expression matrix, with cell barcodes as rows and gene identities as columns. Initial filtering in CellRanger recovered 11,569 cells for the mock-treated sample and 9,799 cells for the cytokinin-treated sample. To ensure that we only used high-quality cells for further analysis, we used the filtered data provided by cellranger. This corresponds to 26,756 mean reads per cell in the mock-treated sample and 32,885 mean reads per cell in the cytokinin-treated sample.

scRNA-seq: data analysis (clustering, identity assignment, differential gene expression and quality control). All analyses were performed in R (version 3.6.0). Pre-processing of the data was done by the scater package (version 1.10.1) according to the workflow proposed by the Marioni lab⁵⁰. Outlier cells were identified on the basis of 2 metrics (library size and number of expressed genes) and were tagged as outliers when they were 4 median absolute deviations away from the median value of these metrics across all cells. Normalizing the raw counts, detecting highly variable genes, finding clusters and creating t-distributed stochastic neighbour embedding (t-SNE) plots were done using the Seurat pipeline (version 3.2.3). Differential expression analysis for marker gene identification per subpopulation was based on the non-parametric Wilcoxon rank sum test implemented within the Seurat pipeline. Clusters with the same cell annotation based on gene expression analysis were combined to generate a more comprehensible dataset. Potential doublets were identified using the DoubletFinder algorithm (version 2.0.0)⁵¹. The number of high-confidence doublets was below 1% (113 out of 11,313 cells for the mock-treated sample and 96 out of 9,640 cells for the cytokinin-treated sample). Relative proportions of cell identities for both the mock- and cytokinin-treated samples were comparable (Supplementary Fig. 3d) to that of a previously published root meristem scRNA-seq atlas¹⁸.

scRNA-seq: gene selection, cloning and plant transformation. Genes were selected on the basis of their cell type-specific differential expression between treatments. Transcriptional reporter lines were obtained by amplifying the upstream region of the transcriptional start and cloning this into the pGGA000 entry vector. The final expression clones were generated by cloning the promoter regions, three fluorescent tags (GFP), a HEAT SHOCK PROTEIN 18.2 (HSP18.2) terminator and a linker into the pFASTR-AG destination vector using Golden Gate cloning. All constructs were verified by Sanger sequencing and transformed into wild type using simplified floral dipping⁵². All primer sequences used for cloning and sequencing can be found in Supplementary Data 3.

Quantification and statistical analysis. All microscopic images were quantified using ImageJ and final figures were composed using Adobe Illustrator. For all expression pattern analyses, at least five independent transgenic lines showed the same expression pattern. For the expression pattern analysis and changes upon DEX treatments, at least three biological repeats with individual lines show similar results. For TCSn quantification, a rectangular area was selected from the quiescent center to the top of the image comprising the pericycle and all cells inwards. The pixel intensities from the entire selected area were analysed on root meristems of 7-day-old seedlings. Confocal images were all made through the quiescent center to ensure that the same position in the root was analysed in all images. Over 20 individual roots were imaged for each line. 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 ($P < 0.001$).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Upon acceptance, the scRNA-seq data will be made accessible via an on-line browser tool (<http://bioit3.irc.ugent.be/plant-sc-atlas/>) and raw data can be accessed at NCBI with GEO number GSE179820. All other data are either in the main paper or the Supplementary Material. Source data are provided with this paper. Requests should be directed to the corresponding authors.

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Author contributions

B.D.R. and B.Y. conceived the project and designed experiments; F.B., L.P., I.P., K.H. and O.N. performed enzymatic assays and cytokinin measurements; J.N. produced and purified BGLU44 protein with the help of J.H.; M.M., K.V., T.E. and Y.S. analysed single-cell data; E.F. and A.B. performed the mathematical modelling; B.Y., M.M., Y.S., W.S. and J.R.W. performed all other experiments; B.D.R. supervised the project; B.Y. and B.D.R. wrote the paper with input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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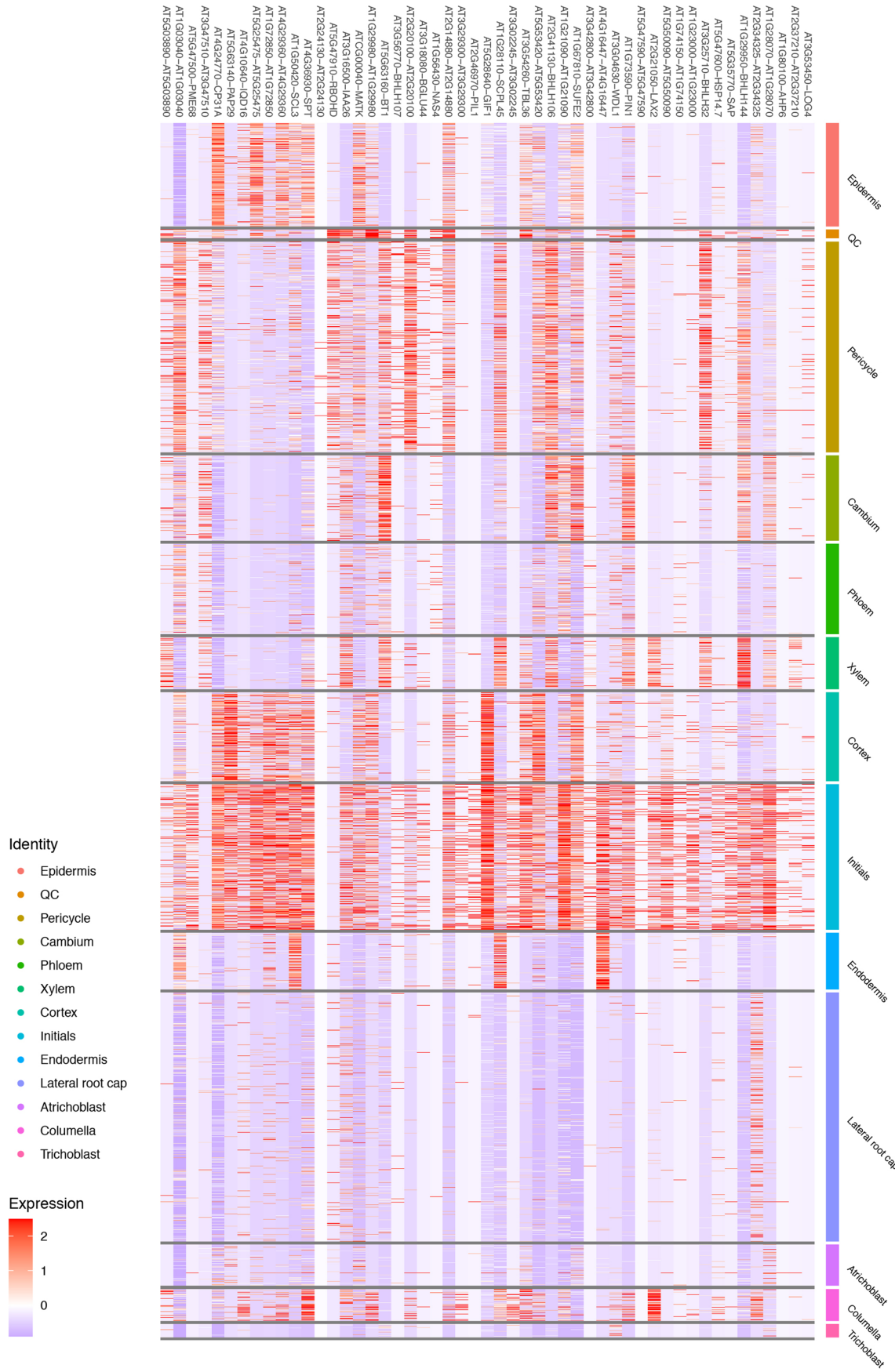
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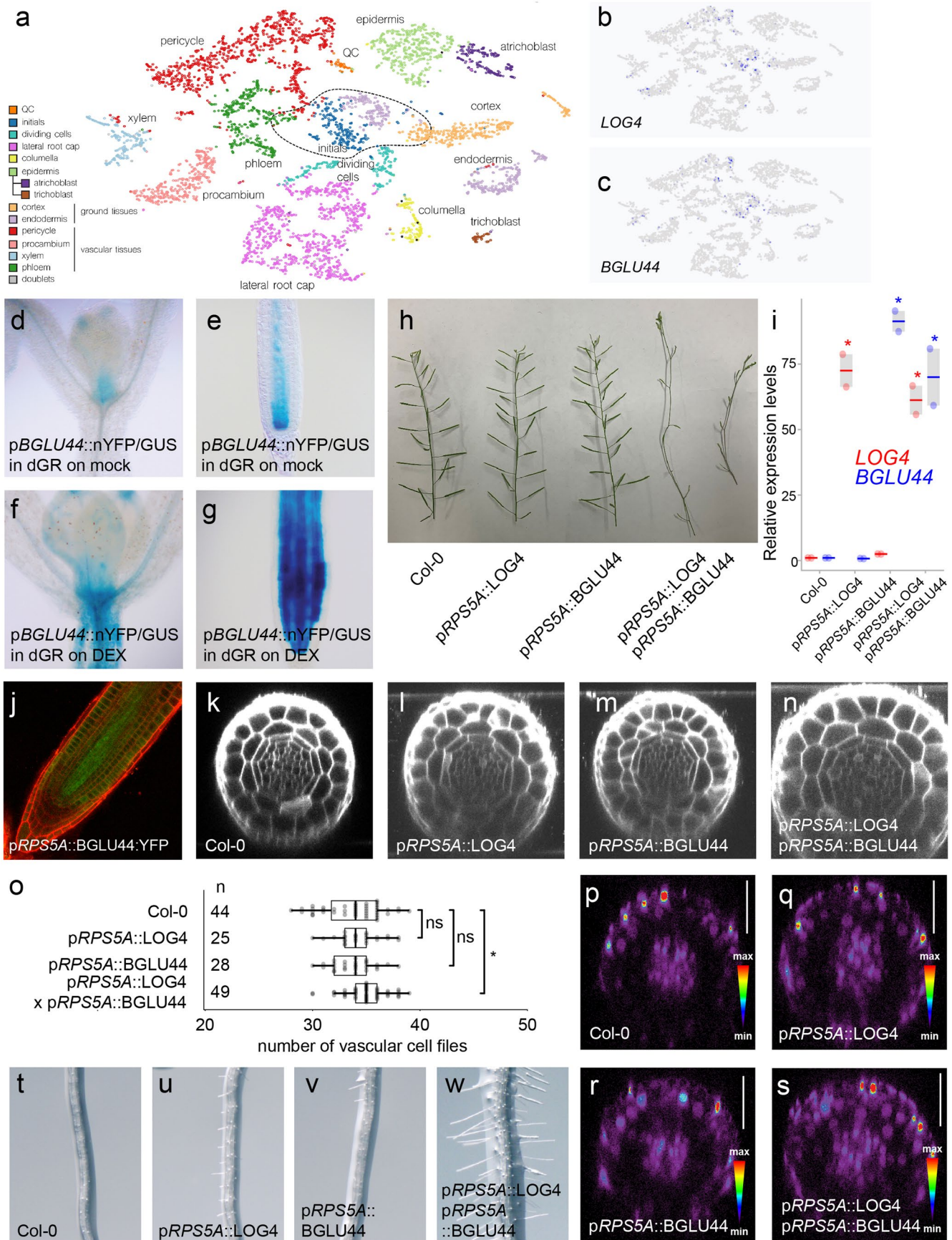
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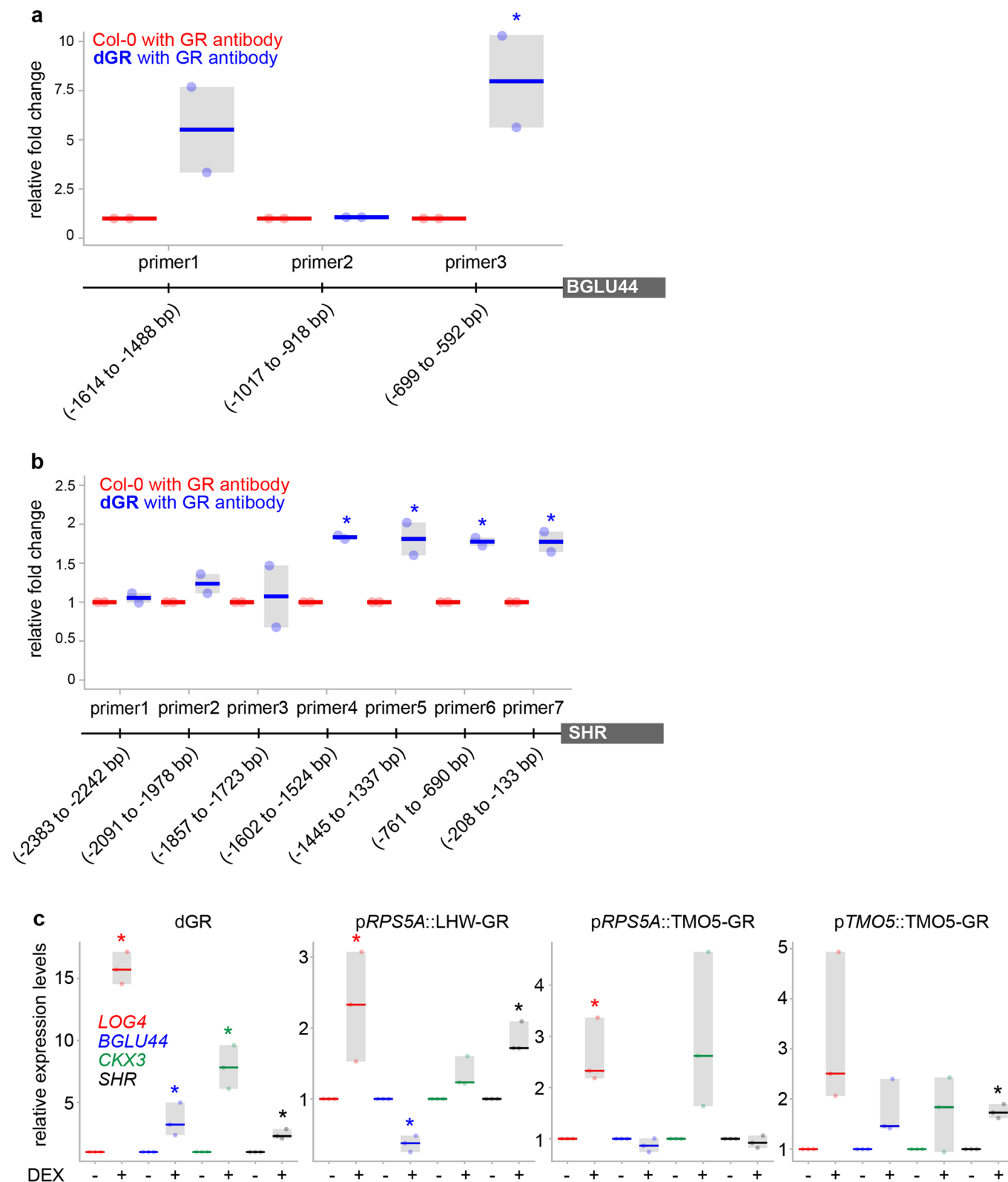


Extended Data Fig. 1 | LOG4 single cell co-expression analysis. Top 50 genes co-expressed with LOG4 according to a single cell atlas of the Arabidopsis root meristem. Expression values are shown for each of the separate tissue types.

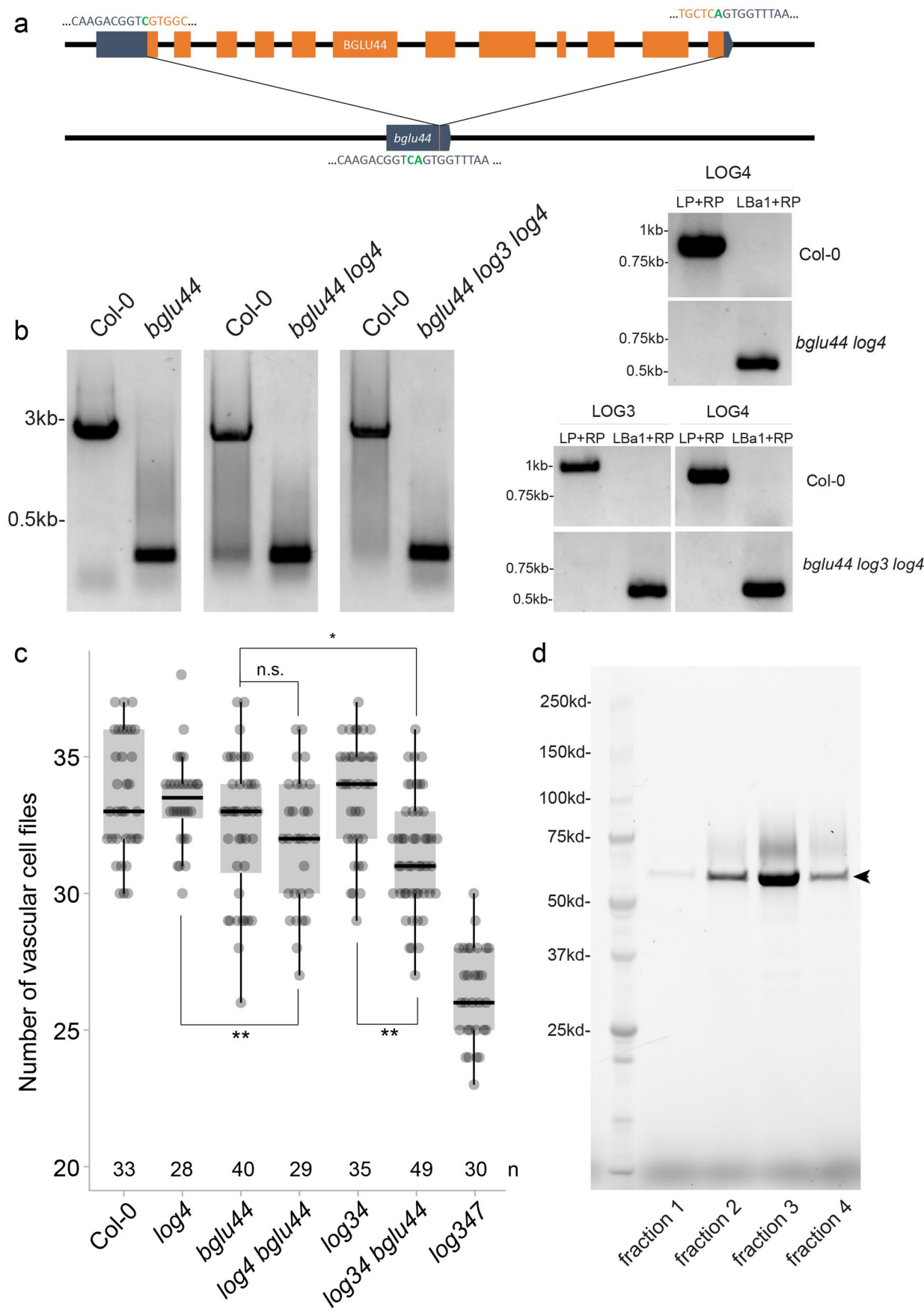


Extended Data Fig. 2 | See next page for caption.

Extended Data Fig. 2 | Characterization of *BGLU44* expression and function. **a-c**, Predicted expression of *LOG4* and *BGLU44* according to a single cell atlas of the Arabidopsis root meristem. Note the predicted co-expression in the feature plots for both genes. **d-g**, Expression of p*BGLU44*::nYFP/GUS in the dGR shoot and root meristems grown on mock medium and transferred to mock or 10 μM DEX for 24h. **h**, Inflorescence stem and siliques phenotypes in the indicated genotypes. **i**, Relative expression levels as determined by RT-qPCR of *BGLU44* and *LOG4* in the indicated lines. Boxes represent the 1st and 3rd quartile and the center line represents the median. Asterisks in graphs indicate significance values as determined by standard two-sided t-tests. Black lines indicates mean values and grey boxes indicate data ranges. **j**, Expression of p*RPSSA*::*BGLU44*:YFP in the root meristem grown on mock medium showing that the *BGLU44*:YFP fusion protein is expressed. **k-o**, Phenotypes and quantification of the number of vascular cell files in the mentioned genotypes. Boxes represent the 1st and 3rd quartile and the center line represents the median. Statistics in n are performed using pairwise comparisons using a standard two sided t-test ($p\text{-value}<0,05$). **p-s**, Confocal images of optical cross sections of root meristems expressing p*TCSn*::ntdTomato reporter in the mentioned genotypes. Scale bars are 10 μm. **t-w**, Root hair phenotypes of the mentioned genotypes. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Supplementary Data 4. The experiments in j was repeated three times independently, with similar results.

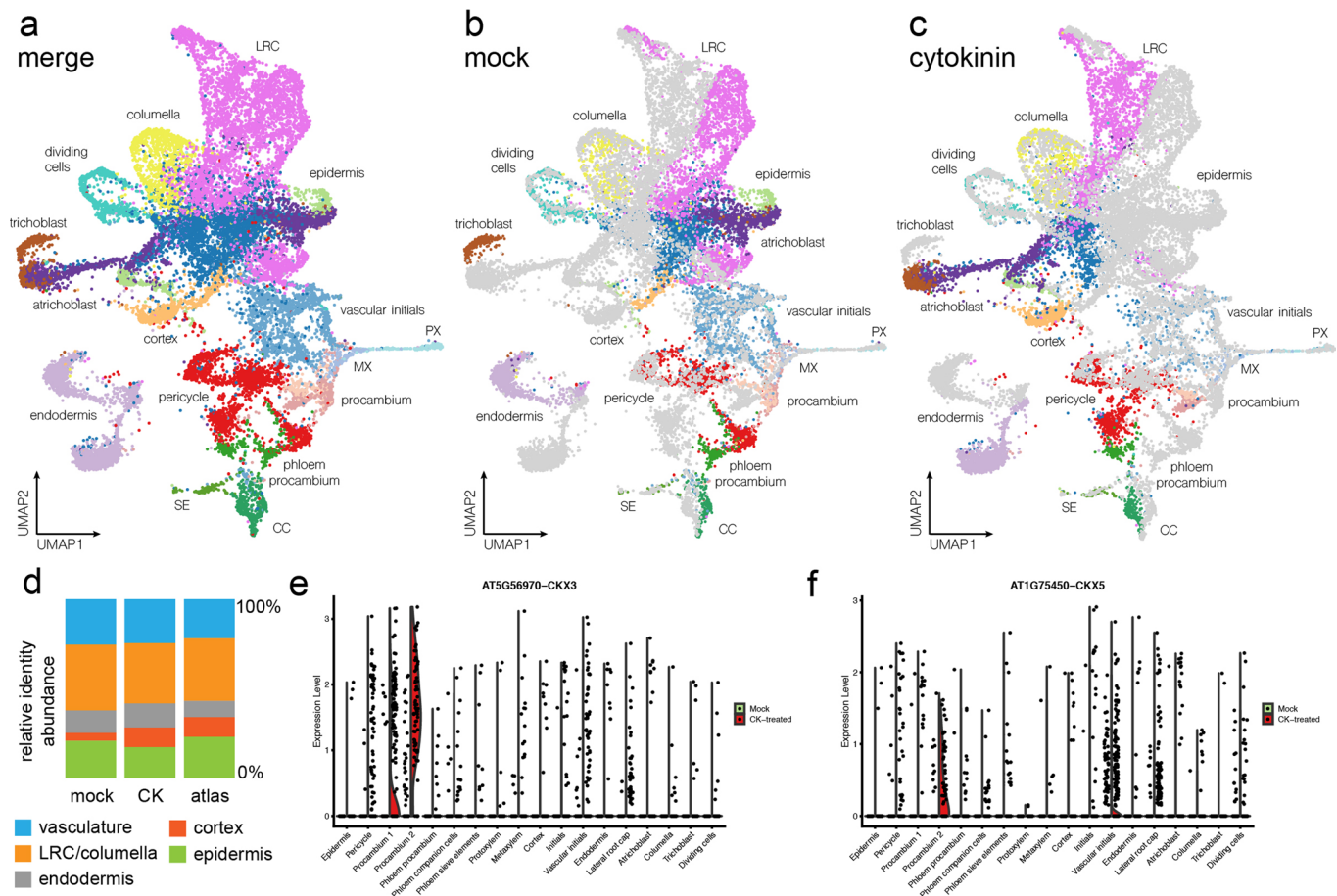


Extended Data Fig. 3 | BGLU44 and SHR are direct TMO5/LHW target genes. **a, b**, ChIP-RT-qPCR experiments using anti-GR antibody on Col-0 and dGR lines. Graphs show the values of two biological repeats; black lines indicates mean values and grey boxes indicate data ranges. n = 2 biological replicates. Asterisks indicate significance values as determined by standard two-sided t-tests. **c**, Relative expression levels as determined by RT-qPCR of *LOG4*, *BGLU44*, *CKX3* and *SHR* in the indicated lines in presence or absence of 10 μM DEX. n = 3 biological replicates. Asterisks in graphs indicate significance values as determined by standard two-sided t-tests. Black lines indicates mean values and grey boxes indicate data ranges. All Boxes represent the 1st and 3rd quartile and the center line represents the median. In all panels, n represents the.

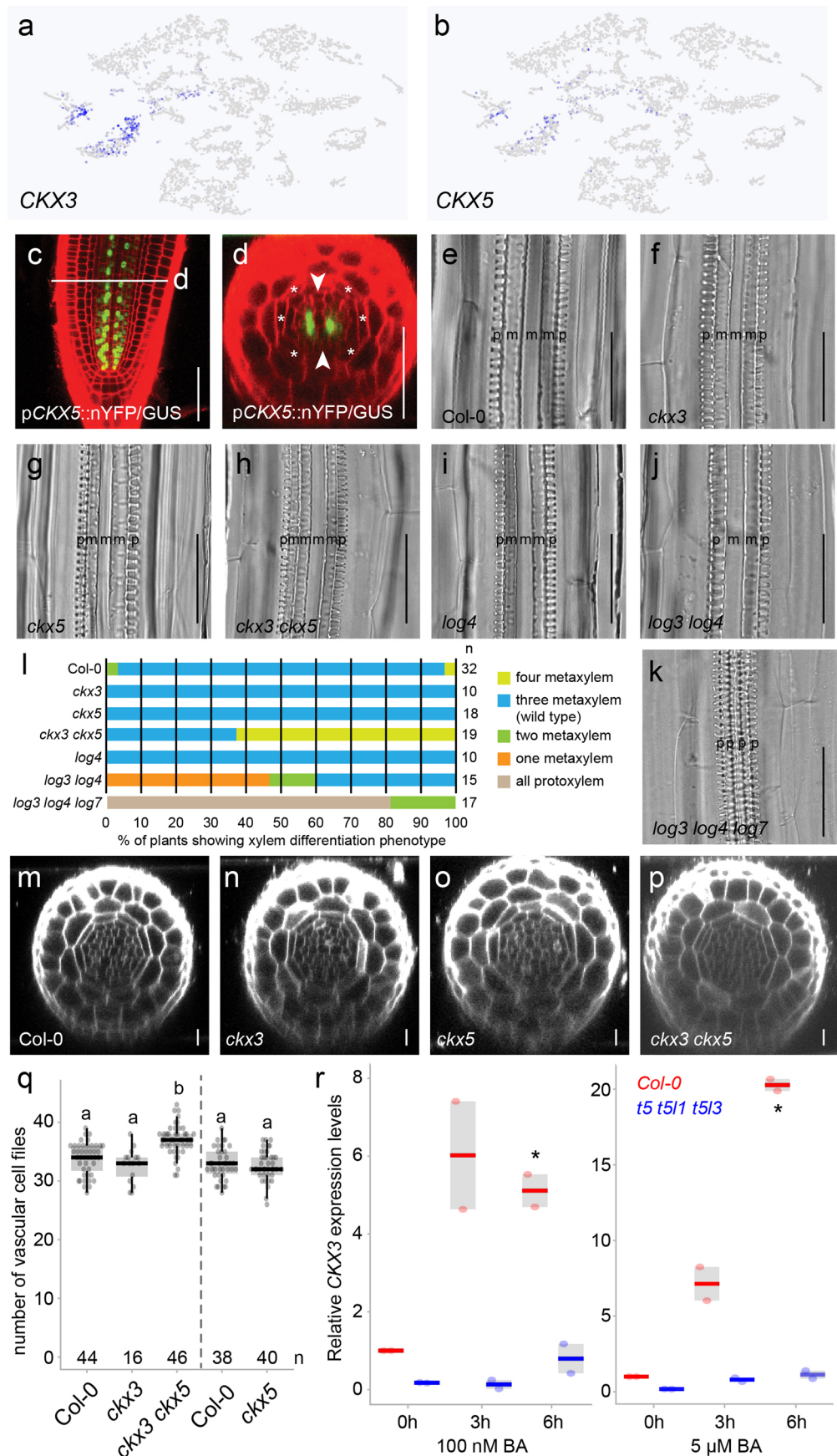


Extended Data Fig. 4 | See next page for caption.

Extended Data Fig. 4 | BGLU44 mutant generation and protein purification. **a**, Position of the deletion in the *bglu44* mutant line. **b**, PCR validation of the *bglu44* mutant and crosses with *log4* and *log34* mutants. **c**, quantification of the number of vascular cell files in the indicated lines. Boxes represent the 1st and 3rd quartile and the center line represents the median. n indicate independent plants, the data and statistics are summarized in Supplementary Data 4. Asterisks in graphs indicate significance values as determined by standard two-sided t-tests (*:p-value<0.05; **: p-value<0.01). **d**, Coomassie gel showing subsequent fractions of purified BGLU44 protein (arrowhead) after ion exchange chromatography. The experiments in b and d were repeated three times independently, with similar results.

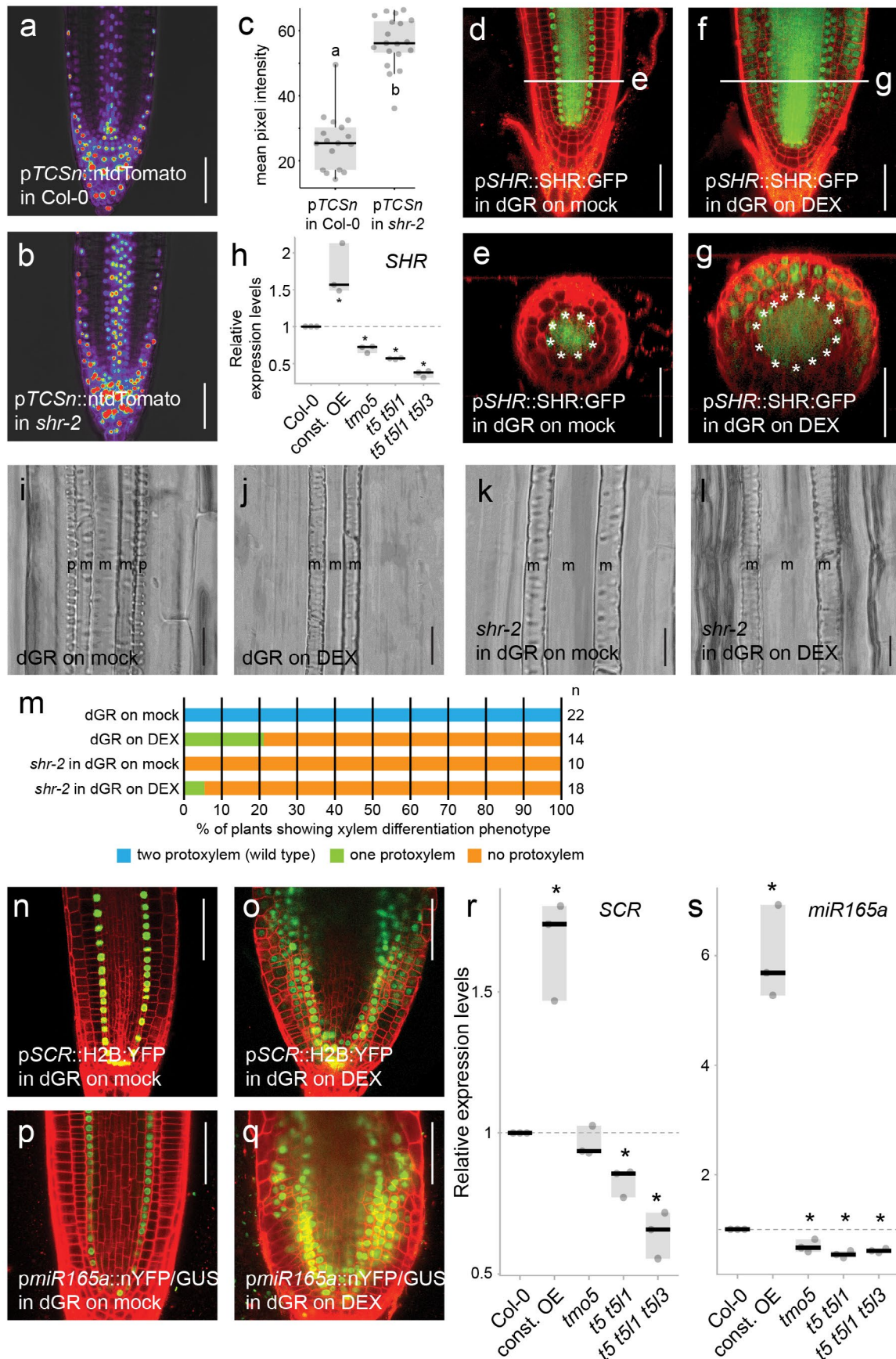


Extended Data Fig. 5 | Single cell transcriptional changes in root meristem cells upon cytokinin treatment. **a-c.** UMAP representation of the merged (a), mock (b) and cytokinin (c) datasets in which cell identities are shown by clusters of different colours. In panels b-c, the cells from the other dataset are indicated in grey to show contribution to each of the cell clusters. LRC: lateral root cap; PX: protoxylem; MX: metaxylem; SE: sieve element; CC: companion cell. The most central dark blue cell cluster is the initial cell cluster. **d.** Relative cell identity abundance in the mock and cytokinin (CK) datasets in comparison to a previously published root meristem atlas⁸. **e,f.** Violin plots showing normalized gene expression of CKX3 (e) and CKX5 (f) by cell type and split by treatment: left side of the violin is mock dataset in green; right side of the violin is CK-treated dataset in red).



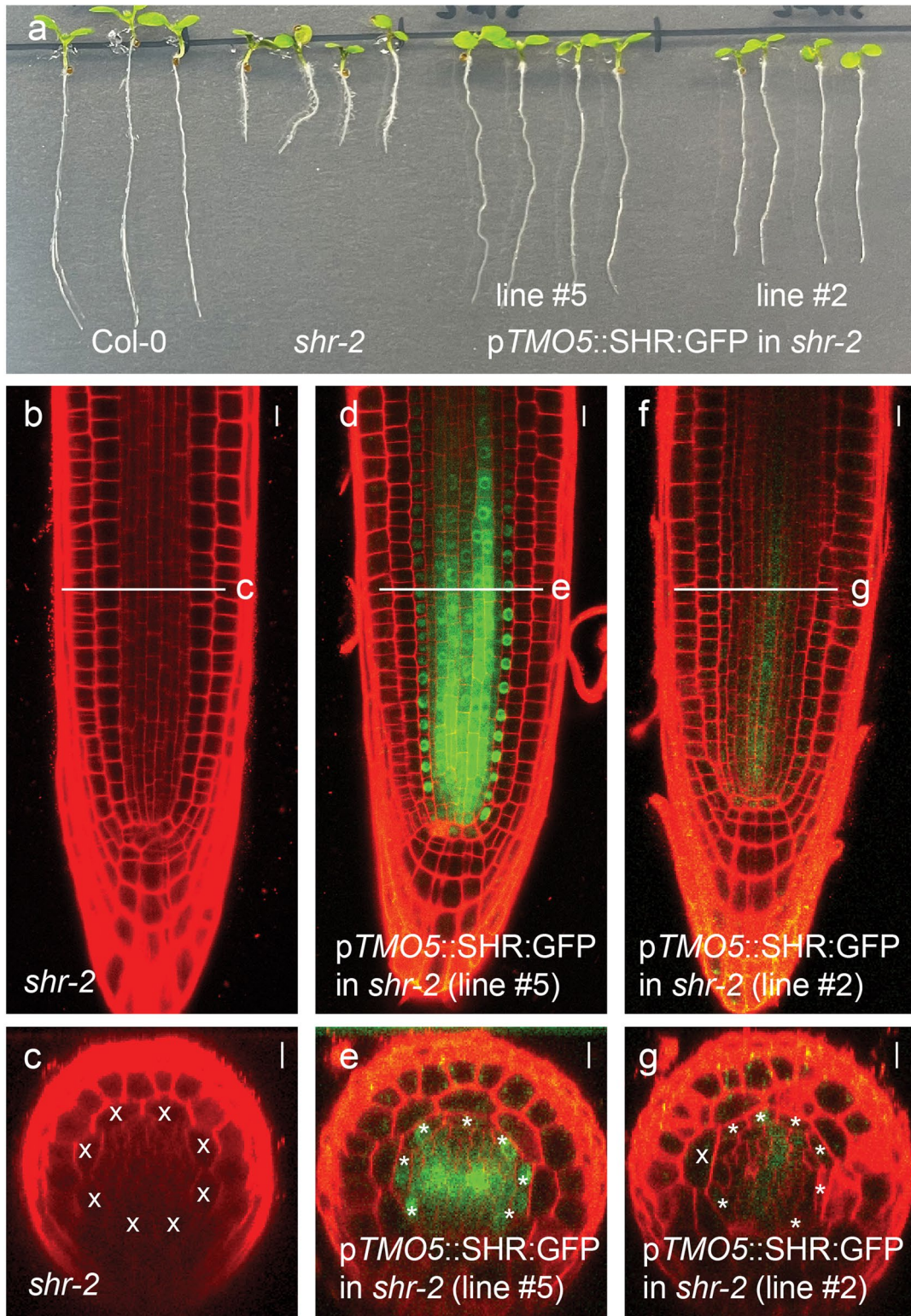
Extended Data Fig. 6 | See next page for caption.

Extended Data Fig. 6 | CKX3 and CKX5 are required for normal vascular development. **a,b**, Predicted expression of CKX3 and CKX5 according to a previously published single cell atlas of the Arabidopsis root meristem¹⁸. Compare to Supplementary Fig. 2a for cell identities. **c,d**, Expression of pCKX5::nYFP/GUS in the root meristem grown on mock medium. **e-k**, Microscopic images of xylem differentiation in the mentioned genotypes. **p**: protoxylem; **m**: metaxylem. **l**, Quantification of the different classes of xylem phenotypes shown in panels e-k. **m-p**, Confocal microscopy images of cross sections through the root meristem showing the vascular cell file number phenotype and quantification of the mentioned genotypes. **q**, Quantification of the number of vascular cell files shown in panels m-p. Lower-case letters on top of the boxplots indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($p<0.001$). Boxes represent the 1st and 3rd quartile and the center line represents the median. Grey dotted line separates data from different experiments. **r**, RT-qPCR analysis showing the relative expression levels of CKX3 on wild type Col-0 and *tmo5* triple mutant backgrounds when grown on mock (0h) or 100 nM or 5 μM BA for the indicated time. Boxes represent the 1st and 3rd quartile and the center line represents the median. $n = 3$ biological replicates. Asterisks in graphs indicate significance values as determined by standard two-sided t-tests. Black lines indicate mean values and grey boxes indicate data ranges. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Supplementary Data 4. The experiments in c-d were repeated three times independently, with similar results.

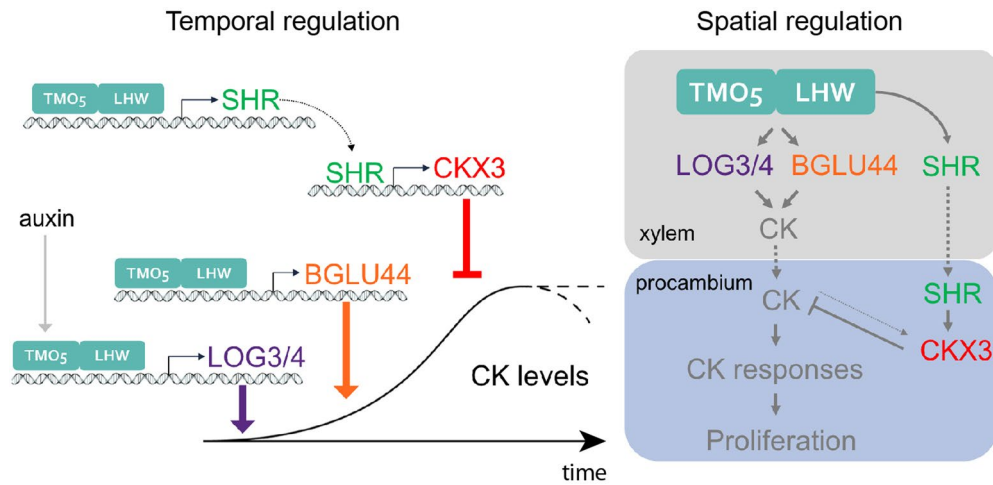


Extended Data Fig. 7 | See next page for caption.

Extended Data Fig. 7 | SHR acts downstream of TMO5/LHW. **a,b**, Confocal images of root meristems expressing pTCSn::ntdTomato reporter in Col-0 and *shr-2* backgrounds. **c**, Quantification of the pTCSn::ntdTomato mean pixel intensity in the mentioned genotypes. Boxes represent the 1st and 3rd quartile and the center line represents the median. n indicate independent plants. Lower-case letters on top of the boxplots indicate significantly different groups as determined by one-way ANOVA with post-hoc Tukey HSD testing ($p < 0.001$). **d-g**, Expression of pSHR::SHR:GFP in the dGR root meristem grown on mock medium and transferred to mock or 10 μ M DEX for 24 h. **h**, Relative expression levels of *SHR* in wild type (Col-0), TMO5/LHW misexpression, and *tmo5*, *tmo5like1* (*t5 t5l1*) and *tmo5 tmo5like1 tmo5like3* (*t5 t5l1 t5l3*) mutant backgrounds. Asterisks indicate significance values as determined by standard two-sided t-tests. Boxes represent the 1st and 3rd quartile and the center line represents the median. n = 3 biological replicates. Black lines indicates mean values and grey boxes indicate data ranges. **i-l**, Microscopic images of xylem differentiation in the mentioned genotypes. p: protoxylem, m: metaxylem. **m**, Quantification of the different classes of xylem phenotypes shown in panels i-l. **n-o**, Expression of pSCR::H2B:nYFP in the dGR root meristem grown on mock medium and transferred to mock or 10 μ M DEX for 24h. **p-q**, Expression of *pmiR165a::nYFP/GUS* in the dGR root meristem grown on mock medium and transferred to mock or 10 μ M DEX for 24h. **r-s**, Relative expression levels of *SCR* and *miR165a* in wild type (Col-0), TMO5/LHW misexpression, and *tmo5*, *tmo5like1* (*t5 t5l1*) and *tmo5 tmo5like1 tmo5like3* (*t5 t5l1 t5l3*) mutant backgrounds. Boxes represent the 1st and 3rd quartile and the center line represents the median. n = 3 biological replicates. Asterisks indicate significance values as determined by standard two-sided t-tests. Black lines indicates mean values and grey boxes indicate data ranges. In panels a-b, d-g, i-l and n-q, scale bars are 50 μ m. In all panels, n represents the number of replicates or data points; all data and statistics are summarized in Supplementary Data 4. The experiments in d-g, n-q were repeated three times independently, with similar results.



Extended Data Fig. 8 | Xylem expressed SHR is sufficient to rescue the *shr-2* mutant. **a**, Seedling phenotypes of the lines indicated. Note the rescue of root length in the pTMO5::SHR:GFP lines in *shr-2* mutant background. **b-g**, Confocal images of root meristems of the indicated lines counterstained with PI (red). Scale bars are 10 μ m. Panels c, e and g are optical cross sections of the images in b, d and f, respectively. Asterisks indicate endodermal cells; x marks mixed ground tissue identity. The experiments here were repeated three times independently, with similar results.



Extended Data Fig. 9 | A model for spatiotemporal control of cytokinin levels. Auxin induces *TMO5* expression and as such triggers activation of the *TMO5/LHW* heterodimer complex. Sequentially, *LOG3/4*, *BGLU44* and *SHR* expression are induced. *SHR* itself binds to the promoter region of *CKX3* to activate its expression. As such, the same transcription factor complex sequentially increases levels of active cytokinin via *LOG3/4* and *BGLU44*; and represses levels active cytokinin via *SHR-CKX3*. The combined activity balances levels of active cytokinin to allow normal vascular proliferation and differentiation. The left side of the figures shows the temporal aspect, while the right side shows the spatial aspect.

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Software and code

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Data collection	Olympus BX53 microscope, Leica SP8, LightCycler480 apparatus (Roche), NightSHADE LB 985 system (Berthold Technologies), Acuity UPLC® System (Waters, Milford, MA, USA)
Data analysis	Demultiplexing of the raw sequencing data was done by the 10x CellRanger (version 3.1.0) software ‘cellranger mkfastq’. Reads were aligned to the <i>Arabidopsis thaliana</i> reference genome (Ensemble TAIR10.40) using ‘cellranger count’. All analyses were performed in R (version 3.6.0). Pre-processing of the data was done by the scater (version 1.10.1) package according to the workflow proposed by the Marioni lab (Lun et al., 2016). Normalizing the raw counts, detecting highly variable genes, finding clusters and creating tSNE plots was done using the Seurat pipeline (version 3.2.3). Leica Application Suit X (version 3.3.0.16799) ImageJ (version,1.53c)

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Sample size	Sample sizes were determined by previous pilot experiments to be sufficient to achieve desired outcomes. Sample sizes are indicated in the Figures, legends and main text.
Data exclusions	No data was excluded in this study. In the single-cell analysis, outlier cells were excluded when they were 4 median absolute deviations (MADs) away from the median value of either library size or number of expressed genes.
Replication	Each experiment was repeated at least twice, and only results representing the consistent outcome are reported.
Randomization	Plant samples used in this study were genetically homogeneous and of the same age with same growth condition. Plants were randomly picked from the larger pools for experiments.
Blinding	The experiments were carried out without prior knowledge of the experimental outcome. Therefore investigators were not blinded, but biological repeats were performed as described above.

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Methods

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<input checked="" type="checkbox"/>	Flow cytometry
<input checked="" type="checkbox"/>	MRI-based neuroimaging

Antibodies

Antibodies used

Anti-GR antibody (polyclonal antibody, Catalog # PA1-516, LOT WB308073 Invitrogen)

Validation

This antibody had been successfully used in Arabidopsis for GR-fusion protein ChIP assay. (Morohashi et al, 2017. <https://doi.org/10.1101/pp.107.104521>)