ARABIDOPSIS HOMOLOG of TRITHORAX1 (ATX1) is required for cell production, patterning, and morphogenesis in root development

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Abstract

ARABIDOPSIS HOMOLOG of TRITHORAX1 (ATX1/SDG27), a known regulator of flower development, encodes a H3K4-histone methyltransferase that maintains a number of genes in an active state. In this study, the role of ATX1 in root development was evaluated. The loss-of-function mutant atx1-1 was impaired in primary root growth. The data suggest that ATX1 controls root growth by regulating cell cycle duration, cell production, and the transition from cell proliferation in the root apical meristem (RAM) to cell elongation. In atx1-1, the quiescent centre (QC) cells were irregular in shape and more expanded than those of the wild type. This feature, together with the atypical distribution of T-divisions, the presence of oblique divisions, and the abnormal cell patterning in the RAM, suggests a lack of coordination between cell division and cell growth in the mutant. The expression domain of QC-specific markers was expanded both in the primary RAM and in the developing lateral root primordia of atx1-1 plants. These abnormalities were independent of auxin-response gradients. ATX1 was also found to be required for lateral root initiation, morphogenesis, and emergence. The time from lateral root initiation to emergence was significantly extended in the atx1-1 mutant. Overall, these data suggest that ATX1 is involved in the timing of root development, stem cell niche maintenance, and cell patterning during primary and lateral root development. Thus, ATX1 emerges as an important player in root system architecture.

Key words: Arabidopsis, lateral root development, patterning, root apical meristem, root development, root morphogenesis, root system.

Introduction

Gene expression changes to active or repressed states and the maintenance of these changes are essential for the orchestration of any developmental programme in living organisms. Trithorax (TrxG) and Polycomb group (PcG) proteins are important regulatory factors that contribute to the mainte-
nance of gene expression states (Pien and Grossniklaus, 2007; Avramova, 2009; Alvarez-Venegas, 2010; Berr et al., 2011; Schuettengruber et al., 2011). Therefore, these proteins repres-
sent critical dynamic factors that act at the epigenetic level to define cell and tissue identities in both animal and plant species. Numerous chromatin modification factors participate in various developmental processes and play essential roles in root development. Mutants in genes encoding two subu-
nits of CHROMATIN ASSEMBLY FACTOR-1 (CAF-1),
**fasciata1-1 (fas 1-1)** and **fas 2-2**, are characterized by aberrant root apical meristem (RAM) organization. The quiescent centre (QC) of these mutants is either absent or difficult to identify (Kaya et al., 2001). In addition, **FAS2** is involved in trichoblast/atrichoblast cell specification (Costa and Shaw, 2006) and **FASI** in lateral root (LR) initiation (Manzano et al., 2012). A histone deacetylase, HDA18, participates in root epidermal cell patterning (Xu et al., 2005), and a histone acetyltransferase, GCN5 (GENERAL CONTROL NONDEPRESSIBLE 5), and the GCN5-associated factor, ADA2b (ALTERATION/DEFICIENCY IN ACTIVATION 2B), act in the PLETHORA pathway and are essential for maintaining RAM activity (Kornet and Scheres, 2009). Furthermore, a SWI2/SNF2 chromatin-remodelling ATPase, BRAHMA, regulates primary root growth in an ABA-dependent manner (Han et al., 2012). **PICKLE** (PKL), a chromatin-remodelling factor with the chromodomain/helicase/DNA-binding domain (CHD3/CHD4), is required for the transition from the embryonic stage to post-embryonic development (Ogas et al., 1999), for repression of LR initiation through auxin-dependent negative regulation of pericycle cell activation (Fukaki et al., 2006), and for RAM maintenance (Aichinger et al., 2011). RAM activity is regulated by the antagonistic activity between a PcG protein, CURLY LEAF (CLF), and PKL (Aichinger et al., 2011).

TrxG and PcG proteins are involved in maintaining the active and repressed states of genes with antagonistic functions (Köhler and Hennig, 2010). Most TrxG proteins exert their function as part of large multimeric protein complexes that have either histone-modifying or nucleosome-remodelling activities. Thus, TrxG proteins are involved in the formation of an open chromatin structure and facilitate transcription by being involved in chromatin remodelling (Breiling et al., 2007). The best-studied member of this group in plants is **ARABIDOPSIS HOMOLOG OF TRITHORAX1 (ATX1/SDG27)**, which is a member of the SET DOMAIN GROUP (SDG) family of genes and encodes a H3K4-histone methyltransferase. **ATX1** participates in flower development by activating flower homeotic genes (Alvarez-Venegas et al., 2003; Pien et al., 2008). However, little is known about the role of TrxG genes in root development, and only recently was it shown that **SDG2**, which encodes another H3K4-histone methyltransferase, is required for root growth and root stem cell niche maintenance (Yao et al., 2013). Here it is shown that **ATX1** is essential for proper RAM organization, RAM activity, and, subsequently, for cell production. The morphogenesis of lateral root primordia (LRPs) of the **atx1-1** loss-of-function mutant was affected at both early and later developmental stages. These data suggest that this TrxG gene is required for cell proliferation-related processes, cell patterning, and morphogenesis of the root.

**Materials and methods**

**Plant materials and growth conditions**

Arabidopsis thaliana (L.) Heynh wild type (Wt) and the **atx1-1** mutant were in the Wassilewskija (Ws) ecotype. The isolation and shoot phenotype of the **atx1-1** mutant have been described (Alvarez-Venegas et al., 2003). Transgenic marker lines **pWOX5::GFP** (Sarkar et al., 2007), **pSCR::GFP** (Heidstra et al., 2004), **pDR5rev::GFP** (Friml et al., 2003), and **Cyclin B1;1::GUS** (Colón-Carmona et al., 1999) have also been described. Seeds were sterilized for 10 min in 20% commercial bleach and 0.08% Triton X-100, washed four times with sterile distilled water, and imbibed at 4 °C for 2 d. Plants were grown in soil (Metromix 200) or in Petri dishes oriented vertically and containing 0.2× Murashige and Skoog (MS) medium prepared from Linsmaier and Skoog medium (L477; Phyto Technology Laboratories, Lenexa, KS, USA), pH 5.7, and supplemented with vitamins (0.1 mg l⁻¹ pyridoxine, 0.1 mg l⁻¹ nicotinic acid, from Sigma-Aldrich, St Louis, MO, USA), 1% sucrose, and 0.8% agar (w/v, Bacto™ Agar; BD Difco, Sparks, MD, USA). All plants were grown at 21 °C, under a 16:8 h light/dark photoperiod with a light intensity of 105 µmol photons m⁻² s⁻¹.

**Auxin treatments and RT-qPCR**

For root treatments and RT-qPCR, Wt and **atx1-1** seedlings were grown for 3 days post-germination (dpg) in vertically oriented Petri dishes containing 0.2× MS medium and then transferred to the same medium or media supplemented with 1 µM or 5 µM indole acetic acid (IAA), and grown for an additional 5 d. For transcript analysis, 7.5 dpg seedlings were treated with 1 µM naphthaleneacetic acid (NAA) for 12 h and then RNA was extracted. IAA and NAA were purchased from Sigma-Aldrich. Total RNA was extracted from roots of Wt and **atx1-1** seedlings using TRIzol reagent (Invitrogen), according to the manufacturer’s instructions. Real-time quantitative reverse transcription–PCR (RT-qPCR) analysis was performed using an iQ5 Multicolor Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA). Reactions were set up using a one-step RT-PCR Kit with SYBR Green (Bio-Rad), according to the manufacturer’s instructions; 100 ng of RNA was used for each reaction, except for the negative control. The primers specific for AuxIAA14 (AT4G14550) were IAA14-Fw CCT CCT GCT AAA GCA CAA GTG and IAA14-Rv CTT CGC CGC TCT TCT GAT TAG C. Data were normalized to the expression of two reference genes, **UBQ10** (At4g05320) and **EF1α** (At5G60390) (Czeckowski et al., 2005), and normalized expression levels were calculated according to Vandesompele et al. (2002). Two biological and six technical replicates were performed.

**Microscopy**

Roots were cleared using an acidified methanol procedure (Malamy and Benfey, 1997) with modifications as described (Dubrovsky et al., 2006), and whole-mount preparations were analysed under a Zeiss Axiosvert 200M microscope (Zeiss, Oberkochen, Germany) equipped with differential interferential contrast (DIC; Nomarski) optics. For β-glucuronidase (GUS) staining, roots were pre-fixed in 0.3% formaldehyde for 20 min at room temperature, washed in 100 mM sodium phosphate buffer, pH 7.4, and stained as described (Hemerly et al., 1993). Photographs were taken using a Photometrics CoolSNAPcf Color Camera (Valley International Corporation, Austin, TX, USA). The roots were fixed in 50% methanol and 10% acetic acid at 4 °C for 5 h and pseudo-Schiff staining was performed as described (Truernit et al., 2008). Then, samples were mounted in NaI-based clearing and mounting solution (Dubrovsky et al., 2009). Live roots were stained with 1 µM or 5 µM propidium iodide dissolved in water. Confocal laser scanning microscopy (CLSM) was performed with a Zeiss LSM 510 Meta (Oberkochen, Germany) microscope using sequential scanning. For the red and green channels, the 543 nm line of a He/Ne laser and the 488 nm line of an Ar laser were used for excitation, respectively. In some instances (indicated in the figure legends), image contrast was improved using the Gaussian blur and Unsharp mask filters in ImageJ (http://rsb.info.nih.gov/ij).
increments measured using ImageJ. LR density, the density of LRPs, length of fully elongated cortical cells, LR initiation index, length of the root apical meristem (RAM), length of the proliferation domain (PD), and the length of the transition domain (TD) were determined on cleared roots as described (Dubrovsky et al., 2009; Dubrovsky and Forde, 2012; Ivanov and Dubrovsky, 2013). Criteria for defining the PD and TD have been described (Ivanov and Dubrovsky, 2013; Lópe-Bucio et al., 2014). Briefly, the PD comprises cells that maintain proliferation activity and the TD comprises cells that have a very low probability of proliferating, but grow at the same rate as cells in the PD and have not yet started elongating rapidly. The domains were determined based on relative changes in the cell lengths observed on cleared root preparations. In the PD, the cell length commonly varies no more than 2-fold, and in the TD, cells are longer than the longest cells in the PD. In the elongation zone (EZ), the cell length starts to increase steadily and simultaneously in all tissues. The point at which this increase can be observed was defined as the distal (rootward) border between the TD and the EZ.

The position of the most distal (rootward) LRP and LR, as well as the number of LRPs in the LR formation and branching zones was determined on cleared root preparations under a microscope equipped with DIC optics. Cortical cell length was determined for 10 cells per root on cleared preparations using an ocular micrometer. The root growth parameters and RAM activity, including cell cycle duration, were evaluated as described (Ivanov and Dubrovsky, 1997; López-Bucio et al., 2014). Briefly, all the parameters were evaluated for each individual root. The cell production rate was calculated based on the rate of root growth and the length of fully elongated cells, and the cell cycle time was evaluated based on cell production and the number of cells in the PD, as described (Ivanov and Dubrovsky, 1997). The number of cells displaced from the cell proliferation domain (Ntrans) during a 24 h period was estimated from the equation Ntrans=24ln2 Npr(Td+Tc+Tr)−1, based on assumptions given in Ivanov and Dubrovsky (1997), where Npr is the number of cells in the PD of the RAM and T is the average cell cycle time (h).

Timing of LR formation
The timing of LR formation, defined as the period from LRPI to LR initiation to LR emergence, was estimated based on the rate of root growth (V, mm h−1) and the length of the LR formation zone (Lp, mm). The latter comprises the portion of the root from the most rootward (i.e. closest to the root tip) LRP to the most rootward emerged LR (Dubrovsky and Forde, 2012). For each individual root, the following data were collected: (i) the root growth increments during the last 3 d; (ii) the distance from the root tip to the most rootward LRP (Lp, mm); and (iii) Lp. The latter two parameters were determined in cleared roots. The measurements were performed using ImageJ. Previously, it was observed that length Lp did not change significantly in seedlings of different ages (JGD, personal observation). Therefore, it was assumed that Lp was the same in a root at the moment of LRP initiation and when the primordium emerged as an LR. The time interval (days) between the two time points was calculated based on the growth increments. Root growth rate (V) during these days was evaluated and LR formation time (Tr, h) was calculated as Tp=VLP−1. When the growth rate was significantly different between the last and the first days of growth recorded, to decrease a calculation error, a fraction of the root portion Lp formed during a certain day of growth was determined and Tp was evaluated as the sum of separately calculated Tp intervals.

LRP symmetry analysis
To estimate the LRP symmetry, only primordia in cleared roots positioned on a slide in the protoxylem plane (i.e. both protoxylem strands were clearly visible in the same focal plane) were analysed (Dubrovsky et al., 2000). The length of the primordium base was measured and did not include pericycle cells at the primordium borders that did not divide periclinally. From the centre of the primordium base line, a perpendicular line was drawn that corresponded to the axis of the prospective LR. From the centre of this axis line, two radii were drawn parallel to the primordium base and measured. If these radii were of equal lengths, the LRP was considered to be symmetrical (asymmetry=0). When the radii were of unequal length, the longer (rL) and shorter (rS) radii were recognized. To estimate the percentage of asymmetry (A) for each primordium, the following equation was used: A=rL−rS(rL+rS)−1 100, where rL is the longer radius length and rS is the shorter radius length. An average of A values was calculated for Wt and atx1-1 LRPs. The statistical analysis was performed using SigmaPlot 12 (Systat Software, San Jose, CA, USA). The number of independent experiments in each case is indicated in the corresponding figure legend. The two-tailed Student’s t-test and Mann–Whitney rank sum test were used.

Results
ATX1 is required for root growth and cell production in the RAM
A subset of histones present in the atx1-1 loss-of-function mutant analysed in this study is known to be modified. Specifically, K4 methylation of histone H3 is significantly lower than in the Wt (Alvarez-Venegas and Avramova, 2005). The mutant exhibits abnormal flower development (Alvarez-Venegas et al., 2003) and retarded root growth (Fig. 1A). The length of the primary root of the atx1-1 mutant at 8 dpg was 60% of that of the Wt (Fig. 1B). Analysis of the longitudinal zonation pattern showed that the RAM was significantly shorter in the mutant (Fig. 1C). The cell proliferation domain (PD) and transition domain (TD) of the RAM (Ivanov and Dubrovsky, 2013) were clearly visible. The reduced RAM length was caused by a decrease in the length of the PD (Table 1). Confocal sections showed that the RAM cells of the atx1-1 mutant were larger than those of the Wt (Figs 1C, 2), suggesting that cell division was delayed in the atx1-1 RAM. To test the hypothesis that cell proliferation was affected in the mutant, the expression of a G2/M transition marker, Cyclin B1;1::GUS (Colón-Carmona et al., 1999), in the atx1-1 background, was analysed (Fig. 1E). Indeed, far fewer RAM cells exhibited GUS activity in atx1-1 than in the Wt, suggesting that cell proliferation activity was compromised in atx1-1.

To determine to what extent the cell proliferation activity of the RAM was affected in the atx1-1 mutant, various parameters related to root growth and RAM activity were analysed (Table 1). Between 7 and 8 dpg, the growth rate of atx1-1 roots was only 35% of that of the Wt. Fully elongated cell length was not affected in the mutant, indicating that decreased RAM activity was the main cause of the retarded root growth. Interestingly, while the length of the TD was the same in the Wt and atx1-1, the length of the atx1-1 PD and the number of cells in this region were both 49% of those in the Wt. As a result, cell production by the RAM was 2.5-fold lower in the mutant than in the Wt (Table 1). The decreased activity of the RAM was also reflected in the increased cell cycle time. The cell cycle was 1.5 times longer in atx1-1 than in the Wt.

RAM maintenance depends on a well-regulated balance between cell proliferation and the transition of cells to
elongation (Ivanov, 1974, 1997; Barlow, 1976; Perilli et al., 2012). Increased or decreased RAM size is thought to signify delayed or accelerated transition to elongation, respectively (e.g. Dello Ioio et al., 2008). To determine if the decreased RAM length of the atx1-1 mutant is related to an increased transition to elongation, the number of cells that start to elongate during the same time period in the mutant and Wt was estimated. It has been predicted that during one cell cycle, ln2 $N_{PD}$ (i.e. $\sim 70\%$ of $N_{PD}$) cells leave the PD of the RAM and become displaced to the TD and EZ of the RAM (Ivanov and Dubrovsky, 1997). Based on the estimated cell cycle duration, this approach was used to evaluate the number of cells that leave the RAM PD during a 24h period. Surprisingly, this analysis showed that the number of cells

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Fig. 1. ATX1 is required for primary root growth. (A) Wild-type (Ws) and atx1-1 seedlings, at 15 days post-germination (dpg). (B) Primary root growth dynamics of Ws and atx1-1 during the first 8 dpg. Values are means ±SD ($n=27–32$). Combined data of three independent experiments are shown. (C) Longitudinal zonation pattern in the primary roots of Ws and atx1-1 seedlings at 8 dpg. Pseudo-Schiff-stained roots were analysed using confocal laser scanning microscopy. The proliferation domain (PD) and the transition domain (TD), which together form the root apical meristem (RAM), and the elongation zone (EZ) are colour-coded. (D and E) $CycB2;1::GUS$ expression in Ws (D) and atx1-1 (E) seedlings at 8 dpg. Representative roots are shown ($n=17–20$ for each genotype). Scale bars=10mm (A), 50 μm (C), and 100 μm (D and E).
leaving the RAM PD is 2.9 times lower in the mutant than in the Wt (Table 1). Overall, this analysis suggests that ATX1 modulates root growth by regulating the cell cycle time, cell production, and the transition from cell proliferation in the RAM to elongation. As RAM activity depends on the stem cell niche, it was thus important to establish whether the stem cell niche and RAM organization were altered in atx1-1.

ATX1 is required for the organization and cell patterning of the RAM and its stem cell niche

It was found that the organization of the QC and initial (stem) cells was irregular in atx1-1 roots. Columella initial cells are recognized based on the absence of starch granules, which are present in differentiated columella cells. Columella initials in atx1-1 formed one tier of cells similar to the Wt. However, in 21 out of 22 (95%) of the atx1-1 roots examined, the QC was abnormal. Whereas QC cells are commonly transversely aligned in Wt roots (Fig. 2A; Dolan et al., 1993), such an alignment was rarely found in the mutant roots, and the QC cells were frequently irregular in shape (Fig. 2B–D). From the second week after germination onwards, cells in the QC of Wt Arabidopsis seedlings of the Ws accession undergo periclinal divisions (Baum et al., 2002). In agreement with this observation, periclinal divisions in the QC cells were observed in 12 of 16 (75%) Wt and 15 of 19 (80%) atx1-1 seedlings at 8 dpg. As a result, the QC was composed of on average 1.8 cells in the longitudinal direction, and no statistical differences were found between the Wt and atx1-1 plants (n=16–19, P=0.883, Mann–Whitney rank sum test). However, the QC height (the QC size in the longitudinal direction) was 19% greater in the mutant than in the Wt (Fig. 2K), indicating that the QC cells in atx1-1 were more expanded than those in the Wt. Cell patterning above the QC was also strongly affected in the mutant, and aberrant oblique divisions in the provascular cylinder and ground tissue were not uncommon (arrowheads in Fig. 2B–D).

The root cap–protoderm (RCP) initial (stem) cell divides asymmetrically to give rise to cells with different cell type identities: protoderm (epidermis) and the lateral root cap (Kuras, 1978; Dolan et al., 1993; Baum and Rost, 1996; Wenzel and Rost, 2001; Cruz-Ramírez et al., 2012). Once the daughter cells have yielded the protoderm and the lateral root cap, the RCP stem cell divides again. When it undergoes a periclinal division (i.e. parallel to the nearest root surface), the division is recognized as a T-division (Kuras, 1978). This sequence of formative and proliferative division events is well coordinated, and T-divisions are therefore regularly distributed in the Wt (Baum and Rost, 1996; Wenzel and Rost, 2001).

This regularity is lost in atx1-1, and T-divisions are frequently observed in close proximity to each other (Fig. 2B, C). Cells in the PD mostly undergo anticlinal divisions (i.e. perpendicular to the nearest root surface). In all root tissues, including those of the provascular cylinder, many instances of atypical oblique divisions were found (Fig. 2C, arrowheads). Together, the irregularly shaped and enlarged QC, the atypical distribution of T-divisions, and the presence of oblique divisions in atx1-1 suggest that a lack of coordination between cell division and cell growth results in abnormal cell patterning in atx1-1 roots.

ATX1 restricts the expression domain of cells with quiescent centre identity

To establish whether these abnormalities affected cell type identities in the stem cell niche, various cell type marker lines were analysed. SCARECROW (SCR), a GRAS family transcription factor, is involved in RAM maintenance and radial patterning. It is expressed in the QC, cortex–endodermis initial (stem) cells, their daughters, and the endodermis (Di Laurentizio et al., 1996; Nakajima et al., 2001; Sabatini et al., 2003). WUSCHEL-RELATED HOMEOBOX 5 (WOX5) is specifically expressed in the QC (Sarkar et al., 2007). To analyse possible changes in QC identity, atx1-1 was crossed with the respective marker lines, and F2 seedlings carrying the marker in the atx1-1 background were selected and propagated. The pSCR::GFP expression domain in atx1-1 primary roots (n=17) was similar to that of Wt roots (Fig. 2F). pWOX5::GFP expression was observed in the QC of both Wt and atx1-1 roots.

However, in 18 of 29 (62%) of the mutant plants analysed, the domain of pWOX5 activity was expanded in the primary RAM compared with the Wt, as pWOX5::GFP was observed in the QC (as for the Wt) and also in provascular cells adjacent to the QC (Fig. 2G, H; Supplementary Fig. S1A–C available at JXB online). Moreover, in the RAM of first-order LRs, the pWOX5::GFP expression domain was expanded to an even greater extent (Fig. 2I, J) and was detected in 18 out
of 22 (82%) LRs analysed. A QC46 QC marker was expressed in the QC in 13 out of 20 (65%) atx1-1 primary roots, and the expression domain was expanded and detected in adjacent provascular cylinder cells (data not shown). Furthermore, it is known that RAM activity depends on established auxin gradients (Blilou et al., 2005; Petersson et al., 2009).

To examine whether auxin response gradients in the RAM were affected in the mutant, the auxin-response marker DR5rev::GFP (Friml et al., 2003) was introduced into the atx1-1 background. Interestingly, DR5rev::GFP expression was unaltered in the atx1-1 RAM (Supplementary Fig. S1D, E). Furthermore, primary root growth in atx1-1 was inhibited by IAA to the same extent as in the Wt (Supplementary Fig. S2A, B). These results suggest that RAM abnormalities are auxin independent.

Collectively, the results indicated that ATX1 is required for the maintenance of QC identity, RAM organization, and cell patterning. In particular, ATX1 contributes to the restriction of WOX5 and QC46 expression to the stem cell niche.

ATX1 is involved in LR emergence by controlling the timing and proliferation of LRP cells

As cell proliferation-related processes and cell patterning in the atx1-1 mutant were affected during primary root development (Figs 1, 2; Table 1), it was of interest to analyse LR development in the atx1-1 mutant. First, investigations were carried out to determine whether LRP initiation was affected. Despite retarded primary root growth, the density of all LR initiation events (including LRs and LRPs) was not affected in the mutant (Fig. 3A). Next, the LR initiation index was estimated; this is a parameter that evaluates the number of LR initiation events per root portion comprising 100 cortical cells of average length in a file (Dubrovsky et al., 2009). As cell length was unaltered in atx1-1 (Table 1), it was assumed that the LR initiation index would also be unchanged. Indeed, no differences was found in the index (Fig. 3B), indicating that LR initiation was not affected in the mutant. However, it was noticed that atx1-1 had a less rooty phenotype (Fig. 1A). To characterize this phenotype quantitatively, the density of LRP3 and LRs within the branching zone, which includes the root portion from the most distal (rootward) LR to the primary root base (Dubrovsky and Forde, 2012), was estimated. The LR density was significantly (2-fold) lower in the atx1 mutant than in the Wt (Fig. 3C). As the density of all LR initiation events (including LRs and LRPs) in the mutant did not differ from those in the Wt (Fig. 3A), but the LR density in the branching zone decreased, it was expected that the primary (G, H) and lateral (I, J) roots. (G, I) Wt plants. (H, J) atx1-1. (K) The QC height in Wt (Ws) and atx1-1 plants. An asterisk indicates a statistically significant difference (P<0.05, n=16–19, Student’s t-test; error bars indicate the SD). In all cases, 8 dpg plants were analysed, and n=16–29 per genotype. (A–D) Pseudo-Schiff-stained, fixed roots were analysed using confocal laser scanning microscopy and shown after the application of Gaussian blur and Unsharpmask filters. (E–J) Red signal is from propidium iodide, which labels cell walls. Scale bars=20 μm (A–D), 50 μm (E–H) and (J) and (J).

Fig. 2. Cell patterning is altered in the primary root apical meristem (RAM) of atx1-1. (A–D) RAM organization in Ws (A) and atx1-1 (B–D) plants. The organization shown in (A) was found in 20 out of 21 Wt plants analysed. In contrast, only one out of 22 atx1-1 plants had a normal RAM organization (B), whereas the RAM was disorganized in the remaining 21 plants analysed (C and D). Square brackets indicate the quiescent centre (QC); T-divisions are indicated by red Ts; arrowheads indicate oblique divisions. (E, F) pSCR::GFP expression in the roots of Wt (E) and atx1-1 (F) plants. (G–J) pWOX5::GFP expression in primary (G, H) and lateral (I, J) roots. (G, I) Wt plants. (H, J) atx1-1. (K) The QC height in Wt (Ws) and atx1-1 plants. An asterisk indicates a statistically significant difference (P<0.05, n=16–19, Student’s t-test; error bars indicate the SD). In all cases, 8 dpg plants were analysed, and n=16–29 per genotype. (A–D) Pseudo-Schiff-stained, fixed roots were analysed using confocal laser scanning microscopy and shown after the application of Gaussian blur and Unsharp mask filters. (E–J) Red signal is from propidium iodide, which labels cell walls. Scale bars=20 μm (A–D), 50 μm (E–H) and (J) and (J).
ATX1 is required for root development

Density of LRP vs. time

Fig. 3. Quantitative analysis of lateral root (LR) formation in wild-type (Ws) and atx1-1 plants. (A) Combined density of LR initiation events (including LRs and LRP). (B) LR initiation index. The density and index were estimated within the branching and LR formation zones of the primary root. (C) LR and LRP density in the branching and LR formation zones. Mean ±SD, n=22. *P<0.001, Student’s t-test. The scheme at the bottom shows the branching zone and the LR formation zone of the seedling’s primary root.

ATX1 is required for LRP morphogenesis independently of auxin response

Even though the incidence of LRP initiation was unaffected in the primary root of the mutant (Fig. 3), a detailed analysis showed that LRP initiation and development were affected in terms of morphogenesis. The first anticlinal divisions in the pericycle usually result in the formation of a core of a few cells that form an LR primordium (five cells in Fig. 5A). In atx1-1, such a core was frequently missing and stage I LRP was much longer than in the Wt (Fig. 5B). Developing LRP within the atx1-1 LR formation zone showed detectable GUS staining (Fig. 4B). Even in primordia where GUS-positive cells were found, the number of such cells was much lower than in the Wt (Fig. 4C–H). Overall, these data suggested that ATX1 regulates the timing of LRP development, apparently through its involvement in cell proliferation. As the data indicated that ATX1 is important for cell patterning in the primary root meristem, and cell proliferation and patterning are frequently coupled, how cell patterning is affected during LRP morphogenesis was next studied.

ATX1 is required for root development

(see details in the Materials and methods). The observations in Arabidopsis show that this is indeed the case (data not shown). LR formation time (i.e. the time from LR initiation to emergence) has not been evaluated for Arabidopsis. Here, it was demonstrated that this is a relatively rapid process that takes on average 38.1 h in the Wt. In atx1-1, however, this time was 1.7-fold greater (Fig. 4A). Slower LR formation in the mutant compared with the Wt may explain the decreased root branching phenotype.

Cell production was affected in the atx1-1 primary RAM (Table 1), and LR formation in the mutant was also slower than in the Wt. Therefore, it was hypothesized that cell proliferation may also be compromised during LRP formation. To test this possibility, the expression of the Cyb1;1::GUS G2/M transition marker (Colón-Carmona et al., 1999) was analysed in the atx1-1 background. Only 23% of LRP within the atx1-1 LR formation zone showed detectable GUS staining (Fig. 4B). Even in primordia where GUS-positive cells were found, the number of such cells was much lower than in the Wt (Fig. 4C–H). Overall, these data suggested that ATX1 regulates the timing of LRP development, apparently through its involvement in cell proliferation. As the data indicated that ATX1 is important for cell patterning in the primary root meristem, and cell proliferation and patterning are frequently coupled, how cell patterning is affected during LRP morphogenesis was next studied.

ATX1 is required for root development

(see details in the Materials and methods). The observations in Arabidopsis show that this is indeed the case (data not shown). LR formation time (i.e. the time from LR initiation to emergence) has not been evaluated for Arabidopsis. Here, it was demonstrated that this is a relatively rapid process that takes on average 38.1 h in the Wt. In atx1-1, however, this time was 1.7-fold greater (Fig. 4A). Slower LR formation in the mutant compared with the Wt may explain the decreased root branching phenotype.

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detected in the Wt. This analysis showed that patterning was affected in both the early and late stages of LRP formation.

As abnormal LRP morphogenesis was common in atx1-1, experiments were conducted to examine how this was related to QC establishment during LRP formation. It was found that in Wt seedlings, at LRP stages V–VII the expression of the QC marker \( pWOX5::GFP \) was restricted to QC cells, while a greater number of cells expressed the marker in atx1-1 (Fig. 5M). Similarly, a greater domain of \( pWOX5::GFP \)-expressing cells was found in recently emerged LRs of the mutant (Fig. 5N). These observations suggest that QC establishment was affected in LRP of the atx1-1 mutant. Morphogenetic abnormalities in LRP development could be related to abnormal stem cell activities during the establishment of a new RAM. As auxin gradients are important for the RAM activity and the QC cell identity (Sabatini et al., 1999; Aida et al., 2004; Blilou et al., 2005), the expression of the auxin reporter, \( DR5rev::GFP \), was also analysed. No apparent change in GFP (green fluorescent protein) expression was found in stage V–VII LRPs or in recently emerged LRs (Fig. 5O). \( AUXIIAA14 \) is a key auxin response gene involved in LR formation (Fukaki et al., 2005). As expected, the level of \( AUXIIAA14 \) expression increased upon auxin treatment; however, the expression of \( AUXIIAA14 \) did not differ in the roots of Wt and atx1-1 plants, either with or without auxin treatment (Supplementary Fig. S2E at JXB online).
Therefore, this analysis showed that ATX1 controls cell patterning during LR development and that the developmental abnormalities of the atx1-1 mutant are apparently unrelated to auxin response gradients.

**Discussion**

Here the role of the most extensively studied *Arabidopsis* TrxG gene, *ATX1* (Avramova, 2009), was explored, and it was shown that, in addition to its known role in flower development, *ATX1* is an important player in root development. The data indicate that *ATX1* is required for primary root growth through its role in maintaining RAM activity and the transition to elongation, but not in cell elongation itself (Table 1). The RAM activity in the mutant was compromised in part by its increased cell cycle time. The fact that neither the transition domain of the RAM nor the fully elongated cells were affected in atx1-1 underlines the importance of *ATX1* for cell proliferation-related processes and cell production. Interestingly, *SDG2* is also required for root growth, and the differences between the *sdg2* mutant and the Col Wt were similar to those between atx1-1 and the Ws Wt (Guo et al., 2010; Yao et al., 2013). It has been shown that *SDG2* is a major histone methyltransferase that contributes to the genome-wide H3K4me3...
modifications in Arabidopsis (Guo et al., 2010). Therefore, this and other histone methyltransferases are expected to be functional in atxl-1. In spite of this, the cell proliferation and cell patterning defects were found in atxl-1 roots. This suggests that different methyltransferases are involved in regulating different aspects of developmental processes and implies non-redundant requirements of ATXI for root development.

It has been proposed that cell proliferation in the RAM and the transition from proliferation to elongation are regulated in the root independently (Ivanov, 1974, 1981, 1997). Decreased RAM length does not necessarily indicate accelerated cell differentiation as is sometimes considered (e.g. Dello Ioio et al., 2008). Evaluation of how many cells during the same time period are displaced from the RAM to the TD and the EZ in the Wt and atxl-1 would provide an estimate of whether the transition to elongation is affected in the mutant. This parameter was evaluated and a significant decrease in atxl-1 was found compared with the Wt (Table 1). These results suggest that ATXI modulates both cell proliferation and the transition to elongation. In studies of floral timing, it has been proposed that co-localization of the binding sites of an activating transcription factor and a Polycomb response element, which result in competition between PcG proteins and a transcription factor, may represent a general mechanism for timing regulation of cell division-dependent processes (Sun et al., 2014). The present findings that ATXI is required to maintain cell cycle timing in the RAM and is involved in regulating the transition to elongation suggest that TrxG members may be involved in controlling the timing-related processes of RAM development.

RAM activity depends on stem cell activity. Descendants of stem cells either differentiate or maintain a proliferation-competent state. This decision is mediated by the balanced activity of PcG and TrxG proteins (Köhler and Hennig, 2010). Abnormal RAM organization in the atxl-1 strongly suggests that ATXI is required for stem cell activity in the root. The facts that the QC was larger in atxl-1 and that pWOX5::GFP and QC46 expression domains were expanded compared with the Wt indicate that the QC identity was compromised in the atxl-1 mutant. This in turn may explain the abnormalities observed in initial (stem) cells, as their activity is dependent on QC cells (Van Den Berg et al., 1995). There is a distinct difference between stem cell organization in sdg2 and atxl-1 mutants: in contrast to atxl-1 (Fig. 2), columella initial (stem) cells are differentiated in sdg2-3 roots (Yao et al., 2013). However, the QC cells in sdg2-3 do not lose their identity, similar to those of atxl-1 (Yao et al., 2013), and both mutants maintain a functional RAM, although with different degrees of abnormalities. In Wt roots, the auxin concentration is maximal in the QC (Petersson et al., 2009) and the distal auxin gradients are involved in maintaining RAM activity (Bilou et al., 2005). In spite of a number of abnormalities in the atxl-1 RAM, the auxin response was unaltered (Supplementary Fig. S1 at JXB online), in contrast to sdg2-3 (Yao et al., 2013). This observation confirms that these TrxG genes perform at least some non-redundant functions.

Overall, the analysis of the role of ATXI in the RAM suggests that it participates in cell proliferation and cell patterning processes. This conclusion was confirmed in analyses of developing atxl-1 LRs. Similar to primary root development, LRP development was slow in the atxl-1 mutant. Considering that LR formation largely depends on cell proliferation and that the cell cycle time during LRP morphogenesis is very short (Dubrovsky et al., 2001), an increase in the period from LR initiation to LR emergence in atxl-1 indicates that cell proliferation in the LRP was also affected. Despite the role of ATXI in cell proliferation demonstrated here, the rate of LR initiation was unaffected in the atxl-1 mutant, signifying that ATXI has differential roles in distinct developmental processes. Nevertheless, LR initiation was affected in terms of early primordium morphogenesis, as abnormally wide primordia were formed (Fig. 5). The chromatin-remodelling factor PICKLE (PKL) is required for primary root growth, as it maintains stem cell activity and the size of the RAM. It is also required to maintain the active state of genes involved in RAM activity, such as PLT1, PLT2, WOX5, and AGL42 (Aichinger et al., 2011). Nevertheless, in the pkl mutant, although pWOX5::GFP expression is reduced, it is restricted to the QC (Aichinger et al., 2011), whereas in the atxl-1 mutant, the pWOX5::GFP expression domain is expanded during both primary root and LR development. The increased pWOX5::GFP expression domain in the developing LRP of atxl-1 plants suggested that this mutant had abnormal or delayed QC establishment that could be responsible for the abnormal primordium morphogenesis.

To conclude, it is suggested that the morphological defects found in the atxl-1 roots are related to the observed defects in cell proliferation. For example, unusually large cells at the LRP boundaries (Fig. 5) could be the result of the increased cell cycle time and continued cell growth. If interphase cell growth is not constant for all proliferating cells, it may result in LRP asymmetry. Similarly, grouped T-divisions in the RAM (Fig. 2) may also be a consequence of increased cell cycle duration and loss of coordination between cell division and growth. As mentioned above, the QC function in the RAM and its establishment during LR formation are compromised in atxl-1. LRP morphogenesis is affected by a number of factors, many of which are auxin related (reviewed in Szymanska-Pułka, 2013). Importantly, the abnormalities identified in the LRP and RAM patterning of atxl-1 were apparently unrelated to the auxin response. It remains to be determined which genes regulated by ATXI are important for root growth and development.

Supplementary data

Supplementary data are available at JXB online.

Figure S1. Expansion of the pWOX5::GFP expression domain and the auxin response in the primary root apical meristem of the atxl-1 mutant.

Figure S2. Effect of auxin on root development in Wt (Ws) and atxl-1 plants.

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