Root stem cell niche organizer specification by molecular convergence of PLETHORA and SCARECROW transcription factor modules

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Continuous formation of somatic tissues in plants requires functional stem cell niches where undifferentiated cells are maintained. In Arabidopsis thaliana, PLETHORA (PLT) and SCARECROW (SCR) genes are outputs of apical–basal and radial patterning systems, and both are required for root stem cell specification and maintenance. The WUSCHEL-RELATED HOMEOBOX 5 (WOX5) gene is specifically expressed in and required for functions of a small group of root stem cell organizer cells, also called the quiescent center (QC). PLT and SCR are required for QC function, and their expression overlaps in the QC; however, how they specify the organizer has remained unknown. We show that PLT and SCR genetically and physically interact with plant-specific teosinte-branched cycloidea PCNA (TCP) transcription factors to specify the stem cell niche during embryogenesis and maintain organizer cells post-embryonically. PLT–TCP–SCR complexes converge on PLT-binding sites in the WOX5 promoter to induce expression.

[Keywords: Arabidopsis; stem cell niche; organizer; quiescent center; PLT–TCP–SCR complexes; WUSCHEL-RELATED HOMEOBOX 5]

Supplemental material is available for this article.

Received March 12, 2018; revised version accepted May 31, 2018.
Risueno et al. 2015]. In addition, SCR [together with its partner protein, SHORT-ROOT (SHR)] is required for asymmetric cell division of the cortex/endodermis initial (CIE) daughter (CEID) cell in the ground tissue, which gives rise to cortex and endodermis cell lineages [Benfey et al. 1993; Scheres et al. 1994; Di Laurenzio et al. 1996; Scheres and Benfey 1999; Helariutta et al. 2000]. Furthermore, SCR is required for the regulation of QC division rate, which determines how rapidly abutting stem cells are replenished [Cruz-Ramírez et al. 2013].

The embryonic initiation of the root stem cell niche in Arabidopsis is marked by a stereotypic transverse asymmetric cell division within the hypophyseal cell during the early to mid-globular embryo stages. The smaller lens-shaped apical daughter acquires QC cell identity, whereas the basal descendant cell becomes distal columella stem cells [Jürgens et al. 1994; Scheres and Benfey 1999; Jürgens 2001; Weigel and Jürgens 2002; Ten Hove and Heidstra 2008; Ten Hove et al. 2015]. Although PLT and SCR genes are expressed in partially overlapping larger domains [of which the QC forms a subset], loss-of-function mutants of SCR and combinations of loss of function of PLT clade members lead to differentiation of the root stem cell niche and decrease the expression of different QC identity markers from embryogenesis onward [Sabati-ni et al. 2003; Aida et al. 2004; Galinha et al. 2007]. Up to now, it has not been revealed how their activities might converge for QC specification in such a narrow domain. Reported target genes of the SHR/SCR pathway [Levesque et al. 2006; Sozzani et al. 2010; Moreno-Risueno et al. 2015] and of the root-expressed PLT genes [Santuari et al. 2016] do not show overall overlap, leaving it unclear whether SCR and PLT regulate identical target genes relevant for stem cell niche function.

In contrast to PLT and SCR [whose expression encompasses larger domains, including the QC], expression of the gene encoding homoeodomain transcription factor WUSCHEL (WUS)-RELATED HOMEOBOX 5 (WOX5) is highly enriched in the QC [Sarkar et al. 2007]. WOX5 is also required for QC division rate control and the maintenance of at least a subset of surrounding stem cells [Sarkar et al. 2007; Pi et al. 2015; Zhang et al. 2015]. Both WOX5 and its shoot-expressed homolog, WUS, are required for the function of organizer cells of roots and shoots, respectively [Mayer et al. 1998; Sarkar et al. 2007]. The mechanisms by which PLT and SCR converge in regulating the WOX5 root expression domain and how this links to specification of the QC have remained unknown.

Class I members of the teosinte-branched cycloidea PCNA (TCP) protein family encode plant-specific transcription factors [Li et al. 2005; Hervé et al. 2009; Martin-Trillo and Cubas 2010; Li 2015]. Class I TCPs are implicated in the coordination of cell proliferation and development, especially during leaf development, lateral branching, and shoot apical meristem formation in several plant species [Aguilar-Martínez and Sinha 2013; Davière et al. 2014]. Loss-of-function mutants in class I TCP genes or EREB factor-associated amphiphilic repression (EAR) motif-fused class I TCP proteins show developmental alterations, suggesting that they are positive regulators of meristem formation [Hervé et al. 2009; Kieffer et al. 2011; Aguilar-Martínez and Sinha 2013; Li 2015].

Here we investigate how the two major PLT and SCR pathways for root stem cell niche specification converge to specify QC cells within the root stem cell niche. We show that both PLTs and SCR interact with specific class I TCP proteins. We provide genetic and molecular evidence that PLT1, PLT3, SCR, TCP20, and TCP21 proteins cooperate for the specification of QC identity and the induction of WOX5 expression in at least four developmental contexts: embryogenesis, primary root formation, secondary root development, and the root regeneration process. Our data connect hitherto separated stem cell pathways through novel protein complexes that regulate the formation of the root stem cell organizer and expression of a key gene involved in its function.

Results

Class I TCP proteins interact with both PLTs and SCR protein through different motifs

PLT1, PLT2, AIL6/PLT3 [referred here as PLT3], and BBM/PLT4 [referred here as PLT4] proteins across the root meristem with maximum levels in the stem cell niche domain, whereas SCR expression is restricted mainly to the QC, cortex/endodermis stem cell, and endodermis [Fig. 1A; Wysocka-Diller et al. 2000; Galinha et al. 2007]. To investigate whether their functions in the stem cell niche might converge at the protein level, we performed yeast two-hybrid screening [Y2H] of an Arabidopsis root cDNA library to search for interacting factors of PLT1, PLT2, PLT3, PLT4, and SCR proteins. Since the full-length coding sequence of PLTs elicited strong autoactivation in yeast cells, we used C-terminal regions, including the double AP2 domain of PLT1 [395 amino acids], PLT2 [380 amino acids], PLT3 [338 amino acids], and PLT4 [376 amino acids], fused to the GAL4 activation domain. In total, 25 putative interactors were identified at least in triplicate for PLT1, 23 were identified for PLT2, 18 were identified for PLT3, 21 were identified for PLT4, and 25 were identified for SCR [Supplemental Table 1]. We focused on two common interactors between the bait proteins. Retransformation full-length cDNAs for one-to-one assays with PLTs or SCR revealed that one of the plant-specific transcription factor TCP family members, TCP20, interacted with PLT1, PLT3, and SCR proteins [Fig. 1B]. We did not find other TCP proteins in the SCR Y2H screening, however, like TCP20, TCP21 is highly expressed across the root meristem. Therefore, we tested the ability of TCP21 to interact with SCR in a one-to-one assay, which flagged TCP21 as a candidate SCR interactor. The Y2H tests did not reveal reproducible interactions between PLT2 and PLT4 with these two TCP proteins. We validated the shared interactors in plant cells using bimolecular fluorescent complementation [BiFC] analysis, where cDNAs driven by the 35S promoter were fused to the N-terminal or C-terminal half of YFP and cotransfected into Arabidopsis mesophyll protoplasts.
TCP proteins possess a centrally located conserved common motif, the TCP domain, which has been implicated in DNA-binding sites as well as TCP–TCP dimerization but not in interactions with other proteins (Cubas et al. 1999; Broholm et al. 2008; Aggarwal et al. 2010; Shutian 2015). To assess whether the PLT and SCR proteins bound to separate or the same TCP interaction domains, we resolved which TCP protein regions are required for PLT and/or SCR interactions. We generated a truncation series of TCPs using three regions— for TCP20, domain A (1–408 base pairs [bp]), domain B (409–600 bp), and domain C (601–945 bp) and, for TCP21, domain A (1–267 bp), domain B (268–480 bp), and domain C (481–720 bp)—represented in Figure 1D. We investigated the interaction strengths of truncated TCPs with full-length PLT3 or SCR in the yeast system. As shown in Supplemental Figure 1, A and B, PLT3 interacted with domain C of TCP20 and TCP21, whereas SCR interacted with domain B of both TCP proteins. These data indicated that TCPs directly interact with PLTs and SCR through different regions C-terminal to the TCP domains. BiFC assays using these truncated TCP fragments confirmed these protein interactions in living plant cells, indicating that TCP20 and TCP21 bind with PLT proteins and SCR through different motifs (Fig. 1E; Supplemental Fig. 1C, D). To further test whether PLT and SCR could simultaneously interact with TCP proteins, we performed coimmunoprecipitation assays after transient expression in Arabidopsis mesophyll protoplasts. TCP20, PLT3, and SCR were fused with 10xMyc, 3xFlag, and 7xHA tags, respectively, driven by 35S promoters and cotransfected into Arabidopsis mesophyll protoplasts. Coimmunoprecipitation assays after anti-Flag pull-down showed that TCP20 interacted with PLT3 and SCR in planta and that TCP20 was required for SCR interaction with PLT3, suggesting that PLTs and SCR assemble in vivo into a TCP-containing complex (Fig. 1F; Supplemental Fig. 1E).

plt, tcp, and scr mutants display synergistic genetic interactions

To assess the potential relevance of PLT and SCR protein interaction mediated through TCP proteins, we investigated genetic interactions in different combinations of
their T-DNA insertion alleles (Danisman et al. 2012). We selected plt1-3, plt3-1, tcp20-1, tcp21-1, and scr-3 mutant lines for the analysis (Supplemental Fig. 2A; Fukaki et al. 1998; Aida et al. 2004; Galinha et al. 2007). Quadruple plt1-3\(^{-/-}\) plt3-1\(^{-/-}\) tcp20-1\(^{-/-}\) scr-3\(^{-/-}\) mutants further enhanced the root phenotype of scr and blocked primary root, but not shoot, growth (Fig. 2A,B; Supplemental Fig. 2C). Specifically, primary root growth and meristem size were reduced in several combinations of loss-of-function alleles of PLT, TCP, and SCR genes (Fig. 2C; Supplemental Fig. 2B–D). We investigated the contribution of individual alleles further by varying plt1-3, plt3-1, and tcp20-1 mutant allele dosage in seedlings homozygous for scr-3\(^{-/-}\) and determined primary root length in genotyped mutant backgrounds. The data revealed that primary root length was affected by PLT and TCP20 dosage in the absence of SCR (Supplemental Fig. 2A; Fukaki et al. 1998; Aida et al. 2004; Galinha et al. 2007). Quadruple plt1-3\(^{-/-}\) plt3-1\(^{-/-}\) tcp20-1\(^{-/-}\) scr-3\(^{-/-}\) mutant (Supplemental Fig. 2D). While the double-homozygous mutant caused severe primary root growth defects, the heteroallelic combination tcp21-1\(^{-/-}\) scr-3\(^{-/-}\) showed an intermediate root length, indicating that SCR function also contributes to root growth in a dose-dependent manner upon TCP21 reduction. Together, our data indicate that a reduced level of each of the four proteins creates dosage sensitivity for the others, which is consistent with their physical interactions being relevant for root growth.

We investigated whether the observed reduction of root growth and meristem cell number in plt, tcp, and scr mutant combinations was accompanied by defects in the root stem cell niche. In the wild type, only differentiated columella cells (CCs) contain large starch granules that are not observed in columella stem cells (CSCs) or in the QC (Fig. 2D). Two days after germination [dag] plt1-3\(^{+/+}\)−classical QC (white) and columella stem cells (CSCs; yellow), respectively. Numbers indicate roots with additional QC divisions of total roots examined. (ND) Not determined. [J–M] pWOX5-GUS expression levels in Columbia-0 (Col-0) (K), tcp20-1\(^{-/-}\) scr-3\(^{-/-}\) (L), and tcp21-1\(^{-/-}\) scr-3\(^{-/-}\) (M) roots. (N,O) GUS expression levels in pWOX5-GUS\(^{+/+}\) wild-type (N) and pWOX5-GUS\(^{-/-}\) plt1\(^{+/+}\) plt3\(^{+/+}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) (O) roots. The images displayed in J–O are representative of at least three independent experiments with >10 seedlings examined that obtained similar results. Bars: A,B, 1 cm; D–I, 30 µm; J–O 40 µm.

Figure 2. The effect of plt, tcp, and scr genetic interaction on growth and stem cell maintenance in the Arabidopsis root. [A,B] Seedlings of wild-type [A] and the plt1-1\(^{-/-}\) plt3-2\(^{-/-}\) tcp20-1\(^{-/-}\) scr-3\(^{-/-}\) quadraple mutant [B] 10 d after germination (dag). (C) Primary root length measurements of indicated wild-type and mutant seedlings from 4 to 10 dag. Values are average lengths (means ± SD) of >25 seedling roots per genotype per time point. (D–I) The root apical meristem of 2-dag seedlings in wild-type [D], plt1\(^{-/-}\) plt3\(^{-/-}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) [E], plt1\(^{-/-}\) plt3\(^{-/-}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) [F], plt1\(^{-/-}\) plt3\(^{-/-}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) [G], plt1\(^{-/-}\) plt3\(^{-/-}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) (H), and plt1\(^{-/-}\) plt3\(^{-/-}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) (I). Arrows indicate the QC (white) and columella cells (CSCs; yellow), respectively. Numbers indicate roots with additional QC divisions of total roots examined. (ND) Not determined. [J–M] pWOX5-GUS expression levels in Columbia-0 (Col-0) (K), tcp20-1\(^{-/-}\) scr-3\(^{-/-}\) (L), and tcp21-1\(^{-/-}\) scr-3\(^{-/-}\) (M) roots. (N,O) GUS expression levels in pWOX5-GUS\(^{+/+}\) wild-type (N) and pWOX5-GUS\(^{-/-}\) plt1\(^{+/+}\) plt3\(^{+/+}\) tcp20\(^{-/-}\) scr-3\(^{-/-}\) (O) roots. The images displayed in J–O are representative of at least three independent experiments with >10 seedlings examined that obtained similar results. Bars: A,B, 1 cm; D–I, 30 µm; J–O 40 µm.
tcp20-1−/+ scr-3−/+ and plt3-1−/+ tcp20-1−/+ scr-3−/+ triple-homozygous mutants displayed ectopic cell divisions in the QC and starch granule accumulation into CSCs (Fig. 2H,I), while the 2-dag scr-3−/+ single mutant still maintains the wild-type stem cell niche anatomy (Fig. 2, cf. D and G). In addition, at 2 dag, the plt1-3−/+ plt3-1−/+ tcp20-1−/+ scr-3−/+ quadruple mutants no longer display the typical cell arrangement associated with a functional stem cell niche and meristem (Fig. 2, cf. D and E). To assess a potentially synergistic role in the stem cell niche for PLT, TCP, and SCR proteins, we generated plt1-3−/+ plt3-1−/+ tcp20-1−/+ scr-3−/+ trans-heterozygous mutant seedlings. Like in wild type, starch granule accumulation in plt1-3−/+ plt3-1−/+ tcp20-1−/+ scr-3−/+ mutants indicates the presence of starch-free CSCs below the QC. However, these mutants revealed extra cell divisions in the QC, which is generally associated with improper QC function (Fig. 2F; Blilou et al. 2009; Ten Hove et al. 2015). We traced back the origin of the abnormal root phenotype of the plt3-1−/+ tcp20-1−/+ scr-3−/+ and plt1-3−/+ tcp20-1−/+ scr-3−/+−/+−/+−/+ triple-mutant combinations at 16-cell and earlier stages revealed no morphological defects.

PLT1 mRNA accumulation in the preglobular stage embryo is restricted to basal cells, which will form the QC in later stage (Blilou et al. 2005). To assess whether the expression patterns of other PLT, TCP, and SCR proteins were consistent with a role in QC specification, we analyzed the expression level and distribution of PLT3, TCP20, TCP21, and SCR proteins during the relevant embryonic stages using functional fusion proteins. SCR, TCP20, and TCP21 protein fusions were specifically enriched in the hypophyseal cell at the dermatogen stage. Also at that stage, weak PLT3 expression was detected (Fig. 3B,C). At the early globular stage, the PLT3, TCP20, TCP21, and SCR proteins are enriched in the QC and neighboring cells at all embryonic stages and become more prominent in the apical region at the heart stage (Fig. 3B [right], C). It is at the early globular stage that WOX5 expression is initiated, consistent with the observed dependency of WOX5 expression on these factors in post-embryonic roots (Figs. 2J–O, 3B). It is of note that both TCP20 and TCP21 are expressed in (precursor) QC cells, but TCP21 expression is more restricted and retained in these cells from the dermatogen stage to the late globular stage (Fig. 3B). Taken together, the overlapping expression profiles of relevant PLT, TCP, and SCR fusion proteins coincide with the defects in the establishment of appropriate cell division planes in the hypophyseal cell that accompany the formation of the stem cell niche.

To test whether abnormal early stem cell progenitor formation interfered with the later development of the root stem cell niche during embryogenesis, we quantified the number of columella cells and frequencies of cell divisions in the QC in mature embryos of wild type, scr-3−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/+−/...
niche during regeneration, we performed QC laser ablations and monitored expression recovery of PLT3, SCR, and TCP20 proteins as well as the QC marker pWOX5::H2B-YFP. Six hours after QC ablation, PLT3 already showed expression in the region of the presumptive regenerating stem cell niche when other regulatory factors (SCR and TCP20) were still undetectable (Supplemental Fig. 3). Six hours later, PLT3 and TCP20 showed pronounced expression in this area, and we detected SCR-YFP and pWOX5::H2B-YFP in the newly regenerated area (Supplemental Fig. 3). These observations and the reported requirement of stem cell niche regeneration on PLT and SCR genes (Xu et al. 2006) suggest a scenario in which the PLT–TCP–SCR expression pattern overlap triggers the selection of organizer cells also in the context of regeneration.

PLT–TCP–SCR genes are redeployed for QC specification during lateral root development

QC specification is reiterated in a different developmental context during the post-embryonic formation of lateral

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Figure 3. Genetic interaction among PLT, TCP, and SCR during embryogenesis. (A) The wild-type (Col-0) and plt1-3+/− tcp20-1−/− scr-3−/− embryos at the dermatogen to globular stages. Original cleared images (left) and merged images with tracings of embryos (right) are shown. Bars, 20 µm. (B) Expression patterns of PLT3-YFP, SCR-YFP, TCP20-YFP, and pWOX5::H2B-YFP at the transition between the dermatogen and early globular stages (left and middle) and expression heat maps during the late globular/transition stages (right) are shown. Bars, 10 µm. (C) YFP signal intensities of PLT1-YFP, PLT3-YFP, SCR-YFP, TCP20-YFP, TCP21-YFP, and pWOX5::H2B-YFP during the octant to transition/heart stages exemplified in B. Box length represents the range in which the central 50% of the values fall, with the box edges at the lower (orange) and upper (gray) quartiles. The whiskers indicate the highest and lowest values. YFP fluorescence intensities (n > 15) in the QC (or its precursor cells in the octant and dermatogen stages) were quantified using ImageJ. (D) Cellular anatomies of the radicle in wild-type and plt1-3+/− tcp20-1−/− scr-3−/−, plt3-1+/− tcp20-1−/− scr-3−/−, tcp20-1−/− scr-3−/−, and plt1-3+/− tcp20-1−/− scr-3−/− homozygous mature embryos. The numbers of embryos that showed improper cell divisions in the QC per examined total embryos are indicated in the respective panels. (Yellow arrows) Position of the QC; (dots) positions of the columella cell layers. Bars, 30 µm.

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and the stable expression of central cells act as QC progenitors based on fate mapping. In the stage III LRP, the round s of periclinal cell divisions in the central cells generated a single layered stage I lateral root primordium (LRP) composed of up to eight to 10 cells of equal size. Two rounds of periclinal cell divisions in the central cells generate the stage III primordium. In the stage III LRP, the central cells act as QC progenitors based on fate mapping and the stable expression of WOX5 in these cells from that stage onward (Supplemental Fig. 4A; Ditengou et al. 2008; Goh et al. 2016; Du and Scheres 2017).

To investigate the potential relevance of PLT–TCP–SCR cooperative action during LRP formation, we monitored LRP development in 7-dag seedlings of wild type and three different plt tcp scr mutant combinations (Fig. 4, Supplemental Fig. 4). All compound mutant LRP s tested as well as the scr-3 single mutant itself were indistinguishable from wild type at stage I and early stage II (Fig. 4A-7,B-7,C-7). However, from late stage II onward, we observed aberrant cell sizes and division planes in the cell files that generate QC progenitor cells in wild type (Fig. 4B1–D, Supplemental Fig. 4A). To correlate the region of morphological abnormalities in compound mutant LRP-to-QC specification, we followed pWOX5-erGFP expression, which we consistently detected in the central cells of interposed cell layers at the emerging state, whereas consistent signals were undetectable in stage I and generally initiated in stage II (Fig. 4A–7,B–7,C–7). In contrast to the strong effects on LRP morphology in the compound mutants, we observed no strong reduction of LRP density in single, double, and triple plt tcp scr mutants (Fig. 4E) but rather an increase in the number of abnormal lateral roots with defective meristems, indicative of the failure to maintain a stem cell niche (Fig. 4D).

We asked whether PLT, TCP, and SCR proteins, such as in the context of the primary root, might cooperatively regulate WOX5 expression during lateral root formation.

Figure 4. The lateral root formation in various plt–tcp–scr mutant combinations. [A–I–C–7] Early developmental stages during lateral root initiation in stage I [A–I–A–7], stage II [B–I–B–7], and emerging [C–I–C–7]. Cell lineage maps are shown in A–I (stage I), B–I (stage II), and C–I (emerging). [B–I–C–I] QC progenitor cells appear from late stage II onward (highlighted in red). Morphologies of stage I, stage II, and the emerging stage present in wild-type Col-0 (A–2,B–2,C–2), scr-3 [A–3,B–3,C–3], plt1-3–/–tcp20-1–/–scr-3–/– (A–4,B–4,C–4), plt3–1–/–tcp20-1–/–scr-3–/– [A–5,B–5,C–5], and tcp21-1–/–scr-3–/– [A–6,B–6,C–6], respectively. pWOX5-erGFP expression is observed in stage II [B–7] and the emerging stage [C–7] but is absent in stage I [A–7] wild-type primordium. Bars, 50 µm. [D] The frequencies (percentage) of the abnormal LRP s on the primary root from 8-dag wild-type and plt–tcp–scr mutant combination lines, as indicated. [Blue bar] Normal primordium; [red bar] abnormal primordium. Error bars show SDs. The letters above the bars (a, b, c) indicate significant differences [one-way ANOVA and Tukey’s test, P < 0.01]. Results are means ± SD, n = 15 per line. [E] Statistical analysis of the number of lateral roots per centimeter of the primary root from the 8-dag wild-type and the series of plt–tcp scr mutant combinations. Results are means ± SD, n = 15, no significant differences by one-way ANOVA. [F–I] Expression levels of pWOX5-GUS−/− in LRPCs of wild-type [F,G] and plt1-3–/–plt3-1–/–tcp20-1–/–scr-3–/– mutant [H,I] backgrounds. [F,H] Stage III. [G,I] Emerging LRP. Bar, 40 µm. [J] Statistical analysis of pWOX5-GUS−/− expression in F–I. Results are means ± SD, n = 46 in wild-type; n = 40 in plt1-3–/–plt3-1–/–tcp20-1–/–scr-3–/–−/−. [**] P < 0.01, compared with the corresponding values of wild-type seedlings; two-tailed t-tests.
To this end, we monitored WOX5 promoter activity in pWOX5-GUS/+ and pWOX5-GUS/+pLT1-3-1/+/pLT3-1-1/+/tcp20-1-1/+/scr-2-1/+ LRP s [Fig. 4F-I]. In wild type, WOX5 expression was observed from stage II onward during lateral root formation and gradually increased in strength [Fig. 4B-7, C-7, F,G; Du and Scheres 2017]. The trans-heterozygous pWOX5-GUS/+pLT1-3-1/+/pLT3-1-1/+/tcp20-1-1/+/scr-2-1/+, however, possessed weaker expression at stage II and later stages [Fig. 4H-J]. Our results suggest that, like in primary roots, the induction of WOX5 expression during lateral root formation depends on joint activity of PLT, TCP, and SCR.

To determine whether PLT, TCP, SCR, and WOX5 expression patterns during LRP development coincide with the morphological and gene expression defects observed in their mutants, we compared the expression dynamics of five fusion proteins [PLT3, SCR, SHR, TCP20, TCP21] and WOX5 promoter activation during lateral root initiation [Supplemental Fig. 4B]. PLT3, TCP20, and TCP21 fusion proteins were detected at stage I, whereas SCR expression was consistently observed in the stage II outer cell layer [Goh et al. 2016; Du and Scheres 2017]. Notably, WOX5 promoter activity was detected from stage II onward in cells where PLT, SCR, and TCP proteins are all expressed. Together with our genetic data, these results indicate that joint activity of PLT–TCP–SCR proteins is essential for the specification of the QC and function of the stem cell niche in two very different developmental contexts: the early embryo and post-embryonic lateral root formation.

Expression overlap and synergistic activity of PLT and SCR proteins in the stem cell niche can limit WOX5 gene expression to the QC

PLT proteins display a graded distribution and may assign stem cell niche states in a dose-dependent manner [Galinha et al. 2007; Mähönen et al. 2014]. We asked whether the distribution of PLT–TCP–SCR proteins in the context of the primary root might explain the domain of QC specification. To this end, we quantified expression levels of the four PLTs [PLT1–PLT4], SCR, and two TCPs [TCP20 and TCP21] in the root meristem [Fig. 5]. We classified 15 cell positions in and around the stem cell niche (Fig. 5N) and calculated fluorescence intensities in PLT-YFP, TCP-YFP, and SCR-GFP protein fusion lines pPLT1::PLT1-YFP, pPLT2::PLT2-YFP, pPLT3::PLT3-YFP, pPLT4::PLT4-YFP, pTCP20::TCP20-YFP, and pTCP21::TCP21-YFP, and pSCR::SCR-YFP using ImageJ [Fig. 5A-G, O-U].

pPLT1::PLT1-YFP, pPLT2::PLT2-YFP, pPLT3::PLT3-YFP, pPLT4::PLT4-YFP, and pSCR::SCR-YFP lines were selected previously for [near] wild-type function activities by complementation tests, and we used these homozygous lines for analysis [Galinha et al. 2007; Cruz-Ramírez et al. 2012]. Both TCP20 and TCP21 are broadly expressed across the root stem cell niche at 2 dag, coinciding with the reactivation of the post-embryonic stem cell niche after seed germination [Fig. 5F, G, L, M, T, U]. Consistent with previous reports, all of the PLT proteins measured revealed graded distributions that peak within the stem cell niche. However, the expression levels among PLTs throughout the root meristem were diverse [Fig. 5A-D], PLT1 and PLT3 fusions, which have the most prominent TCP interaction, peak in QC cells and vascular initial cells [Fig. 5A, C, H, J, O, Q]. PLT2-YFP was more broadly expressed and revealed no significant differences among neighboring cells within the stem cell niches [Fig. 5B, I, P]. PLT4-YFP expression peaked in the QC and vasculature, but its level was low overall [Fig. 5C, R]. SCR-YFP was expressed mainly in the QC and endodermis cell lineages, and its expression level revealed a shallow proximodistal gradient with a peak in the QC, cortex/endodermis stem cells, and their immediate daughters [Fig. 5E, I, O].

With quantitative data on the PLT and SCR protein accumulation levels at hand, we asked whether the synergistic activity of TCP20- and TCP21-interacting PLT proteins with SCR could effectively limit WOX5 gene expression to the QC. Indeed, when fluorescence intensities of PLT3-YFP and SCR-YFP were combined by multiplication, maximal synergy was achieved in the QC, where pWOX5-erGFP is most abundantly expressed [Fig. 5V].

PLTs directly induce WOX5 gene expression by cooperative interaction with TCP and SCR

Our findings implied that PLT–TCP–SCR transcription factors function together in QC function and specification as measured by WOX5 gene expression. wox5-1 mutants display extra cell divisions in the QC and lack a columella stem cell layer [Sarkar et al. 2007], which is similar to the late embryogenesis phenotypes observed in compound plt tcp scr mutant combinations. WOX5 may therefore be a significant downstream effector of PLT–TCP–SCR activity. To uncover the mechanism involved, we asked whether WOX5 could be directly regulated by these transcription factors. Previously, in vivo binding sites for PLT transcription factors have been identified on the WOX5 promoter, and, in addition, WOX5 gene expression is induced upon ectopic dexamethasone [DEX] induction of glucocorticoid receptor [GR]-fused PLT2 [Santucci et al. 2016]. We examined whether the WOX5 promoter could also be [ectopically] activated in a DEX-inducible PLT1-GR and PLT3-GR activation system. After 3 h of DEX induction in both PLT1-GR and PLT3-GR lines, pWOX5-erGFP expression slightly expanded into the CEI and CEID in primary root apical meristems within the domain where SCR and TCP proteins reside [Fig. 6A, left and middle panels]. Induced ectopic WOX5 promoter activity in the SCR expression domain was more conspicuous after PLT3-GR activation in lateral root meristems [Fig. 6A, right panel]. To assess whether PLT1 and PLT3 proteins directly activate WOX5 gene expression, we performed quantitative RT–PCR [qRT–PCR] analysis on 3-h DEX– and cyclohexamide [CHX]-treated seedlings of 3SS::PLT1-GR and 3SS::PLT3-GR. WOX5 mRNA levels were increased by DEX and combined DEX/CHX treatment compared with the DMSO mock and CHX control, respectively, consistent with direct activation of WOX5 by PLT1 and PLT3 [Fig. 6B]. The accumulation of WOX5

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transcript in 35S::PLT1-GR and 35S::PLT3-GR seedlings upon induction suggested that PLT1 and PLT3 act as direct transcriptional activators, consistent with the reported occurrence of PLT2 ChIP-seq (chromatin immunoprecipitation [ChIP] combined with high-throughput sequencing) peaks on the WOX5 promoter (Santuari et al. 2016). To further test this, we identified three putative PLT-binding motifs (I–III) in the ChIP-seq data on a 1.6-kb WOX5 promoter fragment (Fig. 6C). Like other reported WOX5 gene constructs, the 1.6-kb WOX5 promoter is expressed most abundantly in the QC. Each motif was replaced with random sequences by site-directed mutagenesis, fused to YFP, and transformed into Arabidopsis wild-type plants. Motif I substitution slightly decreased...
expression, and motif III substitution fully abolished detectable gene expression [Fig. 6D], suggesting that these PLT-binding motifs are crucial for in vivo WOX5 expression in the QC cells of primary roots. To understand whether PLT–TCP–SCR complex function underlies WOX5 induction, we coexpressed PLT3 together with TCP20 and SCR proteins in Arabidopsis mesophyll protoplasts, commonly applied to investigate transcription factor interactions with promoters [Yoo et al. 2007; Long et al. 2015; Díaz-Triviño et al. 2017], and tested their potential to activate the WOX5 promoter. Protoplasts were transfected with reporter constructs harboring endogenous promoter sequences [wild type] or the variant where the predicted PLT-binding motifs were substituted...
(Amotil [I + II + III]) (Fig. 6E,F). PLT3, TCP20, and SCR genes were cotransfected, and WOX5 promoter activities were determined by measuring luciferase (LUC) intensities. We noted that PLT3 alone could activate the WOX5 promoter (Fig. 6E). Single transfection of SCR and cotransfection of TCP20 and SCR were unable to raise WOX5 wild-type promoter activity above that observed with PLT3 transfection alone (Fig. 6E). However, combining PLT3 with TCP20 and SCR expression drastically enhanced WOX5 promoter activity sixfold over the induction observed with PLT3 alone (Fig. 6E). In addition, the WOX5 promoter with disrupted PLT-binding sites could only be activated around threefold over the vector control when cotransfected with PLT3, TCP20, and SCR, whereas the intact WOX5 promoter was enhanced $\geq 20$ times over the vector control (Fig. 6F). In addition, when driving WOX5 gene expression, the 1.6-kb promoter could partially complement the wox5-1 stem cell maintenance phenotype, whereas the version with disrupted motif III-binding sites could not (Supplemental Fig. 5). Collectively, our data show that predicted PLT protein-binding sites are critical for WOX5 induction and that TCP20 and SCR can act as positive regulatory factors mainly through these PLT-binding sites.

**Discussion**

_A dosage-dependent combinatorial model for stem cell organizer specification_

Here we provide evidence of how two major transcription factor modules required for maintenance of the root stem cell niche—the PLT pathway and the SCR pathway—interact at the molecular level with distinct domains of specific members of the TCP protein family. Reduced levels of the relevant PLT, SCR, and TCP proteins in primary roots and their embryonic progenitors, regenerating roots, and LRPs reveal dose-dependent defects in the specification and activity of organizer cells within the root stem cell niche: the QC cells. Our genetic and protein interaction data indicate that dosage-sensitive combinatorial interactions among PLT–TCP–SCR directly regulate promoter activity of the WOX5 gene, which is the best-studied specific molecular marker for QC activity (Sarkar et al. 2007; Pi et al. 2015; Zhang et al. 2017b). Consistent with this scenario, overexpressed PLTs can only effectively expand the WOX5 expression domain where SCR and TCP also are highly expressed.

Our data reveal how the broader roles of PLT and SCR pathways in meristem function are combined for QC specification in a highly specific subdomain, which had been postulated more than a decade ago (Sabatini et al. 2003; Aida et al. 2004; Bennett and Scheres 2010).

Quantification of expression domains indicates that cooperative interaction between PLT1, PLT3, and SCR at specific gene promoters is sufficient to restrict a molecular response to a handful of cells in a single layer (Fig. 5V). The existence of cell-specific complexes that encompass only a part of the, and not the entire, domain of overlap between transcription factors has been corroborated recently by direct visualization of different SCR complexes in the Arabidopsis root using fluorescence resonance energy transfer-fluorescence lifetime imaging (FRET-FLIM) (Long et al. 2017). Note that our cooperative combinatorial model does not exclude the existence of either additional shared targets or many unshared PLT and SCR target genes. Unshared targets can serve broader roles in root development, such as the progression of cell division and differentiation in the root PLT gradient (Mählönen et al. 2014) and the regulation of division and differentiation of the cortical–endodermal lineage by SCR and its binding partner, SHR (Di Laurenzio et al. 1996; Helariutta et al. 2000; Sabatini et al. 2003; Cui et al. 2007; Cruz-Ramírez et al. 2012; Clark et al. 2016; Long et al. 2017). The many additional nonoverlapping functions are consistent with the observation that the published regulated targets of the PLT and SCR pathways are quite distinct (Levesque et al. 2006; Moreno-Risueno et al. 2015; Santuari et al. 2016).

In the shoot apical meristem, the WOX5 sister protein WUS defines the organizing center for the overlying stem cells, and its expression domain is generated by different genetic programs to maintain the shoot stem cell niche [Meng et al. 2017; Zhang et al. 2017a; Zubo et al. 2017]. During initiation of the embryonic shoot stem cells, a WOX2 module promotes the expression of HD-ZIPIII genes to ensure the appropriate ratio of cytokinin and auxin pathways [Zhang et al. 2017b]. During de novo shoot regeneration, cytokinin B-type Arabidopsis response regulators (ARRs) combine with HD-ZIP III coaxactivators for region-specific activation of the WUS promoter [Zhang et al. 2017a], whereas, in the SAM, ARRs appear to maintain mainly WUS expression [Meng et al. 2017; Zhang et al. 2017a]. _WUS_ expression is restricted by the stem cell-expressed CLAVATA3 (CLV3) peptide and interacting receptor kinase pathways [Clark et al. 1997; Brand et al. 2000; Schoof et al. 2000; Müller et al. 2008; Ogawa et al. 2008; Nimchuk et al. 2011]. Notably, the root WOX5 expression domain is also restricted by signaling through the CLV3/EMBRYO-SURROUNDING REGION40 (CLE40) peptide acting through partially overlapping receptor kinases [Stahl et al. 2009, 2013]. Thus, while positive regulators of the expression domain of WUS/BOX5 organizer domain proteins in roots and shoots may be quite distinct, the signaling systems that maintain homeostasis of organizer and (subsets of) stem cells show similarities.

**Developmental plasticity of stem cell specification**

Somatic stem cells are commonly preserved by signals from “organizer cells,” which provide a microenvironment to maintain the undifferentiated state (Scheres 2007; Biteau et al. 2011; Heidstra and Sabatini 2014; Gaillochet and Lohmann 2015). Over the last two decades, research in animals and plants has revealed examples of considerable plasticity in the specification of organizer cells [Boyer et al. 2006; Wilson et al. 2008; Tian et al. 2011; Dejana et al. 2017; Hoeck et al. 2017]. A dramatic demonstration of this plasticity in plants is the
regeneration of an entirely new stem cell niche upon surgical dissection [Xu et al. 2006; Efroni et al. 2016; Zhang et al. 2017a]. In both roots and shoots, this leads to the re-establishment of a new domain of expression of WOX5/WUS. Relocation of WOX5 expression at the center of the new stem cell niche during root regeneration correlates with shifts in the expression domains of PLT and SCR proteins [Xu et al. 2006]. While it has not yet been clarified exactly how these “organizer upstream” expression domains are established, clues for this are suggested from the known root regulatory networks. First, PLT transcription is a slow response to persistent high auxin and therefore can respond to organ injuries that impede auxin flux and thereby change auxin distribution. Second, SCR regulation requires SHR protein movement from vascular to nonvascular cells and subsequent SHR nuclear retention [Sabatini et al. 1999; Dolan 2001; Nakajima et al. 2001; Long et al. 2015]. This nuclear retention is restricted to a single layer through a dynamic developmental control mechanism [Nakajima et al. 2001; Long et al. 2015] and can be envisaged to be reinstated in the regenerating cells that surround intact vascular cells. Therefore, the normal mechanisms that guide PLT and SCR expression are dynamic in that they continue to depend on mobile signals provided by their cellular context. In this way, tissue severing or de novo stem cell niche establishment can create new regions of PLT–SCR overlap, which—as we show in three different contexts [embryogenesis, lateral root formation, and regeneration]—can lead to combinatorial activation of WOX5. The activity of WOX5 then contributes to the specification of new organizer cells, which maintain the undifferentiated state of neighboring stem cells [Sarkar et al. 2007; Forzani et al. 2014; Pi et al. 2015].

Materials and methods

Arabidopsis genetics, transformation, and growth conditions

All of the *A. thaliana* lines used in this study were in the Columbia-0 (Col-0) background. Arabidopsis mutants plt-1-3, plt-3-1, and scr-3 have been described previously [Fukaki et al. 1998; Aida et al. 2004; Galinha et al. 2007]. tcp20-1 (SALK_016203) and tcp21-1 (SALK_106694) mutants were obtained from the Arabidopsis Biological Resource Center (ABRC). plt-1-3/− plt-3-1/− tcp20-1/− scr-3/− trans-heterozygous mutant seedlings were generated by crossing the quadruple homozygote of plt1-3/− plt3-1/− tcp20-1/− scr-3/− with Col-0 and harvesting these siliques (F1 seeds). Combinations of stacked mutant lines with heterozygous alleles in Supplemental Figure 2 were F1 seedlings and were generated as follows: tcp20-1/− scr-3/− was generated by crossing tcp20bcsr with scr. plt3-1/− tcp20-1/− scr-3/− was generated by crossing plt1tcp20bcsr with tcp20bcsr. plt1-1/− tcp20-1/− scr-3/− was generated by crossing plt1tcp20bcsr with scr23scr. plt1-1/− plt3-1/− tcp20-1/− scr-3/− was generated by crossing plt1plt3tcp20bcsr with plt3scr. plt1-1/− plt3-1/− tcp20-1/− scr-3/− was generated by crossing plt1plt3tcp20bcsr with plt3SCR. plt1-1/− plt3-1/− tcp20-1/− scr-3/− was generated by crossing plt1plt3tcp20bcsr with scr. plt1-1/− plt3-1/− tcp20-1/− scr-3/− was generated by crossing plt1plt3tcp20bcsr with scr. Obtained F1 seedling genotypes were confirmed by PCR using proper primer combinations in Supplemental Table 5. All plants used in this study were grown on soil or 0.5× Murashige-Skoog (MS) medium plate under long-day conditions [16-h light/8-h dark period] at 22°C. Seeds were fume-sterilized [100 mL of bleach supplemented with 3 mL of hydrochloric acid], soaked into 0.1% [w/v] agarose, and plated on 0.5× MS plate [pH 5.8]. Floral dip transformation methods were described previously [Clough and Bent 1998].

Construction of promoter and protein fusions

MultiSite Gateway cloning [Invitrogen, Thermo Fisher Scientific, Inc.] was performed to generate transgenic plants expressing TCP20 and TCP21 protein fusions. Promoter sequences upstream of the ATG start codon of TCP20 (2.56 kb) and TCP21 (2.15 kb) were cloned into pDONR P4-P1R. The coding sequences (CDSs) of TCP20 and TCP21 were amplified and cloned into a Gateway-adapted pGEMEasy221. A YFP-NOS terminator cassette was cloned into pGEMEasy221R3. The three types of entry vectors were recombined into pGentinII [Hellen et al. 2000] expression constructs adapted for Gateway cloning that generate bait resistance in plants. These constructs were transformed into Col-0 and also into tcp20-1 for tcp20-1:TCP20-YFP and into tcp21-1 mutants for tcp21-1:TCP21-YFP, of which the extra cell divisions in the QC at 5 day of these genotypes were complemented. To introduce specific motif mutants into the WOX5 promoter, we applied site-directed mutagenesis [Kunkel 1985], for which sequence information is available in Supplemental Table 5. The resultant WOX5 promoters were subcloned into pDONR P4-P1R and fused with H2B-YFP in-frame using the MultiSite Gateway system. The pWOX5-ergfp line was published previously [Xu et al. 2006]. For BiFC analysis in *Arabidopsis* mesophyll protoplasts, all full-length CDSs without stop codons were cloned into pGEMEasy221 vectors and recombined with the N-terminal half YFP (parC235) or C-terminal half YFP (parC236) and driven under the constitutive cauliflower mosaic virus 35S promoter. For transient expression in *Arabidopsis* mesophyll cells, CDS cassettes of PLT3, SCR, and TCP20 genes were fused in-frame with synthesized 3xFlag, 7xHA, and 10xMyc tags, respectively, and recombined into the plant binary vector pGII227 [hygromycin resistance in planta] for the generation of in-frame fusion with the 35S promoter by Multisite Gateway LR reactions [Invitrogen]. Truncated coding sequences of PLT1, PLT3, TCP20, and TCP21 were amplified by PCR, cloned into pGEMEasy221 vector, and recombinated to pDEST22 or pDEST32 for binding assay in a Y2H system. Truncated TCP fragments for BiFC assays were created in p35S-N-terminal fusion YFP or p35S-C-terminal fusion YFP vectors based on parC235 and parC236, respectively. Detailed information of primers used for genotyping of T-DNA insertion lines and the cloning procedures used in this study are in Supplemental Table 5.

**Y2H screening and binding test**

Y2H screenings of PLT and SCR cointeractors were performed using truncated PLT/C-terminal regions of PLT1 [395 amino acids], PLT2 [380 amino acids], PLT3 [338 amino acids], and PLT4 [376 amino acids], including the double AP2 domain] and full-length SCR CDSs into pDEST32 [Invitrogen] in-frame fused with the Gal4-DNA-binding domain. Each construct was cotransformed with an Arabidopsis root cDNA library cloned in pDEST22 [Invitrogen] into yeast strain P669A [James et al. 1996]. Screens were performed on SD medium without His, Leu, and Trp and plus 30 mM 3-aminotriazole [Sigma-Aldrich]. Interaction studies were performed using the recommended protocols for the ProQuest two-hybrid system [Invitrogen]. For one-to-one interaction studies, at least three experimental and three technical replicates
were performed. Primers used for cloning are listed in Supplemental Table 5.

Transient BiFC assay and transcriptional assay in Arabidopsis mesophyll protoplasts

For BiFC assays, plasmids were constructed as indicated above. The transfection protocols for Arabidopsis mesophyll protoplasts, Renilla luciferase transcriptional analysis, and microscopy have been described previously [Cruz-Ramírez et al. 2012, Diaz-Triviño et al. 2017]. More than 200 cells per combination were observed, and at least two independent biological replicates were accomplished. For transcription assays, pWOX5 was first cloned into pGEMEasy221 and subcloned by Gateway LR recombination into pGWBS5 [luciferase] vector and cotransfected with 35S::PLT3-3xFlag, 35S::TCP20-10xMyc, and 35S::SCR-7xHA constructs. The protoplast transformation method was described previously [Yoo et al. 2007]. Protoplasts were transected by 10 µg of DNA in 10 µL of water. Transfected protoplasts were resuspended with WS solution (154 mM NaCl, 125 mM CaCl₂, 5 mM KCl, 2 mM MES at pH 5.7) and incubated overnight at room temperature. Extracted protein mixtures were scored for firefly and Renilla luciferase activity with the dual-luciferase reporter assay system [Promega]. At least three technical replicates for Figure 6, E and F, were accomplished. P-values were calculated by Tukey’s honest significant difference (HSD) test [Fig. 6E] and two-tailed t-test [Fig. 6F].

Microscopy analysis, image processing, and GUS staining

Arabidopsis roots were mounted in water containing 10 mM propidium iodide (Sigma-Aldrich) and imaged with a confocal laser-scanning microscope. For the embryos, whole-mount immature seeds were dissected from siliques and transfected to a solution containing 5% glucose. Stained embryos and primary and lateral roots by Schiff reagent were mounted in chloral hydrate solution (chloral hydrate:water:glycerol, ratio [w/v/v] 8:3:1) and visualized by confocal optics [Leica SP2 and Zeiss LSM710]. Details of the Schiff stain method are described in Truernit et al. (2008). LRPs and embryos were observed for >20 seedlings and 50 embryos, respectively. Cultures were then transferred to 0.5× MS agar plates containing 10 mM CHX (Sigma) for 15 min. Next, they were immediately transferred to 0.5× MS agar plates containing 10 µM DEX (Sigma) plus 10 µM CHX control for another 3 h. For DEX induction tests, seedlings were similarly germinated on 0.5× MS agar plates for 3 d. Next, they were all transferred to 0.5× MS agar plates containing 10 µM DEX [Sigma] plus 10 µM CHX control for another 3 h. Total root RNA isolation was performed with the plant RNA purification kit [Qiagen]. For each RNA sample, 1 µg of total RNA was treated with DNase I [Fermentas] and reverse-transcribed using SuperScript III first strand synthesis system [Invitrogen] according to the manufacturer’s instructions. Quantitative real-time PCR was performed using LightCycler 480 II [Roche Diagnostics GmbH]. Obtained results were normalized against UBQ10 expression. Primer information is in Supplemental Table 5. At least two technical replicates with two biological replicates were performed for each time point and sample. P-values were calculated by Tukey’s HSD test. The study of spatio-temporal induction of the WOX5 promoter by PLT1 and PLT3 induction was performed using p35S::PLT1-GR and p35S::PLT3-GR seedlings harboring homozygous pWOX5::egfp, respectively. For each line, seeds were germinated on 0.5× MS agar plates for 2 or 5 d for analysis of primary root or lateral root meristems, respectively. For induction, the seedlings were treated with 10 µM DEX for 3 h followed by the fluorescence imaging by confocal microscopy. Seedlings treated with DMSO for 3 h were used as controls.

Communoprecipitation and Western blotting

For transient expression analysis in Arabidopsis mesophyll protoplasts, cells were ground in liquid nitrogen using a mortar and pestle. Powdered materials were mixed with extraction buffer [50 mM Tris-HCl at pH 7.5, 100 mM NaCl, 1% Triton X-100, protease inhibitors]. After thawing, samples were centrifuged at 14,000 rpm for 10 min at 4°C. The resulting supernatant was incubated with anti-Flag antibodies under rotation for 4 h at 4°C. Bound fractions were immobilized on protein G beads [GE healthcare] and washed twice with 200 µL of extraction buffer. Precipitates were eluted from beads with 40 µL of SDS-PAGE loading buffer (1% SDS, 0.1 M dithiothreitol, 0.08 M Tris-HCl at pH 6.8, 5% glycerol, bromophenol blue). Eluted proteins were separated on 10% polyacrylamide SDS gels and transferred to nitrocellulose membranes [Hybond, GE Healthcare] by blotting. All of the primary monoclonal antibodies produced in mice were incubated for 1 h in 3% non-fat milk in PBS-T [0.1% Tween], and subsequent anti-mouse secondary antibodies marked HRP were incubated for another 1 h in PBS-T [0.1% Tween]. The following antibodies were used: anti-Myc antibody (Santa Cruz Biotechnology), anti-Flag antibody

Laser ablations and regeneration experiments

Laser ablations of QC cells were performed on a Leica TCS SP2 confocal microscope using 4-dag seedlings of pPLT3::PLT3-YFP, pSCR::SCR-YFP, pTCP20::TCP20-YFP, or pWOX5::H2B-YFP expressing lines (n = 5) as described previously [van den Berg et al. 1995; Xu et al. 2006]. Briefly, roots were pre-trained with 10 µg/mL propidium iodide on slides to visualize the cellular structure of stem cells and target the QC for laser ablation. After ablation, seedlings were rinsed with distilled water and put back on 0.5× MS plates. Regeneration was monitored every 6 h for 24 h.

RNA isolation and qRT-PCR analysis of PLT-GR

PLT1-GR- and PLT3-GR-induced activation of the WOX5 promoter by qRT–PCR analysis was performed as follows: For DEX + CHX tests, p35S::PLT1-GR and p35S::PLT3-GR seedlings were germinated on 0.5× MS agar plates for 3 d and subsequently transferred to 0.5× MS agar plates containing 10 µM CHX [Sigma] for 15 min. Next, they were immediately transferred to 0.5× MS agar plates containing 10 µM DEX (Sigma) plus 10 µM CHX control for another 3 h. For DEX induction tests, seedlings were similarly germinated on 0.5× MS agar plates for 3 d. Next, they were all transferred to 0.5× MS agar plates containing 10 µM DEX for another 3 h. Total root RNA isolation was performed with the plant RNA purification kit [Qiagen]. For each RNA sample, 1 µg of total RNA was treated with DNase I [Fermentas] and reverse-transcribed using SuperScript III first strand synthesis system [Invitrogen] according to the manufacturer’s instructions. Quantitative real-time PCR was performed using LightCycler 480 II [Roche Diagnostics GmbH]. Obtained results were normalized against UBQ10 expression. Primer information is in Supplemental Table 5. At least two technical replicates with two biological replicates were performed for each time point and sample. P-values were calculated by Tukey’s HSD test. The study of spatio-temporal induction of the WOX5 promoter by PLT1 and PLT3 induction was performed using p35S::PLT1-GR and p35S::PLT3-GR seedlings harboring homozygous pWOX5::egfp, respectively. For each line, seeds were germinated on 0.5× MS agar plates for 2 or 5 d for analysis of primary root or lateral root meristems, respectively. For induction, the seedlings were treated with 10 µM DEX for 3 h followed by the fluorescence imaging by confocal microscopy. Seedlings treated with DMSO for 3 h were used as controls.
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