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RESEARCH ARTICLES

Spatiotemporal Regulation of Lateral Root Organogenesis in *Arabidopsis* by Cytokinin[™]

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The architecture of a plant's root system, established postembryonically, results from both coordinated root growth and lateral root branching. The plant hormones auxin and cytokinin are central endogenous signaling molecules that regulate lateral root organogenesis positively and negatively, respectively. Tight control and mutual balance of their antagonistic activities are particularly important during the early phases of lateral root organogenesis to ensure continuous lateral root initiation (LRI) and proper development of lateral root primordia (LRP). Here, we show that the early phases of lateral root organogenesis, including priming and initiation, take place in root zones with a repressed cytokinin response. Accordingly, ectopic overproduction of cytokinin in the root basal meristem most efficiently inhibits LRI. Enhanced cytokinin responses in pericycle cells between existing LRP might restrict LRI near existing LRP and, when compromised, ectopic LRI occurs. Furthermore, our results demonstrate that young LRP are more sensitive to perturbations in the cytokinin activity than are developmentally more advanced primordia. We hypothesize that the effect of cytokinin on the development of primordia possibly depends on the robustness and stability of the auxin gradient.

INTRODUCTION

The fundamental task of the root system is to mediate an interaction between plant and soil and to provide the plant with water and nutrients. The architecture of the root system is determined by endogenous mechanisms that constantly integrate environmental signals, such as availability of nutrients and moisture or salinity, and, accordingly, modulate the primary root growth and branching. The developmental flexibility of root systems, which is extremely important for efficient soil exploitation and survival of plants under less favorable conditions, is largely determined by its postembryonic capacity to branch.

In Arabidopsis thaliana, lateral roots initiate from selected pericycle cells that acquire attributes of founder cells, subsequently undergo a series of anticlinal divisions, and produce a few short initial cells. After initiation, coordinated cell division

Overall, the root system architecture depends largely on the spatiotemporal regulation of individual lateral root initiation (LRI) events. Notably, processes leading to initiation seem to be activated much earlier than the morphological detection of the first anticlinal divisions. Priming, proposed to occur in the basal root meristem by recurrent accumulations of the plant hormone auxin in selected protoxylem cells, precedes founder cell establishment and LRI (De Smet et al., 2007; De Rybel et al., 2010). Several concepts have recently been put forward to assess the mechanisms behind the repetitive initiation events. Either correlation between gravity responses and LRI that regulate root bending (De Smet et al., 2007; Lucas et al., 2008a) or the mechanical stretching of cells as a consequence of the root bending by gravistimulation (Laskowski et al., 2008) have been proposed to underlie the regulation of LRI. Another hypothesis considers regular (gravity-independent) oscillations in a set of genes, including transcriptional regulators, as the primary mechanism recurrently activating processes that result in LRI

and differentiation give rise to lateral root primordia (LRP) that continue to grow and emerge through the cortex and epidermal layers of the primary root; finally, the lateral root apical meristems take over the growth regulation (Malamy and Benfey, 1997; Péret et al., 2009; Benková and Bielach, 2010).

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(Moreno-Risueno et al., 2010). Spatially, LRI is restricted to a developmental window, namely, a distinct region within the primary root with the highest probability for LRI events. This developmental window was defined as a root growth period during which pericycle cells in the young differentiation zone maintain a state permissive for LR founder cell specification. Therefore, the term comprises specific time, space, and cell state (Dubrovsky et al., 2006).

Plant hormones are important endogenous regulatory molecules that control plant growth and development and mediate interactions with the environment to modulate developmental responses. Auxins and cytokinins are two classes of plant hormones that are key in the regulation of lateral root organogenesis. Auxin promotes the earliest events related to lateral root organogenesis, including priming (De Smet et al., 2007), founder cell specification (Dubrovsky et al., 2008), and initiation (Benková et al., 2003; Dubrovsky et al., 2008), as well as the later phases of primordium formation and emergence (Benková et al., 2003; Swarup et al., 2008). Critical for the auxin activity is the dynamic regulation of its distribution mediated through the polar auxin transport machinery (Vanneste and Friml, 2009) and the downstream signaling pathway (Paciorek and Friml, 2006; Quint and Gray, 2006; Calderon-Villalobos et al., 2010). Interference with either of them (polar auxin transport or signaling) leads to severe defects in LRI and development (Casimiro et al., 2001; Fukaki et al., 2002; Benková et al., 2003; Dharmasiri et al., 2005; Okushima et al., 2005; Swarup et al., 2008; De Smet et al., 2010).

By contrast, cytokinin constrains both LRI and development (Li et al., 2006; Laplaze et al., 2007). Plants overproducing cytokinin exhibit reduced lateral root formation, whereas transgenic plants overexpressing the *CYTOKININ OXIDASE/DEHYDROGENASE* (*CKX*) gene coding for the enzyme that degrades cytokinin by oxidation have an increased number of lateral roots (Werner et al., 2003; Laplaze et al., 2007). Accordingly, perturbations in either perception or signal transduction of cytokinins affect lateral root organogenesis (Mason et al., 2005; Riefler et al., 2006; To et al., 2007).

The antagonistic roles of auxin and cytokinin in the regulation of lateral root organogenesis imply that their activities must be mutually tightly controlled. In view of the inhibitory effects of cytokinin on LRI, spatiotemporal regulation and restriction of its activity might be crucial for the recurrent LRI. Here, we show that the early phases of lateral root organogenesis, including priming and initiation, take place in the root zone with elevated levels of biologically active cytokinins, but with repressed cytokinin responses. Accordingly, enhanced cytokinin activity prior to the morphologically visible initiation events, in the zone encompassing the basal meristem activity, most efficiently inhibited LRI; by contrast, increased levels of cytokinins in the mature differentiated parts of the roots interfered with the development. but not the initiation, of lateral roots. Enhanced cytokinin responses detected in the pericycle cells neighboring the LRP and the analysis of mutants defective in cytokinin biosynthesis or response indicate that this cytokinin activity might be important to prevent LRI in close proximity to existing LRP. Furthermore, our results demonstrate that primordia in the early phases of development are more sensitive to perturbations in the cytokinin pathway than those in later phases. We hypothesize that the stage-dependent effect of cytokinin on primordium development is determined by robustness and stability of the auxin gradient that might be enhanced in developmentally more mature primordia.

RESULTS

LRI Occurs in the Root Zone with High Active Cytokinin Levels

Particular phases of lateral root development take place in distinct zones of the primary root. Priming, an accumulation of auxin in protoxylem cells, and LRI are spatially restricted to the zone near the root meristem, including the basal meristem and developmental window (Dubrovsky et al., 2001, 2011; De Smet et al., 2007), whereas primordium organogenesis and emergence occur in the mature differentiated zones (Swarup et al., 2008). Cytokinin acts as an endogenous inhibitor of lateral root organogenesis and counteracts the stimulatory effect of auxin (Li et al., 2006; Laplaze et al., 2007). However, thus far, the mechanisms behind this crosstalk are unclear. As a first step, we investigated whether cytokinin levels vary along the primary root axis and correlate with lateral root organogenesis. Cytokinin levels were examined in three zones defined by the developmental phases of lateral root organogenesis: the zones of LRP initiation, formation, and LRP emergence through the cortex (Figure 1A). The LRP initiation zone was recognized as the root area where early phases of initiation take place, including priming in the basal meristem, founder cell establishment, and the first stage of LRP development in the developmental window. The LRP formation zone was defined as the zone between the first-stage LRP and the first emerged lateral roots. The root zone from the first emerged lateral roots to the junction between root and hypocotyl was recognized as the LR emergence zone (for details, see Methods). Quantification of cytokinin metabolites in the three different root segments revealed that levels of the cytokinin bases isopentenyladenine and cis-zeatin were slightly higher in the root segments corresponding to the LRP initiation zone than those in the more proximal parts (Figure 1A), whereas trans-zeatin was distributed evenly through the whole root. Cytokinin ribosides are transport forms of cytokinin, which might be activated and therefore represent sources of rapidly available active cytokinins (Kudo et al., 2010). The trans-zeatin riboside, cis-zeatin riboside, isopentenyl adenosine, and dihydrozeatin riboside were distributed similarly to the corresponding free cytokinin bases (Figure 1A). On the contrary, cytokinin-inactive derivatives, such as trans-zeatin-9-glucoside and cis-zeatin-9-glucoside, are conjugated forms that cannot be reactivated into biologically active cytokinins. A high proportion of these biologically inactive derivatives accumulated in the older root segments, corresponding to the LRP emergence zone (Figure 1A).

Altogether, the analysis of the cytokinin levels in roots suggests that spatiotemporally regulated metabolic processes, such as degradation or inactivation by conjugation, contribute to a differential distribution of cytokinin derivatives along the primary root. With respect to lateral root organogenesis, the early phases, including initiation and primordium formation, take place in the root zone where biologically active or activatable

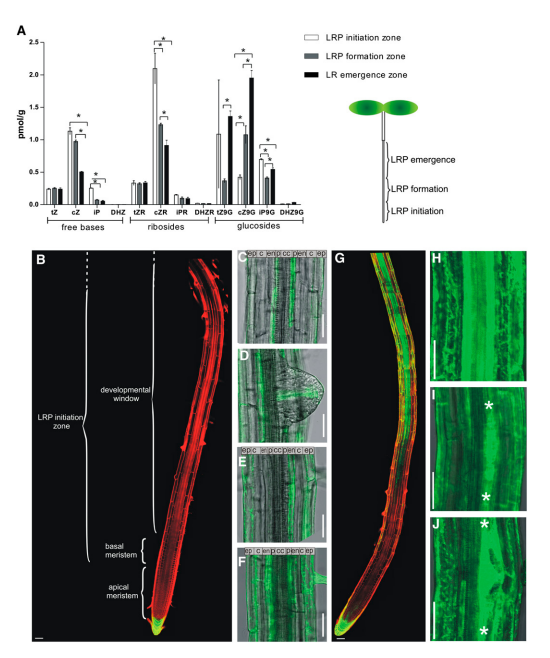


Figure 1. Distribution of Cytokinin and Cytokinin Response in Roots.

(A) Quantification of cytokinin levels in the zones of LRP initiation zone, formation, and emergence. Error bars denote se (*P < 0.05; Student's t test; n = 3).

(B) to (F) TCS:GFP expression in 8-d-old untreated roots detected in the root cap, but not in the root meristem, basal meristem, and developmental window (B); in the individual xylem pole pericycle cells between developing young primordia (C); in the provasculature tissue in emerging LRPs (D); in the endodermal cells adjacent to early-stage LRPs (E); and continuously in the endodermis of the emerging primordium zone (F).

(G) to (J) *TCS:GFP* expression in 8-d-old roots treated for 16 h with 10 μM benzyl adenine induced in the provasculature of the root, but not basal, meristem (G) and in all tissues of the differentiation zone ([G] and [H]), including primordia at all developmental stages ([I] and [J]). Plasma membranes were visualized by propidium iodide staining. c, cortex; cc, central cylinder; cZ, *cis*-zeatin; cZ9G, *cis*-zeatin-9-glucoside; cZR, *cis*-zeatin riboside; DHZ, dihydrozeatin; DHZ9G, dihydrozeatin-9-glucoside; DHZR, dihydrozeatin riboside; en, endodermis; ep, epidermis; iP, isopentenyladenine; iP9G; isopentenyladenine-9-glucoside; iPR, isopentenyl adenosine; p, pericycle; tZ, *trans*-zeatin; tZ9G, *trans*-zeatin-9-glucoside; tZR, *trans*-zeatin riboside. White asterisks indicate borders of LRPs. Bars = 50 μm.

cytokinin derivatives accumulate, but the later developmental phases, such as emergence, occur in the zone with accumulation of biologically inactive cytokinin conjugates.

Cytokinin Responses Are Minimal in the Basal Root Meristem

The main limitation of analytical approaches, such as liquid chromatography-mass spectrometry-based cytokinin quantifications, is the lack of tissue and cellular resolution. Therefore, reporters sensitive to defined hormones, such as synthetic auxin response element DR5 and domain II (DIIVENUS) for auxin (Ulmasov et al., 1997; Vernoux et al., 2011) or synthetic EIN3responsive promoter (EBS) to ethylene (Stepanova et al., 2007), have helped elucidate the cellular action of hormones. To monitor the cytokinin responses in cells and tissues involved in lateral root organogenesis (i.e., the basal meristem, the developmental window zone, the pericycle cells, and the individual stages of LRP and cells adjacent to LRPs), we used the cytokinin-sensitive two-component output sensor (TCS) reporter, which harbors concatemerized B-type Arabidopsis response regulator binding motifs and a minimal 35S promoter. Only cytokinins activate TCS-mediated transcription, whereas other plant hormones, such as auxin, abscisic acid, and gibberellic acid, do not (Müller and Sheen, 2008). Cytokinin-dependent expression of a TCS:LUCIFERASE (LUC) reporter was shown to depend on well-established cytokinin signaling components (Müller and Sheen, 2008).

Detailed examination revealed that the expression of the TCS: GREEN FLUORESCENT PROTEIN (GFP) reporter was not or hardly detectable in the basal meristem zone where priming events take place and throughout the root zone corresponding to the developmental window (Figure 1B). In the pericycle, TCS:GFP was expressed in individual xylem pole pericycle cells between the LRP in the same cell file or across existing LRP (Figures 1C, 2A, and 2D; see Supplemental Figures 1A and 1B online). In the proximal root zone, where LRPs are more developed, the TCS signal in the pericycle disappeared (Figure 1F; see Supplemental Figure 1A online). The TCS:GFP expression occurred in the endodermal cells adjacent to the early-stage LRP (Figure 1E) and persisted in the endodermis of the root zone with emerging primordia and mature lateral roots, but no GFP signal was detected in the pericycle (Figure 1F; see Supplemental Figure 1A online). In the LRP body, the cytokinin reporter was expressed the earliest at stage VI, where it was confined to the provascular cells (Figure 1D).

Expression of the *TCS* reporter was induced by exogenous cytokinin treatment in all cell layers of the mature differentiated root, including LRP at all developmental stages (Figures 1G to 1H), but not or to a very limited extent in the zone corresponding to the basal meristem and the early differentiation zone. These results suggest that the reduced expression of the *TCS* reporter in the basal meristem and developmental window might result from a local repression of cytokinin signaling that cannot be overcome easily by enhanced cytokinin stimulus (Figures 1G to 1J).

Thus, the TCS:GFP reporter expression revealed that cytokinin signaling was strongly repressed in cells and tissues in which

priming and early initiation phases of the lateral root organogenesis took place. By contrast, the cytokinin response was enhanced in the pericycle xylem pole cells between two existing LRP where organogenesis typically was repressed.

Cytokinin Responses Are Enhanced in the Pericycle Cells Near Young LRP

In pericycle cells, a local increase in auxin response had been shown to correlate with founder cell specification (Dubrovsky et al., 2008). In contrast with the findings on auxin, the cytokinin reporter exhibited an enhanced response in the xylem pole pericycle cells between existing LRP (Figure 1C; see Supplemental Figure 1 online), where LRI was compromised (Dubrovsky et al., 2001; Lucas et al., 2008b). The question arises whether enhanced cytokinin signaling might be part of a mechanism preventing new initiation events near the existing primordia. To this end, we compared the expression of the TCS reporter in pericycle cells of 3-d-old roots without detectable LRI, of 4-d-old roots with one, or occasionally, two primordia initiated per root, and of 7-d-old roots with three to four primordia per centimeter root length. In 3-d-old seedlings, no TCS expression was found in the pericycle cells (cf. Figures 2A and 2C); in 4-d-old roots, TCS:GFP was detected in a few pericycle cells in the proximity of the initiated primordia (cf. Figures 2A and 2D); and in 7-d-old roots, the increased TCS: GFP expression correlated with the lateral root formation (Figure 2A; see Supplemental Figure 1 online). To examine the dynamics of the cytokinin responses following LRP initiation, we analyzed the TCS:GFP expression in real time. In parallel, the auxin signaling profile with DR5:3XVENUS-N7 was monitored, the expression of which had been identified as the earliest sign of LRP initiation (Dubrovsky et al., 2008). Within 5 h after initiation, as defined by the expression of DR5:3XVENUS-N7 in nuclei of a few short initials, the TCS-mediated expression of GFP could be detected in pericycle cells neighboring the LRP. This signal gradually increased with time (Figure 2F).

To test whether enhanced cytokinin activity in the pericycle cells was related to primordium formation or to exclude that the enhanced cytokinin response was an inherent characteristic of the pericycle, we examined the TCS:GFP expression in roots lacking LRP due to inhibited auxin transport (Casimiro et al., 2001). The TCS:GFP signal in the pericycle of roots grown for 7 d on 5 μ M naphthylphthalamic acid (NPA) was dramatically reduced (cf. Figures 2A and 2B). These results suggested that the enhanced cytokinin response in pericycle cells might be linked to the LRP formation process itself and that inhibition of LRP formation interfered with cytokinin responses in the pericycle. In this respect, the TCS:GFP expression strongly differed from the xylem pole pericycle marker J0121, the expression of which in pericycle cells was not affected by NPA treatment (Casimiro et al., 2001) (see Supplemental Figure 2 online). Additionally, cytokinin measurements revealed an increase in cytokinin metabolites, such as the cytokinin bases trans-zeatin and isopentenyladenine, transzeatin riboside-O-glucoside, cis-zeatin riboside-O-glucoside, transzeatin riboside monophosphate, cis-zeatin riboside monophosphate, and isopentenyl adenosine monophosphate, in roots of seedlings germinated on NPA. Thus, the reduced TCS:GFP expression might not be the consequence of reduced cytokinin levels in NPA-treated roots (Figure 2E).

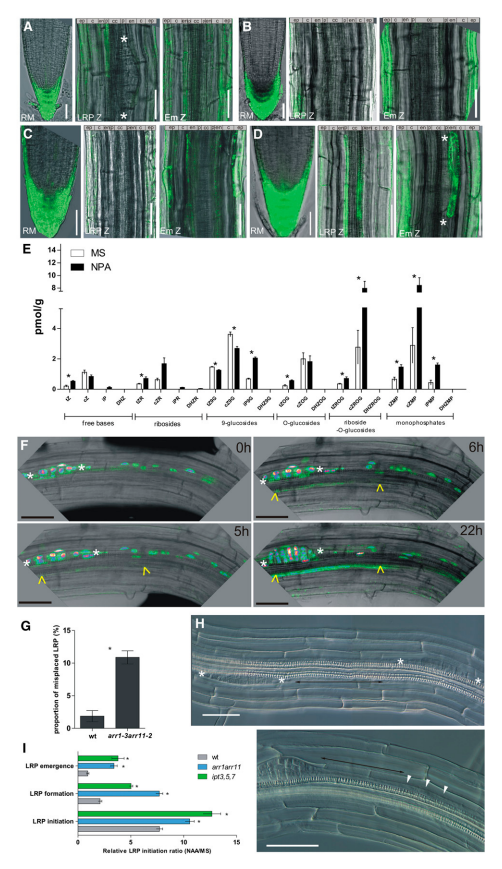


Figure 2. Cytokinin Response in the Pericycle Cells.

In summary, our results suggest that an enhanced cytokinin response in the pericycle is strongly linked to LRP formation and that chemical perturbations of LRP initiation dramatically reduce the *TCS*-mediated expression in the pericycle, which contrasts with the stable expression of pericycle cell identity markers.

Spacing of LRP Initiation Is Defective in Roots with Compromised Cytokinin Response

In Arabidopsis, LRPs initiate in an acropetal manner with the youngest primordia distally positioned from the older LRPs. This architectural pattern of the root system results from strictly spatiotemporally regulated LRP initiation occurring in defined parts of the root behind the root meristem, designated the developmental window (Dubrovsky et al., 2001, 2011). In this zone, LRPs initiate with the highest probability when compared with the proximal part of the root. Accordingly, pericycle cells in the developmental window are more sensitive to exogenous auxin, which promotes LRP initiation, than are pericycle cells of the older, more proximal part of the root (Dubrovsky et al., 2011). Thus, we hypothesize that enhanced cytokinin activity in the pericycle cells between LRPs might be part of the mechanism restricting new initiation events in the proximity of existing LRPs. Therefore, we examined how genetic perturbations in cytokinin levels and signaling affect the positioning of primordia and the sensitivity of pericycle cells from the developmental window to LRP initiationtriggering signals, such as auxin. The mutant ipt3 ipt5 ipt7 with loss-of-function of the ISOPENTENYL TRANSFERASE (IPT) genes that encode the rate-limiting enzyme in cytokinin biosynthesis exhibits reduced cytokinin levels in the roots (Miyawaki et al., 2006). Detailed analysis of the ipt3 ipt5 ipt7 mutant revealed that LRP initiation was not significantly changed (6.54 ± 0.67 LRP/cm versus 6.03 \pm 0.13 LRP/cm in the control). However, when mutant roots were exposed to exogenous auxin, a massive LRP initiation was triggered along the whole root, in contrast with wild-type roots, where the highest LRI occurred in the zone corresponding to the developmental window (Figure 2I). Next, we analyzed mutants lacking the functional B-type *Arabidopsis* response regulator factors that mediate the signal transfer downstream of the cytokinin receptors (To and Kieber, 2008). Both *ARR1* and *ARR11* were expressed in roots and pericycle cells, as confirmed by quantitative RT-PCR analysis (see Supplemental Figure 3A online; for details, see Methods). To test the impact of the lack of ARR1 and ARR11 functions on the cytokinin response, we transiently expressed the *TCS:LUC* reporter in protoplasts isolated from mesophyll cells of the single *arr1* and *arr11* and double *arr1 arr11* mutant plants. The *TCS:LUC* induction decreased significantly in the mutants when compared with controls, indicating that the cytokinin response depended on the ARR1 and ARR11 activity (see Supplemental Figure 3B online).

Detailed phenotype analyses revealed that the arr1 arr11 double mutant was defective in LRP spacing. In 18 examined arr1 arr11 roots, encompassing 267 LRP in total (8.55 ± 0.24 LPR/cm and 2.42 ± 0.2 lateral roots/cm), 31 examples of LRP initiations were found in very close proximity (two LRP one pericycle cell apart), a situation hardly occurring in control seedlings in which four misplaced LRP were detected among 218 LRPs (6.44 \pm 0.31 LRP/cm and 1.74 \pm 0.05 lateral roots/ cm, n = 18 roots) (Figures 2G and 2H; see Supplemental Figure 3D online). Furthermore, similarly to ipt3 ipt5 ipt7, the LRP initiation in arr1 arr11 mutants was significantly increased along the longitudinal root axis in response to auxin treatment and less restricted to the developmental window zone compared with the wild type (Figure 2I). The enhanced sensitivity of pericycle cells in the proximal part of the arr1 arr11 roots to auxin was discernible when roots lacking LRP as a consequence of NPAinhibited auxin transport were treated with auxin. Whereas the auxin-induced LRP initiation clustered to the zone behind the root tip in the wild-type roots, the corresponding zone in the arr1 arr11 roots was enlarged (see Supplemental Figure 3C online).

Analyses of mutants affected in cytokinin metabolism and signaling indicate that enhanced cytokinin activity in pericycle cells

Figure 2. (continued).

(A) and (B) TCS:GFP signal in pericycle cells of 7-d-old roots without treatment (A) and on media supplemented with 10 µM NPA (B).

(C) and (D) TCS:GFP expression in pericycle cells of 3-d-old roots without LRPs (C) and 4-d-old roots with one or two initiated LRPs. Weak TCS signal present in xylem pole pericycle cells and in the endodermis adjacent to LRP (D).

(E) Cytokinin measurements in roots, excluding the root meristem of seedlings grown on control medium (white) and in the presence of 10 μ M NPA (black). Error bars denote se (*P < 0.05; Student's t test; n = 3).

(F) Real-time monitoring of *TCS:GFP x DR5:3XVENUS-N7* reporter in pericycle cells after LRP initiation. White asterisks and yellow arrowheads indicate LRPs at the early initiation stage in pericycle cells visualized by nuclear-localized *DR5:3XVENUS-N7* and the *TCS:GFP* signal, respectively.

(G) and **(H)** Defective LRI positioning in 7-d-old seedlings of the *arr1 arr11* mutant. Proportion of misplaced per total number of LRPs (wild type, n = 218; $arr1-3 \ arr11-2$, n = 267) from 18 roots each. Error bars denote se (*P < 0.05; Student's t test) **(G)**. One pericycle cell distance between two LRPs **(H)**. **(I)** Auxin-induced LRP initiation in $arr1 \ arr11$ and $ipt3 \ ipt5 \ ipt7$ mutants. In the mutant background, 25 h of treatment with 1 μ M 1-naphthaleneacetic acid (NAA) induces LRP initiation along the whole root, whereas in control roots, induction is restricted mainly to the LRP initiation zone. Error bars denote se (*P < 0.05; Mann-Whitney test; n = 15 roots).

White asterisks indicate borders of LRPs. c*Z*, *cis*,zeatin; cZ9G; *cis*-zeatin-9-glucoside; cZMP, *cis*-zeatin riboside monophosphate; cZOG, *cis*-zeatin-O-glucoside; cZR, *cis*-zeatin riboside; cZROG, *cis*-zeatin riboside-O-glucoside; DHZ, dihydrozeatin; DHZ9G, dihydrozeatin-9-glucoside; DHZMP, dihydrozeatin riboside monophosphate; DHZOG, dihydrozeatin-O-glucoside; DHZR, dihydrozeatin riboside; DHZOG, dihydrozeatin riboside; DHZR, dihydrozeatin riboside; DHZNG, dihydrozeatin riboside; DHZNG, dihydrozeatin riboside; Em Z, zone of LRP emergence; iP, isopentenyladenine; iP9G; isopentenyladenine-9-glucoside; iPMP, isopentenyl adenosine; LRP Z, zone of LRP initiation; RM, root meristem; tZ, *trans*-zeatin; tZ9G, *trans*-zeatin-9-glucoside; tZMP, *trans*-zeatin riboside monophosphate; tZOG, *trans*-zeatin-O-glucoside; tZROG, *trans*-zeatin riboside-O-glucoside; tZR, *trans*-zeatin riboside; wt, the wild type. Bars = 50 μm.

might be important to prevent LRI in close proximity to existing LRP. Decreased cytokinin levels and responsiveness in pericycle cells increased the frequency of defects in LRP spacing and significantly enhanced the sensitivity to LRI-triggering signals, such as auxin, of pericyclecells in the proximal root zone.

Cytokinin and Auxin Responses Are Spatially Complementary

Our results show that the TCS-monitored cytokinin signaling, similarly to the DR5-monitored auxin signaling, was tightly linked to LRP organogenesis. However, a comparison of the cytokinin and auxin responses revealed complementary rather than overlapping patterns. Whereas a strong auxin response was detected in the founder cell identity-acquiring pericycle cells and in stage I LRP (Figure 2F; see Supplemental Figure 4A online) (Dubrovsky et al., 2008), the TCS reporter was expressed in pericycle cells neighboring existing LRP (Figure 2F; see Supplemental Figures 1 and 4A online). Similarly, at later developmental stages, DR5 expression maxima were detected at the LRP tips, and the cytokinin reporter expression was restricted to the provasculature (see Supplemental Figure 4B online). Detailed examination of the TCS reporter in the root tip indicated that the cytokinin response was enhanced in columella cells (see Supplemental Figure 4C online), but, in contrast with DR5 that is strongly expressed in the central columella cells, the TCS:GFP signal was the strongest in the outer columella cells surrounding the DR5 domain (see Supplemental Figure 4C online). As shown previously, inhibition of the auxin transport by NPA enlarged the DR5 expression domain in the columella (Sabatini et al., 1999). In the NPA-treated roots, the cytokinin response domain shifted accordingly toward the cell layers of the outer lateral root cap, thus again surrounding the extended domain of auxin response maxima in the central columella (see Supplemental Figure 4C online). Mutual regulation of the expression of auxin and cytokinin reporters by both auxin and cytokinin was further tested with a transient expression assay in Arabidopsis protoplasts. As expected, expression of the cytokinin reporter TCS:LUC was upregulated by cytokinin and not affected by auxin (see Supplemental Figure 4D online) (Müller and Sheen, 2008). However, the cytokinin stimulatory effect on TCS was significantly compromised by the simultaneous application of auxin. Vice versa, auxin, but not cytokinin, stimulated the expression of the DR5:LUC auxin reporter and simultaneous treatment with cytokinin reduced the auxin induction (see Supplemental Figure 4D online). These data suggest that auxin and cytokinin tend to occupy complementary domains and to modulate their activities mutually.

Cytokinin Accumulation in the Priming Zone Represses LRI

Similarly to inhibition of LRI and development by exogenous cytokinin (Li et al., 2006; Laplaze et al., 2007), endogenous increase of cytokinin levels by ectopic expression of the cytokinin biosynthesis gene IPT in differentiated xylem pole pericycle cells reduces the LRP density by $\sim 50\%$ (Laplaze et al., 2007). Recent studies have demonstrated that distal root zones, covering the basal meristem and the developmental window, are the sites at which the earliest decisions take place on priming and LRI. Therefore, to test whether an increase in

cytokinin levels in these zones might affect LRI, we expressed *IPT* in a tissue-specific manner with the *GAL4>>UAS*-based two-component activator-reporter system (Laplaze et al., 2005). Activator lines were selected based on their *GFP* reporter expression pattern and their insensitivity to exogenous cytokinin to avoid a feedback response due to *IPT* expression and cytokinin accumulation. An overview of the lines and their expression pattern is summarized in Table 1. Detailed root phenotypes, including root growth, LRI, and development were analyzed in the F1 generation and, as control, activator lines were crossed with the wild type Columbia-0.

All the lines with *IPT* expression in the root differentiation and early elongation zones, including epidermis/cortex (*J2601>>IPT*, *N9193>>IPT*, *J2092>>IPT*, *M0028>>IPT*, and *J0951>>IPT*), cortex/endodermis (*N9094>>IPT*), provasculature (*J1701>>IPT*), or pericycle (*J0121>>IPT*) exhibited inhibited primary root growth (Figures 3 and 4A, Table 1). By contrast, increased *IPT* expression in the mature differentiated root zones (*J0671>>IPT* and *J3611>>IPT*) did not interfere with root elongation (Figures 3 and 4A, Table 1).

Next, we examined the effect of tissue-specific activation of IPT expression on LRI. In all the cases in which IPT was expressed in the distal root zones of the basal meristem encompassing the epidermis/cortex cells (J2601>>IPT, N9193>>IPT, J2092>>IPT, and M0028>>IPT), cortex/endodermis (N9094>>IPT and N9193>>IPT), or provasculature (J2351>>IPT and J1701>>IPT), LRI was inhibited significantly (Figures 3 and 4B, Table 1; see Supplemental Figure 5 online). By contrast, IPT expression in the mature differentiated root zone (J0671>>IPT and J3611>>IPT) did not affect LRI (Figures 3 and 4B, Table 1; see Supplemental Figure 5 online). Surprisingly, the line driving IPT expression directly in the differentiated xylem pole pericycle cells (J0121>>IPT) exhibited a reduced total number of LRP per root (see Supplemental Figure 5 online) but did not exhibit dramatically altered LRI density (LRP per cm of root length; Figure 4B), despite an almost 40-fold increase of free trans-zeatin and a large increase of other cytokinin metabolites, as detected in J0121>>IPT roots (Figure 4C). Detailed analysis of the J0121 expression pattern revealed that under our conditions, the earliest expression could be detected in pericycle cells that had already left the basal meristem (Figure 3). Thus, condition-dependent variability of expression patterns might be the reason for the less efficient inhibition of LRI compared with previous reports (Laplaze et al., 2007).

Notably, in the lines in which the inhibition of LRI was the strongest (N9094>>IPT, N9193>>IPT, J2351>>IPT, and J2092>>IPT) (Figure 4B), the IPT gene was not necessarily expressed at the early stages of LRP, suggesting that cytokinin might act prior to the asymmetric anticlinal division events to inhibit LRP formation. Accordingly, line J1103, exhibiting IPT activation in primordia from stage I on, was not affected in LRI (Figure 4B, Table 1; see Supplemental Figure 5 online). Similarly, lines with IPT expression in the mature part of the root (J0671>>IPT and J3611>>IPT), in which mostly postinitiation phases of lateral root organogenesis take place, did not show any initiation defects (Figures 3 and 4B).

In summary, cytokinin inhibited LRI most efficiently before the morphologically visible initiation events, in the zone encompassing the basal meristem, but did not interfere with LRI when

Table 1. Overview of the GAL4-GFP Enhancer Trap Line Expression

Organ		J2601	N9193	J2092	N9094	M0028	J0121	J0951	J2351	J1701	J1103	J0671	J3611
Root m	eristem												
	Col	N.E.	N.E.	xxx N.E.		XXX	N.E.	N.E.	XXX	XXX	N.E.	N.E.	X
	Pvas	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	XXX	XXX	N.E.	N.E.	N.E.
	En	N.E.	XX	N.E.	XXX	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.
	С	N.E.	XXX	N.E.	XXX	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.
	Ep	N.E.	XXX	xxx N.E.		XX	N.E.	xx	N.E.	N.E.	N.E.	N.E.	N.E.
	Lrc	XX	N.E.	XX	N.E.	N.E.	N.E.	xx	N.E.	N.E.	XX	N.E.	N.E.
Basal n	neristem												
	Pvas	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	XX	x-fl	N.E.	N.E.	N.E.
	Per	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.
	En	N.E.	XX	N.E.	XXX	N.E.	N.E.	N.E.	N.E.	N.E.	Х	N.E.	N.E.
	С	XX	XXX	N.E.	XXX	Х	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.
	Ep	XX	XXX	xxx N.E.		XXX	N.E.	xx	N.E.	N.E.	N.E.	N.E.	N.E.
Develop	omental w	indow											
•	Vasc	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.	Х	X	N.E.	N.E.	N.E.
	Per	N.E.	N.E.	N.E.	N.E.	N.E.	xxx	N.E.	N.E.	N.E.	N.E.	N.E.	N.E.
	En	XX	XX	Х	XXX	N.E.	N.E.	xxx	N.E.	N.E.	XX	N.E.	N.E.
	С	XX	XXX	X	XXX	XXX	N.E.	xxx	N.E.	N.E.	N.E.	N.E.	N.E.
	Ep	XX	XXX	xxx N.E.		XXX	N.E.	xx	N.E.	N.E.	N.E.	N.E.	N.E.
Mature	root												
	Vasc	N.E.	N.E.	N.E	N.E	N.E.	N.E.	N.E.	XXX	N.E.	N.E.	N.E.	N.E.
	Per	XXX	N.E.	N.E	N.E	N.E.	xxx	xx	N.E.	N.E.	N.E.	XX	N.E.
	En	N.E.	XX	XX	XXX	N.E.	N.E.	xx	N.E.	N.E.	XX	XX	Xxx
	С	XXX	XXX	XX	XXX	xxx	N.E	xx	N.E.	N.E.	N.E.	XXX	Xx
	Ep	XXX	XXX	xxx N.E.		xxx	N.E	xx	N.E.	XX	N.E.	х	N.E.
LRP	•												
		I	N.E.	II on	Em	I on	Up to III	Up to III	IV on	II on	I on	N.E.	N.E.

C, cortex; Col, columella; En, endodermis; Em, emergence; Ep, epidermis; Lrc, lateral root cap; Per, pericycle, Pvas, provasculature; Vasc, vasculature; fl, floem; N.E., not expressed; x, expressed (number of crosses reflects expression intensity).

increased in the mature part of the root or at early stages of LRP. Notably, in no line did ectopic *IPT* expression fully inhibit LRI, suggesting that mechanisms regulating LRI might be quite robust and markedly resist the cytokinin inhibitory stimulus.

Cytokinin Affects LRP Development in a Stage-Specific Manner

Next, we analyzed the impact of the tissue-specific IPT expression on LRP development. When IPT was expressed prior to LRI (N9193>>IPT and N9094) and/or at the very early stages of LRP development (J2092>>IPT, J2601>>IPT, J0951>>IPT, M0028>>IPT,, J0121>>IPT, and J1103>>IPT) (see Supplemental Figure 6 online), the inhibition was dramatic, whereas the developmental defects were less strong (J1701>>IPT and J2351>>IPT) when IPT was expressed at later developmental stages (from stage V on). In the endodermis and cortex of mature roots (J0671>>IPT and J3611>>IPT), the activated IPT expression affected primordium development only moderately and delayed LRP outgrowth (see Supplemental Figure 6 online). Accordingly, an overall decrease in cytokinin levels in the ipt3 ipt5 ipt7 loss-of-function mutants enhanced LRP development and increased the proportion of emerged LRP (see Supplemental Figure 3E online).

To follow and evaluate the short-term cytokinin effects on primordium development, we established time-lapse experiments. Primordia at defined developmental stages were exposed to 2 μM cytokinin and followed for 12 h. Typically, under control conditions, 90% of the LRP progressed from stage I/II (Malamy and Benfey, 1997) to stage III/IV (n = 20; Table 2). After cytokinin treatment, 90% of the LRP were arrested (n = 20). Although in 20% of the LRP one to two divisions could still be detected, the primordia did not progress into the next developmental stage. Similarly, when applied to LRP at stage III/IV, cytokinin dramatically inhibited their development, whereas, upon mock treatment, LRP from stage III/IV progressed to developmental phase VII/ emergence (n = 14) and 70% of the cytokinin-treated primordia were fully arrested in their development (n = 20; Table 2). However, the behavior differed when LRP at stage V/VI were exposed to cytokinin: Under control conditions, such primordia rapidly developed and emerged through the cortex (n = 12; Table 2), but application of cytokinin did not cause an arrest and, consequently, all primordia continued to develop (n = 10; Table 2). Altogether, these data show that increased cytokinin levels prior to initiation and during early developmental phases have the most pronounced impact on the progress of LRP organogenesis, in agreement with previously published data (Laplaze et al., 2007).

A graded distribution of auxin with a maximum at the primordium tips is crucial for proper LRP organogenesis, and, during primordium formation, the auxin maximum is gradually established as the result of a coordinated modulation of the auxin flux toward the tips of newly formed primordia (Benková

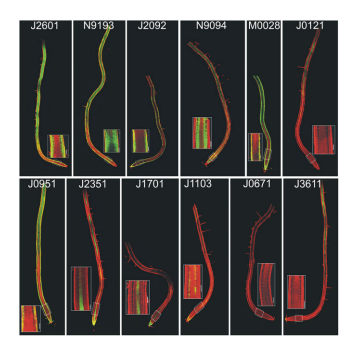


Figure 3. Expression Pattern of GAL4-GFP Enhancer Trap Lines.

Activator lines exhibit expression in the root differentiation and early elongation zones, including epidermis/cortex (J2601, N9193, J2092, M0028, and J0951), cortex/endodermis (N9094), provasculature (J1701), or pericycle (J0121), while in lines J0671 and J3611, expression occurs in the mature differentiated root zones. Insets display enlarged basal meristems. Bars = $50~\mu m$.

et al., 2003; Kleine-Vehn et al., 2009). Therefore, we tested whether cytokinin affected LRP development by influencing the auxin response. In vivo time-lapse monitoring of the *DR5:RFP* (for red fluorescent protein) auxin reporter during LRP organogenesis revealed that cytokinin rapidly reduced the auxin response and prevented auxin accumulation at the LRP tips when applied on the primordia at the early developmental phases, whereas older LRP with established *DR5* maxima at the tips were more resistant to cytokinin treatment (Figures 5A versus 5B). These results indicate that the cytokinin might have a stronger impact on the LRP in the early phases without established auxin maxima than on the developmentally more progressed LRP with an established and stabilized auxin gradient.

DISCUSSION

Auxin and Cytokinin Activities during Lateral Root Organogenesis

The activity of the auxin signaling pathway and the auxin distribution along the root determines stem cell niche and root meristem activity (Sabatini et al., 1999; Blilou et al., 2005), and recurrent local accumulation of auxin in protoxylem cells in the basal meristem has been linked to priming (De Smet et al., 2007). Auxin plays a role in founder cell specification, LRP initiation (Dubrovsky et al., 2008), primordium patterning and

development (Benková et al., 2003), and noninvasive emergence of primordia through surrounding tissues (Swarup et al., 2008). In addition, auxin minima have been linked with the developmental window (Dubrovsky et al., 2011). In general, auxin and its proper distribution have been correlated with organ initiation and proper organ patterning.

Cytokinin has a function in the regulation of root meristem maintenance (Dello loio et al., 2007, 2008) and lateral root organogenesis (Li et al., 2006; Laplaze et al., 2007). Interestingly, monitoring of tissues and cells exhibiting enhanced TCS reporter expression has revealed that cytokinin and auxin response maxima scarcely overlap within the root. For example, whereas DR5 is detected in pericycle founder cells (Dubrovsky et al., 2008), the TCS reporter is expressed in pericycle cells between two, or across, existing LRP. Cytokinin response maxima in the pericycle are strongly attenuated in young roots or roots without primordia because of inhibited auxin transport, linking cytokinin activity in the pericycle with LRP organogenesis. Perturbations in cytokinin signaling in arr1 arr11 lossof-function mutants increased the frequency of abnormally positioned LRP, hinting at a role for cytokinin in preventing LRI near existing LRP. Accordingly, defects in LRP positioning occur in lines overexpressing CKX (Werner et al., 2003). Reduction in cytokinin levels and responsiveness in pericycle cells significantly enhanced the sensitivity to LRI-triggering signals, such as auxin, of pericycle cells in the proximal root zone when compared with controls. At this point, we can only speculate about the mechanism that locally stimulates cytokinin responses in LRP in the vicinity of pericycle cells. Possibly, cytokinin production itself in the primordia might be triggered and diffused laterally afterward. Alternatively, an originally noncytokinin signal could be released from primordia, stimulating cytokinin production and activity in pericycle cells in the vicinity of LRPs. Complex expression patterns of a whole set of genes involved in cytokinin metabolism, such as IPT (Miyawaki et al., 2004), the cytokinin-activating gene LONELY GUY (Kuroha et al., 2009), or CKX (Werner et al., 2003), imply that levels of active cytokinin might be under rapid tissue-specific metabolic regulation.

Besides in the pericycle, in other root tissues, auxin and cytokinin response maxima are also rather complementary. In the primordia, *DR5* is typically expressed at the primordium tips from the early stages on (Benková et al., 2003), whereas the cytokinin response is strongly suppressed during the early stages of development and is detectable only in provascular cells of emerging LRP. Similarly, in the primary root tip, *DR5* expression exhibits response maxima in the central columella cells, whereas *TCS* occupies the expression domain in the outer columella cells (Müller and Sheen, 2008). Enhancement of the auxin maxima by chemical treatment shifts the cytokinin response zone toward the outer columella cells, suggesting that both pathways might mutually delimit their activities.

Role of Cytokinin in the Spatiotemporal Control of LRI

The acropetal architecture of the root system in *Arabidopsis* results from initiation events limited to a restricted zone behind the root tip in the simultaneously growing primary root. As a consequence, new initiation events occur distally from the

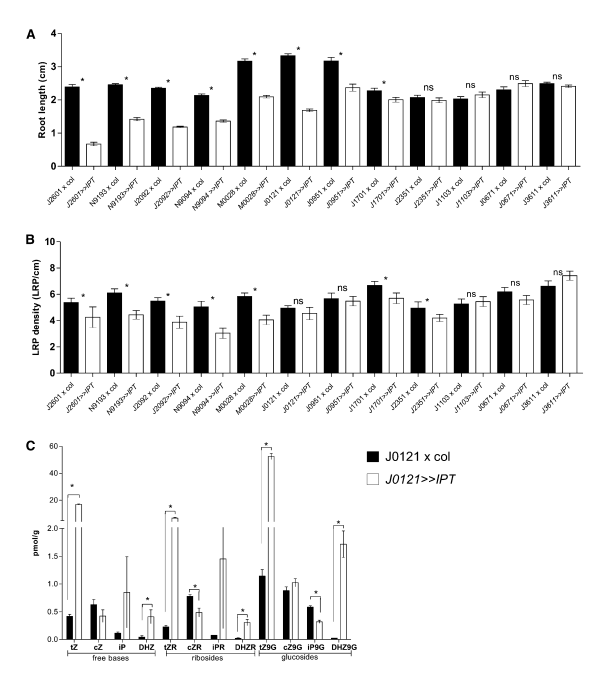


Figure 4. Spatiotemporal Effect of IPT Expression on Root Growth and LRI.

(A) Inhibited root growth in lines with *IPT* expressed in the root differentiation and elongation zones (*J2601>>IPT*, *N9193>>IPT*, *J2092>>IPT*, *N9094>>IPT*, *M0028>>IPT*, *J0121>>IPT*, and *J1701>>IPT*). Error bars denote se (*P < 0.05; Mann-Whitney test, *n* = 10 roots).

(B) LRP density in lines with *IPT* expression in the basal meristem. The lines *J2601>>IPT*, *N9193>>IPT*, *J2092>>IPT*, *N9094>>IPT*, *M0028>>IPT*, *J1701>>IPT*, and *J2351>>IPT* exhibit reduced LRP density. Error bars denote se (*P < 0.05; Mann-Whitney test; *n* = 10 roots). ns, not significant.

(C) Cytokinin levels in control *J0121>>Col* and *J0121>>IPT* roots. Error bars denote se (*P < 0.05; Student's *t* test; *n* = 3).

cZ, *cis*-zeatin; cZ9G, *cis*-zeatin-9-glucoside; cZR, *cis*-zeatin riboside; DHZ, dihydrozeatin; DHZ9G, dihydrozeatin; tZ9G, *trans*-zeatin-9-glucoside; tZR, *trans*-zeatin riboside.

MS		Cytokinin (2 µM Benzyl Adenine)				
Progress of LRP Development After Application	No. of Events	Progress of LRP Development after Application	No. of Events			
From I to I	1	From I to I	12			
From I to II	3	From II to II	7			
From I to III	5	From II to III	1			
From I to IV	1	From I to IV	0			
From I to V	1	From I to V	0			
From II to II	1	From II to II	0			
From II to III	1	From II to III	0			
From II to IV	5	From II to IV	0			
From II to V	1	From II to V	0			
From II to Em	1	From II to Em	0			
Fully arrested/total	2/20	Fully arrested/total	19/20			
From III to III	1	From III to III	7			
From III to V	1	From III to IV	2			
From III to Em	2	From IV to IV	7			
From IV to IV	1	From IV to V	1			
From IV to VII	3	From IV to E	3			
From IV to Em	6	From IV to Em	0			
Fully arrested/total	2/14	Fully arrested/total	14/20			
From V to VI	1	From V to VII	1			
From V to VII	0	From V to Em	7			
From V to Em	5	From VI to VII	1			
From VI to Em	2	From VI to Em	1			
Fully arrested/total	0/8	Fully arrested/total	0/10			

previous ones toward the root apex. Although the mechanisms ensuring such an initiation pattern are still not understood, it might be anticipated that to restrict initiation to a certain root zone, pericycle cells are regulated spatiotemporally at a stage competent for cell divisions and/or recurrent LRI-inducing signals. Both mechanisms might indeed contribute to early initiation events. Previous research on the characterization of xylem pole pericycle cells (Beeckman et al., 2001) and on the identification of the priming zone (De Smet et al., 2007; De Rybel et al., 2010) and developmental window (Dubrovsky et al., 2006) suggest that during primary root growth, pericycle cells in a certain root zone are maintained in an LRI-competent stage. In addition, recurrent enhancement of auxin signals contributes to the initiation of the lateral root developmental program (De Smet et al., 2007; Moreno-Risueno et al., 2010).

Em, emergence.

In view of its inhibitory effect on LRI, the overall cytokinin activity might be expected to be attenuated in the zone where the earliest phases of LRI take place. Indeed, as demonstrated by the *TCS* reporter analysis, the cytokinin response is minimal in the priming and developmental window zones. Interestingly, analysis of cytokinin derivatives has not revealed any dramatic decrease in cytokinin levels in the parts encompassing the priming zone and the developmental window. Although detection of such a tissue-specific reduction in cytokinin levels might be under the resolution limit of the analytical approach used, the observation that exogenous cytokinin could not enhance the *TCS* reporter expression suggests that an important part of the regulation mechanism strongly suppresses the cytokinin signaling pathway.

LRI is inhibited severely by an increase in cytokinin activity in the zone encompassing the basal meristem and the developmental window, but elevated cytokinin levels in more distant root zones are insufficient, although they still interfere with lateral root organogenesis. These data suggest that cytokinin might act prior to any visible signs of LRP initiation to counteract positive LRI

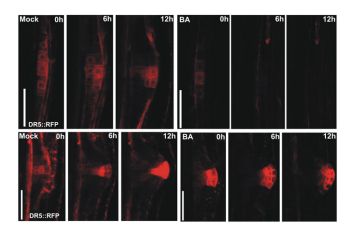


Figure 5. Cytokinin Effects on LRP Development.

Real-time monitoring of LRP development on control medium (Mock) and in the presence of 0.1 μ M benzyl adenine, when cytokinin was applied at stage I **(A)** or stage IV **(B)**. *DR5:RFP* reporter was used to visualize auxin maxima and developmental stages of LRPs. Bars = 50 μ m.

signals. Moreover, cytokinin inhibition of LRI was in most of the cases accompanied by root growth reduction, indicating tight interconnection between these two developmental processes. Interestingly, despite the strong *IPT* overexpression, none of the ectopic increases in endogenous cytokinin levels in defined root zones fully prevented LRI, implying that LRI-regulating mechanisms markedly resist the inhibitory effects of cytokinin.

Role of Cytokinin in the Regulation of Early LRP Development

Overall, the cytokinin response, as monitored with the *TCS* reporter, was strongly repressed during the early phases of lateral root organogenesis and detected in the primordium provasculature only later in older primordia. Increased cytokinin responses during the early phases of lateral root organogenesis, including priming, either by endogenous expression of *IPT* or exogenous treatment, revealed that primordia are more sensitive to increased cytokinin levels in the early phases of development than in the later phases. Similar developmental phase-dependent cytokinin sensitivity has been reported previously, demonstrating that endogenous cytokinin levels fluctuate during the early stages of primary root development and that a developmental window (between 2 and 3 d after germination) specifies the increased sensitivity of the root toward endogenous cytokinin (Kuderová et al., 2008).

During primordium formation, an auxin maximum is gradually established at the primordium tips, as a result of the coordinated repolarization of the auxin flux toward the tips of newly formed primordia (Benková et al., 2003; Kleine-Vehn et al., 2009). The auxin gradient formation is critical for the proper development of the primordia, and genetic or chemical interference with their auxin distribution results in defective primordium organogenesis (Benková et al., 2003). Spatiotemporal manipulation of cytokinin levels at defined stages of primordium development revealed that enhancement of the cytokinin activity during the early phases strongly interferes with the auxin gradient formation, whereas at the later stages, cytokinin only mildly affects the established auxin gradients. Thus, stage-dependent effects of cytokinin on primordium development might rely on the robustness and stability of the possibly higher auxin gradient in more developed primordia.

METHODS

Plant Lines and Growth Conditions

The transgenic *Arabidopsis* thaliana lines were *DR5:GFP*, *DR5:3XVENUS-N7* (Heisler et al., 2005), and *DR5:RFP* (Marin et al., 2010). The *GAL4-GFP* enhancer trap lines *J2601*, *N9193*, *J2092*, *N9094*, *M0028*, *J0951*, *J1701*, *J0671*, *J1103*, *J2351*, and *J3611* and the cytokinin signaling mutant *arr1-3 arr11-2* (N6980) were obtained from the Nottingham Arabidopsis Stock Centre (http://nasc.nott.ac.uk/). The mutant line *ipt3 ipt5 ipt7* had been described previously (Miyawaki et al., 2006) as well as the *ProUAS:IPT*, *J0121*, *TCS:GFP*, and *TCS:LUC* lines (Laplaze et al., 2007; Müller and Sheen, 2008). The GAL4 enhancer trap lines were crossed with *ProUAS:IPT* and the wild type Columbia-0. The F1 generation was analyzed. Seeds were sterilized with chloral gas, plated in Petri dishes on 0.8% agar 0.5× Murashige and Skoog (MS) medium

containing 1% Suc, stored for 2 d at 4°C, and grown vertically at 18°C under a long-day photoperiod. Seedlings were harvested and processed 3, 4, or 7 d after germination.

Phenotypic Analysis, Microscopy, and Statistics

For the confocal laser scanning microscopy, a TCS SP2 AOBS microscope (Leica) was used. Seedlings were stained for 5 min in 0.01 mg mL $^{-1}$ propidium iodide. Images were processed in Adobe Photoshop. Plant material was cleared as described (Malamy and Benfey, 1997). Root lengths were measured on scanned slides with the ImageJ software (NIH). LRPs were counted with a differential interference contrast microscope (BX51; Olympus). At least 15 seedlings were analyzed, and experiments were repeated twice independently. LRP density was calculated as the number of LRP per total root length. The Excel statistical package (Microsoft) and GraphPad Prism software were used for data analysis, and Mann-Whitney nonparametric and Student's t tests were applied.

For real-time analysis of LRP development, 6-d-old seedlings were placed in chambered cover glasses (Nunc Lab-Tek) and covered with 0.2-mm-thin square blocks of solid MS (mock) media with or without cyto-kinin. LRPs were scanned at 3- or 5-min time intervals for 8 to 12 h by a confocal microscope (FV10 ASW; Olympus). The frequency of the positioning-defective LRPs was calculated as the percentage of primordia initiated at distances shorter than those of the control situation from the total initiation events per root.

For the phenotypic analysis after auxin treatment, 7-d-old seedlings grown on MS medium or 10 μM NPA were treated with liquid MS medium with or without 1 μm NAA for 25 h. Seedlings were cleared and the lateral root number was scored in the zones of LRP initiation, formation, and LRP emergence (see below), and the ratio was calculated between the LRP induced by auxin and during incubation in MS medium.

Cytokinin Measurements

Approximately 150 mg of plant tissue was ground in liquid nitrogen and 1 pmol of deuterium-labeled cytokinin internal standards was added to each sample prior to the analysis (Olchemin). After overnight extraction, the samples were purified with a cation exchanger (SCX cartridge; Agilent) and by immunoaffinity extraction with a broad-spectrum monoclonal anticytokinin antibody (Novák et al., 2003). The eluates from the immunoaffinity columns were evaporated to dryness and redissolved in 20 μL of the initial mobile phase. Cytokinins were separated by ultraperformance liquid chromatography (Acquity UPLC; Waters) on a C18 reversed-phase chromatographic column (BEH 18; 1.7 μm; 2.1 × 50 mm; Waters). The mobile phase consisted of 15 mM ammonium formate, pH 4 (A) and methanol (B). For the cytokinin separation, a gradient elution was used (0 min, 10% B; 0 to 8 min, 50% B; flow rate 0.25 mL min⁻¹) with a column temperature of 40°C. The column eluate was introduced into a triple-stage quadrupole mass spectrometer (Xevo TQ MS; Waters), quantified by multiple reaction monitoring of [M+H]+ and the appropriate product ion by Masslynx software (Waters) with a standard isotope dilution method (Novák et al., 2003). The optimal dwell time, cone voltage, collision energy, and exact transition for each cytokinin had been optimized previously (Novák et al., 2008).

For cytokinin measurements in the zones defined by the developmental phases of lateral root organogenesis, namely LRP initiation, formation, and LRP emergence through the cortex, the borders between the three zones were recognized under a confocal microscope for five seedlings 7 d after germination. The zone lengths were measured and an average value was used. The zone of the LRI corresponded to $\sim\!\!6$ mm from the root tip, the zone between 6 and 12 mm from the root tip to the LRP formation zone and above 12 mm from the root tip to the lateral root emergence zone.

Fluorescence-Activated Cell Sorting of Plant Protoplasts

Cells were sorted as described (Birnbaum et al., 2005). Approximately 5000 seeds (per replicate) of each GFP line used in the experiments were sterilized with bleach and plated on MS media. To allow rapid harvesting, seeds were arranged in rows on square plates at a density of \sim 500 seeds per row on top of nylon mesh (Nitex 03 100/47; Sefar). Five-day-old roots were cut off \sim 1 cm from their tip. Dissected roots were placed in protoplasting solution B (50 mL of solution A [see below], 750 mg cellulase, and 50 mg pectolyase) inside 70- μ m cell strainers placed in small Petri dishes and incubated for 1 h at room temperature with agitation. Protoplasted cells were collected from Petri dishes and concentrated by spinning down (at \sim 120 relative centrifugal force). The supernatant was aspirated and the cell pellet was resuspended in 1.5 mL of solution A (2 mL of 1 M KCl 1 M, 400 μ L of 1 M MgCl₂, 400 μ L of 1 M CaCl, 0.2 g BSA, 78 mg MES, and 21.86 g mannitol; final volume, 200 mL, pH 5.5, with 1 M Tris).

GFP-expressing cells were isolated on a fluorescence-activated Epics Altra cell sorter (Beckman Coulter) fitted with a 100-μm nozzle at a rate of 2000 to 5000 events per second and a fluid pressure of 30 p.s.i. Protoplasts from Columbia wild-type plants that did not express *GFP* were used as a negative control for establishing sorting criteria based on the following cell properties: live protoplast clusters with intact membranes were selected based on a high forward to side scatter ratio and GFP-positive cells by their emission intensity in the green channel (\sim 530 nm) above the negative controls. Cells were sorted directly into the RLT lysis buffer (Qiagen), mixed, and immediately frozen at -80° C for later RNA extraction. An autofluorescence filter was established by eliminating cells that fluoresced at an equal intensity in the green and orange (\sim 575 nm) channels.

RNA Extraction of Sorted Cells and Quantitative RT-PCR

RNA was extracted with the RNeasy Micro kit (Qiagen) from sorted cells. A DNase treatment with the RNase-free DNase Set (Qiagen) was performed for 15 min at 25°C. Poly(dT) cDNA was prepared from 1 μg total RNA with the iScript cDNA synthesis kit (Bio-Rad) and quantified with a LightCycler 480 (Roche) and SYBR Green I Master (Roche) according to the manufacturer's instructions. PCR was performed in 384-well optical reaction plates heated for 10 min to 95°C to activate the hot-start Taq DNA polymerase, followed by 40 cycles of denaturation for 60 s at 95°C and annealing/extension for 60 s at 58°C. Targets were quantified with specific primer pairs designed with the Beacon Designer 4.0 (Premier Biosoft International). Expression levels were normalized to UBIQUITIN10 (UBQ10) expression levels. All qRT-PCR experiments were done at least in triplicate. The following primers were used: 5'-CACACTCCACT TGGTCTTGCGT-3' and 5'-TGGTCTTTCCGGTGAGAGTCTTCA-3' for UBQ10; 5'-TTGAAGAAACCGCGTGTCGTCT-3' and 5'-CCTTCTCAACG CCGAGCTGATTAA-3' for ARR1; and 5'-TAATGATGTCGGTGGACGGC GAAA-3' and 5'-AAGCTCCGTGTTGCACTCCCTTCA-3' for ARR11.

Transient Expression in Protoplasts

The DR5rev promoter was amplified from the original plasmid (Friml et al., 2003) using the primers DR5_FOR (5′-ATCCCCCGACGGTATCGCAGCC-CAGG-3′) and DR5_REV (5′-TGTAATTGTAATTGTAAATAGTAATTG-3′). The amplified fragment was extracted and purified from gel and cloned in the pENTR5′-TOPO vector using the TA Cloning kit (Invitrogen) according to manufacturer's instructions. The *DR5:LUC* construct was created by combining pEN-L1-L-L2 and an entry clone holding the DR5 sequence combined by a MultiSite Gateway LR reaction with pm42GW7 as destination vector. *TCS:LUC* was created by PCR amplification of the TCS, minimal 35S promoter, and TMV Ω sequence fragment from TCS:GFP (Müller and Sheen, 2008), using the primers 5′-GTGCCAAGCTTTGCTAGCAAAATC-3′ and 5′-TGTAGGCCTGTAATTGTAATTGTAAATAG-3′. The resulting fragment was digested with Stul and HindIII. A second PCR amplification including the

LUC-coding sequence and nopaline synthase 3'-untranslated region sequence of GCC:LUC:NOS was performed using the primers 5'-ACAGGC CTACATGGAAGACGCCAAAAACATAAAG-3' and 5'-TACGAATTCTCATG TTTGACAGCTTATCATC-3'. The resulting fragment was digested with Stul and EcoRl, and both digested PCR fragments were then ligated into the HindIII-EcoRI-digested pUC18 to result in TCS:LUC.

Mesophyll protoplasts were isolated from rosette leaves of 4-week-old Arabidopsis plants grown on soil under controlled environmental conditions in continuous light at 22°C (arr1 and arr11 versus wild-type control analysis) or in a 16-h/8-h light/dark cycle (hormone treatments for TCS and DR5 reporter genes). Protoplasts were isolated as described (Wu et al., 2009) with modifications (Wehner et al., 2011): The surface of Arabidopsis leaves was cut with a razor blade and placed in a Petri dish with 5 mL enzyme solution (1% cellulase Onozuka R10 [Yakult], 0.3% macerozyme Onozuka R10 [Yakult], 0.4 M mannitol, 20 mM KCl, 10 mM CaCl₂, and 20 mM MES, pH 5.7). After a 4-h digestion, the released protoplasts were filtered into a 50-mL tube and collected by centrifugation. The supernatants were discarded and the protoplasts were resuspended in 5 mL W5 (154 mM NaCl, 125 mM CaCl₂, and 5 mM KCl). The protoplasts were counted with a hemocytometer under a light microscope and kept on ice for at least 30 min. For polyethylene glycol transfection, protoplasts were adjusted to a final concentration of 3 imes 10^5 mL with MMG solution (0.4 M mannitol, 15 mM MgCl₂, and 4 mM MES, pH 5.7) (Yoo et al., 2007). Protoplasts (3 \times 10⁴ cells) were cotransfected with 20 µg of a reporter plasmid containing the LUC reporter gene and 2 µg of a normalization construct expressing Renilla LUC (rLUC). Transfections were done by adding an equal volume of a freshly prepared solution of 40% (w/v) polyethylene glycol 4000 (Fluka) with 0.2 M mannitol and 0.1 M CaCl₂. After 10 min of incubation, 0.44 mL W5 solution was added and mixed by pipetting. Transfected protoplasts were collected by centrifugation and gently resuspended in 50 μ L W5 solution (for hormone treatments, 0.5 μ M benzyl adenine and/or 1 μM NAA was added to the buffer), then transferred to 12-well plates coated with 1% BSA and incubated during 16 h in darkness. Protoplasts were collected and LUC was measured with the dual luciferase reporter assay system (Promega). Luciferase cell culture lysis reagent (Promega) was added to the protoplasts and mixed by pipetting. After 5 min of incubation on ice, 100 µL lysate was used to measure the LUC activity according to the manufacturer's protocol. The mean value ($\pm sd$) was calculated from two measurements, and each experiment was repeated at least three times. To normalize the values, rLUC was used.

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: AT3G63110.1 (IPT3), AT5G19040.1 (IPT5), AT3G23630.1 (IPT7), AT3G16857.1 (ARR1), AT1G67710.1 (ARR11), and EF090415 (GCC1: LUC:NOS).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Spatial Distribution of Cytokinin Response along the Root.

Supplemental Figure 2. Xylem Pole Pericycle Cell Identity in Control and NPA-Treated Roots.

Supplemental Figure 3. LRP Initiation Spacing Defective in Roots with Compromised Cytokinin Responses.

Supplemental Figure 4. Cytokinin and Auxin Response Distribution in Roots

Supplemental Figure 5. Effect of IPT Expression on LRI.

Supplemental Figure 6. Spatiotemporal Effect of *IPT* Expression on LRP Development.

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AUTHOR CONTRIBUTIONS

A.B. and E.B. designed the project. A.B., K.P., P.M., J.D., C.C., B.M., W.G., and P.T. performed the experiments and analyzed data. A.B. and E.B. wrote the article.

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Spatiotemporal Regulation of Lateral Root Organogenesis in Arabidopsis by Cytokinin

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