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Uncovering the physiological and cellular effects of uranium on the root system of Arabidopsis thaliana

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ABSTRACT

Uranium (U) is a naturally occurring radionuclide that is toxic for plants. The aim of this study was to gain insights into the physiological and cellular responses of roots to U stress. We analyzed the effects of uranyl nitrate on the architecture and physiology of Arabidopsis thaliana roots using different staining procedures and reporter genes. Also, we examined the homeostasis of inorganic phosphate (Pi) and Fe during U stress. We showed that, at a sub-toxic dose, U stimulated the apex mitotic activity, resulting in improved primary root growth and reduced secondary root formation. At a toxic level, U arrested primary root growth and increased the formation of secondary and higher-order lateral roots. U stress was linked with a depletion of Pi and a redistribution of Fe in root tissues, together with the production of ROS and nitric oxide. Also, U triggered the deposition of the defense polymers callose and lignin. These results showed that part of the radionuclide effects are linked with the signaling cascade of Pi sensing in the root apex. Other mechanisms involved in U toxicity are likely related to perturbations of Fe homeostasis and direct deleterious effects of U on root components.

KEYWORDS

Callose, iron, lignin, phosphate, root development, uranium.
1. INTRODUCTION

Uranium (U) is a non-essential trace metal element and radionuclide that is naturally present at 3 mg/kg on average in the Earth’s crust (Anke et al., 2009). It is mainly redistributed in the environment by anthropogenic activities including U and phosphate mining, nuclear industry, military activities, and soil fertilization (Vandenhole, 2002). Uranium may locally accumulate to concentrations that pose potential risks for ecosystems, agrosystems, and ultimately human health. Indeed, U is chemotoxic and potentially radiotoxic (natural U has a low specific activity) for all living organisms. Uranium is not essential for plants but it is taken up from the soil, incorporated into the biomass, and consequently enters the food chain. Thus, contamination of soils by U and its absorption by plants represent a significant health risk for human beings (Anke et al., 2009; Schnug and Lottermoser, 2013). After ingestion, most U is excreted within a few days and the small fraction (0.2-5%) that is absorbed into the bloodstream is deposited preferentially in bone and kidneys, where it can cause diseases (e.g. Thiebaut et al., 2007; Vidaud et al., 2012; Schnug and Lottermoser, 2013).

According to the hard and soft acid base Lewis concept, uranyl (UO$_{2}^{2+}$) is a hard acid that reacts with hard bases such as phosphate, carbonate and hydroxyl groups (Pearson, 1963). This property, combined with other parameters such as the pH or the presence of organic acids, influences uranyl speciation in soil and water, its absorption by plants, and its interference with plant physiology and development (e.g. Ebbs et al., 1998; Vanhoudt et al., 2008; Misson et al., 2009; Laurette et al., 2012a,b). Uranium interferes with plant nutrition, photosynthesis, and induces an oxidative stress (e.g. Vanhoudt et al., 2011a,b,c, 2014; Aranjuelo et al., 2014; Saenen et al., 2014, 2015; Tewari et al., 2015). Also, it has been shown that U perturbs iron and phosphate homeostasis and may lead to phosphate starvation (Misson et al., 2009; Doustaly et al., 2014; Berthet et al., 2018).

It is well known that the root system architecture is modulated in response to changes in nutrient availability and the presence of non-essential toxic metals in the rhizosphere (for a review, see Potters et al., 2009). The molecular and cellular events that trigger morphogenic responses of the root system have been described in details for several trace elements including copper (Lequeux et al., 2010), aluminum (Kopittke et al., 2015) or chromium (Eleftheriou et al., 2015), but not for U. Plants are able to integrate internal nutritional status together with external cues to activate the right modifications for the adaptation to a new environment. Messenger molecules are important for such responses (Corpas and Barroso, 2015; Mittler, 2017). For instance, reactive oxygen species (ROS) and nitric oxide (NO) are acting in synergy to induce the modulation of root growth and even trigger programmed cell death (Correa-Aragunde et al., 2004; Locato et al. 2016). Other actors of this response are hormones. Auxin plays an important role in plant development and is a major regulator of the root system architecture. At the root apex level it maintains the meristem activity and is also implicated in the initiation and development of lateral roots (De Smet et al., 2007; Petrasek and Friml, 2009; Laskowski and Ten Tusscher, 2017). Auxin is also the link between environmental signals and root development (Kazan, 2013).

Nevertheless, the effect of U on root development and architecture is poorly described. It has been shown that U can stimulate or inhibit the primary root growth as a function of its concentration and
speciation (Misson et al., 2009). Also, U induces the accumulation of hydrogen peroxide and NO at the primary root apex (Tewari et al., 2015). These data are not sufficient per se to understand the cascade of cellular events that trigger changes in root development during U stress. In order to fill this gap, we analyzed the impact of U on the physiological and cellular processes that control root development and architecture. We showed that U modifies primary and secondary root growth through the modulation of the meristem mitotic activity. Also, we could dissect for the first time the sequence of cellular mechanisms that are induced by U in roots. These events comprise the induction of Pi deficiency, changes of Fe distribution in root tissues, the production of ROS and NO as messengers, and the accumulation of the cell wall polymers callose and lignin to control spreading of the toxic metal.

2. MATERIALS AND METHODS

2.1 Plant growth conditions

Wild-type Arabidopsis thaliana (ecotype Columbia, Col-0) and the following transgenic lines were used in this study: PIN2::PIN2-GFP (Blilou et al., 2005), DR5::GFP (Lequeux et al., 2010), and CYCB1::GUS (Lequeux et al., 2010). Seeds were surface sterilized and stored in water for 2 days at 4°C in the dark. After stratification, seeds were sown into homemade 3D printed hydroponic floating plates containing conical holes filled with 0.65% (w/v) agar (Agar plant type A, Sigma-Aldrich) solubilized in distilled water. Plates were thermoprinted using the amorphous polymer acrylonitrile butadiene styrene using an Ultimaker 2 3D-printer (Ultimaker BV, The Netherlands). Plates were transferred to plastic containers filled with 200 mL of a nutrient solution composed of 0.88 mM K$_2$SO$_4$, 2 mM Ca(NO$_3$)$_2$, 1 mM MgSO$_4$, 0.25 mM KH$_2$PO$_4$, 10 μM H$_3$BO$_3$, 0.1 μM CuSO$_4$, 0.6 μM MnSO$_4$, 0.01 μM (NH$_4$)$_6$Mo$_7$O$_{24}$, 10 μM ZnSO$_4$, 10 μM NaCl, 20 μM Fe-EDTA, and 0.25 mM MES, pH 5.8 (Meyer et al., 2015). Seedlings were grown for 6 days in a growth chamber (21°C, light intensity of 80 μmol photons m$^{-2}$ s$^{-1}$, 16 h of light per day). They were then transferred for up to 9 days onto a no phosphate (NP) or a low phosphate medium (LP, 25 μM KH$_2$PO$_4$) supplemented or not with 10 to 50 μM of uranyl nitrate (UO$_2$(NO$_3$)$_2$). Prior to any subsequent analysis, seedlings were rinsed once with a solution of 10 mM Na$_2$CO$_3$ and then twice with distilled water to remove uranyl adsorbed at the root surface (Doustaly et al., 2014).

2.2 Analysis of root architecture

Roots of freshly harvested seedlings were transferred into square Petri dishes filled with 0.5% (w/v) agarose for scanning with a GS-800 scanner (BioRad). The primary root length and the number of secondary roots were measured with the ImageJ software.

2.3 Cellular and molecular staining procedures

Cell viability at the root apex was assessed using a dual staining with fluorescein diacetate (FDA) and propidium iodide (PI) (Jones et al., 2016). The staining solution was prepared by mixing FDA (Thermo Fisher Scientific) and PI (Sigma-Aldrich) commercial solutions in NP or LP medium to obtain final concentrations of 5 μg/mL FDA and 10 μg/mL PI. Seedlings were stained for 20 min in the dark and
rinsed with NP or LP medium. FDA staining was observed with a FITC filter (excitation 480 nm; emission 543 nm) and PI with a Texas Red filter (excitation 580 nm; emission 610 nm). After acquisition, the green and red channels were fused with the Zen microscopy software (Zeiss).

Iron staining was achieved using the Perls method followed by a 3,3'-diaminobenzidine (DAB) intensification (Roschzdttardtz et al., 2009). Seedlings were stained for 45 min in a 2% (v/v) HCl and 2% (w/v) K$_2$Fe(CN)$_6$ solution and rinsed in water. For DAB intensification seedlings were transferred to a methanol solution containing 0.065% (w/v) NaN$_3$ and 0.3% (v/v) H$_2$O$_2$ and incubated for 1 hour. Finally, seedlings were rinsed with 0.1 M potassium phosphate buffer, pH 7.4, and incubated in the same buffer containing 0.00005% (w/v) CoCl$_2$, 0.00002% (v/v) H$_2$O$_2$ and 0.025% (w/v) DAB. The intensification was stopped after 5 min by rinsing seedlings with water.

ROS spatial and temporal distribution was observed using 2,7-dichlorofluorescein diacetate (Thermo Fisher Scientific) at the concentration of 2 µM in NP or LP medium. Seedlings were stained for 20 min in the dark and then rinsed in NP or LP medium. Stained seedlings were observed with a FITC filter with the same exposure time of 120 ms.

Nitric oxide accumulation was followed with 4,5-diaminofluorescein diacetate (Abcam) prepared in DMSO and diluted to a final concentration of 10 µM in the staining solution (20 mM HEPES, pH 7.2). Seedlings were stained for 10 min in the dark and then rinsed in the HEPES buffer. Stained seedlings were observed with a FITC filter with the same exposure time of 400 ms.

Callose staining was performed according to the method of Schenk and Schikora (2016). Seedlings were rinsed with a solution of 150 mM KH$_2$PO$_4$, stained for 25 min in the dark in a staining solution containing 0.01% (w/v) aniline blue and 150 mM KH$_2$PO$_4$, and then rinsed with a solution of 150 mM KH$_2$PO$_4$. Stained seedlings were observed with a DAPI filter (excitation 365 nm; emission: 460 nm) with the same exposure time of 200 ms.

Lignin, or Wiesner, staining was performed according to Pradhan Mitra and Loqué (2014). A 3% (w/v) solution of phloroglucinol (Prolabo) freshly prepared in ethanol was mixed with 37% (w/v) hydrochloric acid in a ratio of 2:1. Seedlings were directly stained on a microscope slide with the Wiesner solution for 5 min and observed under differential interference contrast light.

For β-glucuronidase activity staining seedlings were fixed in cold 90% (v/v) acetone for 20 min and incubated overnight at 37 °C in a GUS staining solution containing 0.2% (v/v) Triton X-100, 50 mM Na$_2$HPO$_4$/Na$_2$HPO$_4$, pH 7.2, 2 mM potassium ferrocyanide, 2 mM 5-Bromo-4-chloro-1H-indol-3-yl β-D-glucopyranosiduronic acid. After 24 hours of incubation, seedlings were observed under differential interference contrast light.

Stained seedlings or root sections were mounted onto microscope slides in 10% (v/v) glycerol and observed under a Leica Axioscope A1 microscope with the suitable filter.

2.4 Optical section fluorescence microscopy

Structured illumination for optical sectioning fluorescence microscopy has been conducted for the observation of DR5::GFP and PIN2::PIN2-GFP seedlings using an Axiomager Z3 Apotome microscope (Zeiss). The DR5::GFP signal was observed under a GFP filter with the same time of exposure of 120 ms.
2.5 Determination of U by inductively coupled plasma mass spectrometry (ICP-MS)

Seedlings were harvested after 3 days of treatment and digested at 90°C for 4 hours in 65% (w/v) HNO₃. Mineralized samples were diluted in 0.5% (v/v) HNO₃ and analyzed using an iCAP RQ quadrupole mass instrument (Thermo Fisher Scientific GmbH, Germany). ^238^U concentration was determined using a standard curve and corrected using an internal standard solution of ^172^Y added online. Data integration was done using the Qtegra software (Thermo Fisher Scientific GmbH, Germany).

2.6 Quantification of inorganic phosphate

Seedlings treated for 3 days were harvested, frozen in liquid nitrogen, and grinded into a fine powder. Soluble inorganic phosphate (Pi) was extracted using the protocol of Kanno et al. (2016). Samples were suspended in a buffer containing 170 mM MES pH 5.8 and 100 mM dithiothreitol (10 µL of buffer per mg of powder) and incubated for 10 min at 4°C. After centrifugation, 20 µL of the supernatant was mixed with an acidic molybdate solution (1.755% (w/v) (NH₄)₆Mo₇O₄H₂O in 3 M H₂SO₄). After 10 min of incubation, the malachite green solution containing 0.35% (w/v) polyvinyl alcohol and 0.035% (w/v) malachite green was added. Samples were incubated for 2 hours and the absorption measured at 610 nm with a Tecan Spark plate reader. Pi contents were determined using a calibration curve made with a solution of KH₂PO₄.

2.7 Statistical analysis

Statistical analysis was conducted using the R software (R Core Team, 2017) and the nparcomp package (Konietschke et al., 2015). Each parameter was compared to the control LP medium with a non-parametric Dunnett test coupled with a Fisher asymptotic approximation method. Confidence level was set at 95% (p-value <0.05).
3. RESULTS

3.1 Uranium affects cell viability in the primary root apex of Arabidopsis

We developed an experimental setup to analyze the physiological and cellular effects of U on the root system architecture of A. thaliana. The toxicity of metals on plant roots is frequently analyzed using agar-solidified culture media on which seedlings are grown in a vertical orientation (e.g. Howden and Cobbett, 1992; Hugouvieux et al., 2009; Lequeux et al., 2010). We first used this procedure to determine which concentrations of uranyl nitrate induce significant effects on the growth and architecture of primary roots of A. thaliana Col-0 seedlings. It turned out that a high concentration of U (above 100 µM) was necessary to significantly inhibit root growth of 4-day-old seedlings in these conditions (data not shown). This low sensitivity was likely due to a low availability of U because of strong interactions of uranyl ions with the hydroxyl moieties of the agar molecule and with traces of inorganic phosphate (Pi) that are present in this gelling agent (Misson et al., 2009; Jain et al., 2009). Then, we designed an alternative experimental setup using hydroponics for a better control of U bioavailability. In this system, Col-0 seeds were germinated and grown for 6 days in a hydroponic culture medium containing sufficient amounts of all nutrients, including 250 µM Pi (see Methods for details). At this stage the primary root was about 0.5 cm long and suitable to analyze the effects of U stress. Since Pi is a ligand of U, 6-day-old seedlings were transferred onto a medium with a low Pi content (25 µM Pi, LP medium) and treated with uranyl nitrate concentrations ranging from 0 to 50 µM. Seedlings were also analyzed in a medium containing no Pi at all (NP medium). We assessed U toxicity by following cell viability over time using a dual staining procedure in which living cells were stained in green with fluorescein diacetate and dead cells were stained in red with propidium iodide. No modification in cell viability was observed in NP, LP and LP plus 10 µM uranyl nitrate over the 9 days of the experiment (Fig. 1). At 20 µM U (U20 condition), a weak toxicity was observed during the first two days of the treatment, with mainly living cells and some dead cells (Fig. 1). After 9 days in U20 the cell viability was similar to control apices but the root apex was swollen, as in NP, as compared to the LP medium. A strong cellular toxicity of the radionuclide was observed at 30 µM of uranyl nitrate (U30 condition) and above (Fig. 1). In these conditions, cell viability started to decrease within the first 24 hours of U stress, with significant cell death in the outermost cell layers (epidermis and cortex) of the meristematic and elongation zones (see Supplemental Fig. S1 for the organization of the root apex in the different conditions). At days 2 and 3 in U30, cell death spread to the whole primary root apex and abnormal lateral roots emerged in the differentiation zone (Fig. 1). Secondary roots were able to elongate in U30 but higher concentrations of uranyl nitrate fully abolished their growth. Given these results, we selected the U20 and U30 conditions as sub-toxic and toxic doses of uranyl nitrate, respectively, to characterize the effects of the radionuclide on the root system of Arabidopsis.
Figure 1. Uranium affects cellular viability in the primary root apex of Arabidopsis. A. thaliana Col-0 seeds were germinated and grown for 6 days in hydroponics with a complete nutrient solution. Seedlings were then transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 0 to 50 µM of uranyl nitrate (U0-U50). Cell viability of the primary root apex was monitored after 1, 2, 3, 6 and 9 days of treatment. Living cells were stained in green with fluorescein diacetate and dead cells in red with propidium iodide. Observations were made at magnification x200 using an epifluorescence microscope. Fluorescein diacetate and propidium iodide were observed under FITC and Texas Red filters, respectively. Images are representative of n=8 samples analyzed at each time-point in two independent experiments. Scale bar = 100 µm.

3.2 Uranium perturbs phosphate and iron homeostasis in the root

It has been reported that U interferes with Pi and Fe homeostasis at the root and shoot levels (Vanhoult et al., 2011a; Doustaly et al., 2014; Berthet et al., 2018) and that Pi has a central role in U immobilization in root tissues (Misson et al., 2009; Laurette et al., 2012a). To have a better understanding of U toxicity in our experimental conditions we measured U, Pi and Fe. U was determined by ICP-MS following sample mineralization in concentrated nitric acid, whereas Pi was measured spectrophotometrically using the molybdate/malachite green reagents following extraction of soluble compounds (Kanno et al., 2016). Seedlings treated with 30 µM U for 3 days accumulated twice more of the toxic metal than seedlings treated with 20 µM U (Fig. 2A). This significant difference could explain the important changes in root apex cell viability observed between U20 and U30 conditions (Fig. 1). Measurements of soluble Pi indicated that U stress at 20 or 30 µM uranyl nitrate induced a Pi starvation in seedlings similar to the...
one measured in the NP condition. In these three conditions, the soluble Pi content was decreased by 55 to 65% as compared to the control LP medium (Fig. 2A).

Figure 2. Uranium disrupts phosphate homeostasis in Arabidopsis seedlings. Six-day-old Col-0 seedlings grown in Pi-sufficient conditions were transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30) for 3 days. A- The amount of U in seedlings was determined by ICP-MS following sample mineralization in concentrated nitric acid. B- The amount of soluble Pi in seedlings was measured spectrophotometrically using the molybdate/malachite green reagents following extraction of soluble compounds in a MES/DTT buffer (Kanno et al., 2016). The distribution of data is displayed in Tukey’s boxplots with the median as the solid line inside the box, the first and third quartiles as the bottom and top lines of the box, and whiskers with maximum 1.5 interquartile range of the lower and upper quartile, respectively. Outliers are plotted as individual dots. Each distribution represents n = 3-6 pools of 8 seedlings. Statistical significance was determined using multiple nonparametric comparisons with Dunnett’s test with p < 0.05. Letters indicate significant differences.

Since Pi starvation was not proportional to U content in the seedlings, we measured the concentration of both elements in the culture medium. Aliquots were taken at the surface of the medium, which corresponds to the zone of mineral root absorption. We observed that both Pi and U concentrations decreased rapidly and simultaneously during the first 3 days of the experiment (Supplemental Fig S2). After one day, the amount of Pi and U decreased by 60-70% in the root absorption zone, regardless of the initial concentration of uranyl nitrate. At day 3, there was less than 5 µM Pi available for seedlings and the concentration of U was about 2 µM and 6 µM in U20 and U30 conditions, respectively (Fig S2). These results most probably correspond to the formation of U-Pi complexes in the medium and their subsequent precipitation. Also, the similar extent of Pi depletion measured in the U20 and U30 media could explain that Pi starvation in the corresponding seedlings was comparable (Fig. 2B) and not proportional to the amount of absorbed U (Fig. 2A).

We used the Perls-diaminobenzidine (DAB) histological staining procedure (Roschztarttz et al., 2009) to investigate the effect of U on the distribution of labile (non-heme) Fe^{2+}/Fe^{3+} in the root apex. We observed similar patterns in the NP and U20 conditions with the accumulation of Fe in the stem cell niche and the elongation zone (Fig. 3). In the U30 condition, the distribution of Fe was markedly modified with a very strong Perls-DAB staining observed at the primary root apex and along the entire root axis.
Together these data showed that the U20 and U30 conditions lead to very different distributions of labile Fe in the root.

Figure 3. Uranium induces Fe redistribution in the root of Arabidopsis. Six-day-old Col-0 seedlings grown in Pi-sufficient conditions were transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30) for 3 days. The distribution of labile Fe$^{2+}$/Fe$^{3+}$ in roots was analyzed using the Perls-DAB staining. Observation of the dark-brown coloration typical of the Perls stain and DAB intensification (Roschzttardtz et al., 2009) was made at magnification x200 and x50 with an optical microscope under bright field. Images are representative of n=4 samples analyzed at each time-point in two independent experiments. Scale bar = 50 µm for x200 and 200 µm for x50.

3.3 Uranium modifies the root architecture

To analyze the effect of U on the root architecture we measured the primary root length and the number of secondary roots during 9 days after the transfer of 6-day-old Pi-replete seedlings to challenging media. In the NP medium, the primary root length was significantly decreased as compared to the LP condition and the difference was observed since the first day of treatment (Fig. 4AB). Also, the number of lateral roots measured at day 9 was significantly higher in NP than in the LP condition (Fig. 4C). These results were expected since the typical response to Pi starvation is a decreased primary root growth and an increased lateral root development (Péret et al., 2011). At 20 µM uranyl, the primary root length was not different from the control LP condition during the first 3 days of treatment. However, significant changes were observed at days 6 and 9 with longer primary roots in the U20 condition (Fig. 4AB). Also, the number of secondary roots was significantly reduced in U20 as compared to the LP condition (Fig. 4C). At 30 µM uranyl, the primary root growth was completely inhibited in the first 24 hours of treatment and we could not observe any reversibility of the inhibition over time (Fig. 4AB), even when U and Pi were markedly reduced in the culture medium (Fig S2). In the U30 condition, the number of secondary roots was significantly higher than in LP medium and comparable to the one measured in
NP (Fig. 4C). However, there was a marked difference between these two conditions since secondary roots elongated in NP whereas they were rapidly arrested in U30 (Fig. 4A). Together, these data showed that the changes of the root architecture induced by uranyl at sub-toxic or toxic doses are not similar to the one induced by Pi deprivation on its own.

Figure 4. Uranium modulates the root architecture of *Arabidopsis*. Six-day-old Col-0 seedlings grown in Pi-sufficient conditions were transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30). A- Graphical representation of the typical root architecture observed after 9 days of treatment. Scale bar = 0.5 cm. B- Kinetic of primary root growth. Primary root length was measured at regular time intervals. Curves represent mean ± SD with n = 20 to 60 seedlings per condition. Root length in NP and U30 was significantly different from LP from days 1 to 9; for U20, significant difference was observed at days 6 and 9. C- Production of lateral roots. The number of lateral roots per seedling was counted after 9 days of treatment. Each distribution represents n = 12 to 35 seedlings. Statistical significance was determined using multiple nonparametric comparisons with Dunnett’s test with p < 0.05. Letters indicate significant differences.

3.4 Uranium modulates the mitotic activity in the root apex

To gain insight into the mechanisms by which U modifies primary root growth and lateral root initiation we used an *Arabidopsis CYCB1::GUS* reporter line. Cyclin B1 is essential for the G2/M cell cycle transition and its expression gives clues about the relative mitotic activity (Doonan, 1996). The activity of β-glucuronidase (GUS) was detected in the primary root apex in the control LP condition and it was less important in NP (Fig. 5). This is in agreement with the reduced root growth observed in the NP condition (Fig. 4B) and the reported consequence of Pi starvation (Svistoonoff *et al.*, 2007). At 20 µM
uranyl, the mitotic activity was markedly increased during the first day of treatment and was maintained at a very high level till the end of the experiment (Fig. 5). In U30, GUS staining was also increased in the primary root apex during the first day of U stress, but this situation was transient since no more activity could be detected beyond day 2 (Fig. 5). At these stages the CYCB1::GUS reporter was detected in lateral roots that were initiated close to the primary root apex. An Arabidopsis line expressing a 35S::GUS construct was used as a control. Uranyl nitrate at 20 or 30 µM had no effect on GUS activity in this line (data not shown), indicating that the results we obtained are specific to the activity of the CYCB1 promoter. Together, these data indicated that changes in the root architecture under U stress are related to perturbations of the mitotic activity in the primary and secondary root apices.

Figure 5. Uranium modulates the mitotic activity in the root apex of Arabidopsis. A. thaliana Col-0 seeds expressing a CYCB1::GUS fusion were germinated and grown for 6 days in hydroponics with a complete nutrient solution. Seedlings where then transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30). GUS activity in root apices was observed after 1, 2 and 3 days of treatment. Observations were made at magnification x200 under an optical microscope under bright light. Images are representative of n=10 samples analyzed at each time-point in two independent experiments. Scale bar = 100 µm.

3.5 Uranium disrupts the transport and gradient of auxin in the root

Auxin is known to play a major role in the maintenance of the meristem activity and the development of lateral roots (De Smet et al., 2007; Petrasek and Friml, 2009; Laskowski and Ten Tusscher, 2017). We analyzed auxin distribution in the root apex under U stress using Arabidopsis seedlings expressing the...
auxin-reporter DR5::GFP construct. In LP, NP and U20 conditions, the pattern of DR5::GFP expression was similar and restricted to the stem cell niche and root cap (Fig. 6A). The activity of the DR5 promoter was less important in the U30 condition, with a decreased GFP signal after 1 day of treatment and no more detectable signal after 2 days of treatment (Fig. 6A).

Figure 6. A toxic dose of uranium disrupts auxin homeostasis in the primary root apex of Arabidopsis. A. thaliana lines expressing either DR5::GFP or PIN2::PIN2-GFP were germinated and grown for 6 days in hydroponics with a complete nutrient solution. Seedlings were then transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30) for one or two days before observation. A- Expression of DR5::GFP in the root apex. The auxin-responsive DR5::GFP reporter is an indicator of auxin distribution. B- Expression of PIN2::PIN2-GFP in the root apex. PIN2 is the main auxin efflux transporter at the root apex, making the hormone returning toward the shoot. Epidermis (E) and cortex (C) cells are indicated by arrow heads. Optical sections were observed with an Axioimager Z3 Apotome microscope (Zeiss) at magnification x200 under a GFP filter and/or in bright field. Images in panel B were obtained with a digital zoom of 2.6. Images are representative of n=6 to 10 samples analyzed at each time-point in two independent experiments. Scale bar is 100 µm in A, 50 µm in B.

To study the inverted fountain mechanism that controls auxin distribution in the root apex we analyzed the expression of PIN2, one of the main transporters involved in this process (Petrasek and Friml, 2009). To this aim Arabidopsis seedlings expressing PIN2::PIN2-GFP were challenged with uranyl nitrate. The PIN2 transporter is normally present in the epidermis and cortex cells and this pattern was found in the LP, NP and U20 conditions (Fig. 6B). In the U30 condition, the pattern of PIN2-GFP expression was markedly modified with diffuse and less intense fluorescence signals after one day of treatment (Fig.
Together these data indicated that toxic levels of uranyl disrupt the transport and gradient of auxin in the root apex.

3.6 Uranium induces the accumulation of ROS and the synthesis of NO in the root

Uranium stress is known to induce the production of reactive oxygen species (ROS) and, hence, to trigger antioxidant defense mechanisms in plants (e.g. Vanhoudt et al., 2008; Vanhoudt et al., 2011bc; Aranjuelo et al., 2014). Also, it has been shown that the production of hydrogen peroxide and nitric oxide (NO) is linked with the toxicity of the radionuclide in roots and leaves (Tewari et al., 2015). To investigate the pattern of ROS production in the primary root of young Arabidopsis seedlings challenged with U, we used the fluorogenic reagent 2′,7′-dichlorodihydrofluorescein diacetate that produces a green fluorescence when oxidized in the cytosol. Our results showed that, while no specific fluorescence was observed in the LP and NP conditions, ROS were detected in primary root apices exposed to U (Fig. 7A). In the U20 condition, a small and transient increase in fluorescence was observed in the elongation zone after 24 hours of treatment. In the U30 condition, a strong oxidative burst was observed after 6 hours of treatment, mainly in the elongation zone (Fig. 7A). Then, the production of ROS spread to the entire root tip after 24 hours of treatment and, finally, the fluorescence signal decreased to background level after 3 days of stress.

We analyzed NO content in the primary root of Arabidopsis during U stress using the green fluorescent indicator 4,5-diaminofluorescein diacetate (Fig. 7B). We showed that in the LP, NP and U20 conditions there was no noticeable change in fluorescence, while the U30 treatment triggered a strong production of NO. Indeed, a strong fluorescence was detected in the entire root tip after 6 hours of U stress, whereas punctuated and dense signals were observed in the elongation zone at 24 hours, before the signal decreased to the background level at 72 hours (Fig. 7B). These results suggested that NO acts as a messenger molecule in roots exposed to a toxic dose of U.
Figure 7. Uranium induces the production of ROS and NO in the primary root apex of *Arabidopsis*. Six-day-old seedlings grown in Pi-sufficient conditions were transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 μM Pi) containing 20 or 30 μM uranyl nitrate (U20, U30) for 6 to 72 hours. A- Detection of ROS using 2',7'-dichlorodihydrofluorescein diacetate. B- Detection of NO using 4,5-diaminofluorescein diacetate. Observation of both fluorescent reporters was done at magnification x200 with an epifluorescence microscope using a FITC filter. Exposure times were set at 120 ms (A) and 400 ms (B). Images are representative of n=4 to 20 samples analyzed at each time-point in two to four independent experiments. Scale bar = 100 µm.

3.7 Uranium induces the accumulation of defense polymers in the root

Callose is a polymer that controls the aperture of plasmodesmata and, thus, regulates symplasmic exchanges between plant cells (for a review, see Nedukha, 2015). The presence of callose at the root apex during U stress was investigated using aniline blue staining. Callose production was detected specifically in response to U, with no fluorescent signal in the LP and NP conditions (Fig. 8A). In the U20 condition, callose deposition was more important in the elongation zone after one day of exposure to U than after three days (Fig. 8A). In the U30 condition, the signal at day 1 was stronger than in U20 and enclosed the whole primary root apex with strong callose deposition in the meristematic zone (Fig. 8A). The fluorescent signal then disappeared after 3 days of treatment, making the callose accumulation/degradation spatiotemporal pattern similar to the one observed for ROS accumulation (Fig. 7A).
Figure 8. Uranium induces the synthesis and accumulation of defense polymers in the primary root of *Arabidopsis*. Six-day-old seedlings grown in Pi-sufficient conditions were transferred to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30) for 1 or 3 days. A- Staining of callose using aniline blue. Aniline blue fluorescence was observed using an epifluorescence microscope under a DAPI filter. Time of exposure was set at 200 ms. B- Staining of lignin using the Wiesner solution. Lignin staining was observed with an optical microscope under bright field. Observations were made at magnification x200. Images are representative of n=5 to 10 samples analyzed at each time-point in two to four independent experiments. Scale bar = 100 µm.

Lignin is a complex polymer of the cell wall that mainly accumulates in root tissues at the xylem vessels and the endodermis. Lignin contains many functional groups (e.g. hydroxyl, carboxyl) that can bind essential and toxic divalent metal ions and control their transfer between the apoplasm and the symplasm (reviewed in Liu *et al.*, 2018). We analyzed the effect of U on lignin deposition in roots using the Wiesner staining procedure. While no lignin staining was observed in LP, NP and U20 conditions, the accumulation of the polymer was detected in the stele of the elongation zone after 24 hours of U30 treatment (Fig. 8B). At day 3 in U30, lignin staining was even stronger and covered all cell files of the elongation zone except the epidermis and the cortex (Fig. 8B). Together these data suggested that lignin accumulation in the root tip has a role to address the stress induced by toxic doses of U.
4. DISCUSSION

In this study, we deciphered the physiological and cellular consequences of U stress on the root system of Arabidopsis seedlings. Since both root development and U bioavailability are influenced by many environmental cues, including the physicochemical properties and mineral composition of the rhizosphere, we set up experimental conditions to specifically analyze the effects of U. To this aim our hydroponic system comprised a complete mineral nutrient solution in which Pi was reduced from 250 µM (Pi-sufficiency) to 25 µM (Pi-limitation) during U stress. This Pi-limiting condition is in the same order of magnitude as Pi bioavailability in soil solution (10 µM) (Abel, 2017) and limits the formation of uranyl phosphate complexes that decrease U bioavailability and toxicity (Vanhoudt et al., 2008; Misson et al., 2009; Laurette et al., 2012b; Doustaly et al., 2014; Berthet et al., 2018). In these conditions, viability staining of root apices identified two concentrations of uranyl nitrate inducing a contrasted response to the radionuclide (Fig. 1).

At an initial concentration of 20 µM in the medium U induced significant changes in the root architecture, with an improved primary root growth and a reduction of secondary root formation (Fig. 4). Enhanced primary root growth was correlated with a very strong stimulation of the mitotic activity of the primary root apex, as early as the first day of U stress (Fig. 5). Such a positive effect of a stressor at low dose is called hormesis (for a review, see Poschenrieder et al., 2013). Hormesis has already been described in Arabidopsis plants challenged with U (Misson et al., 2009; Vanhoudt et al., 2011a). It could result from an overcompensation of the early perturbation in homeostasis, e.g. ROS production and induction of stress-induced antioxidants, leading to cellular responses that are beneficial for growth (Poschenrieder et al., 2013). Contamination of the medium with 30 µM uranyl nitrate revealed U toxicity with the arrest of primary root growth since the first day of stress (Fig. 5) and cell death that spread all over the primary apex after three days (Fig. 1). The U30 condition was also characterized by an increased formation of secondary and higher-order lateral roots (Fig. 4).

The changes of root architecture observed in the presence of U were not similar to, but shared features with, the ones found in the NP condition, suggesting a relationship between U toxicity and Pi deficiency. This was confirmed through monitoring Pi content in seedlings. Depletion of the medium with Pi (NP condition) or contamination of the low Pi medium with 20 µM uranyl (U20 condition) triggered a similar Pi deficiency in seedlings (Fig. 2B). In the U30 condition, Pi deficiency was slightly but significantly more important, in agreement with the increased amount of U incorporated into seedlings as compared to U20 (Fig. 2A). Phosphate deficiency was correlated with a depletion of Pi in the medium (Fig. S2), likely due to the formation of U-Pi precipitates, and to the interaction of U with Pi homeostasis in the seedlings. Indeed, it has been shown previously by 31P NMR spectroscopy that U triggers an important depletion of the cytosolic pools of Pi and phosphorylated metabolites (Berthet et al., 2018).

Recent advances have been made in the understanding of the sensory mechanisms that monitor external Pi and regulate Pi homeostasis during root development (reviewed in Abel, 2017). The different physiological parameters we analyzed in roots challenged with U indicate that part of the radionuclide effects are associated with Pi deprivation and the complex interaction between Pi sensing and Fe homeostasis in root tips. Indeed, both U treatment at a sub-toxic dose (Fig. 3) and Pi deficiency (Müller...
et al., 2015) induced Fe accumulation in the root apex and the transition zone (located between the meristem and the elongation zone). The pattern of Fe distribution was correlated with the sites of production of ROS and callose deposition upon Pi deprivation (Müller et al., 2015). We observed a similar relationship between Fe (Fig. 3), ROS (Fig. 7A) and callose deposition (Fig. 8A) in the root apices of seedlings challenged with U. Cell-type-specific callose deposition was shown to inhibit symplastic communication through plasmodesmata and to modulate the maintenance of the root apical meristem upon Pi deprivation (Müller et al., 2015). Although similar patterns of callose deposition were observed in Pi-deprived (Müller et al., 2015) or U-contaminated (Fig. 8A) conditions, the first leads to meristem arrest and stop of primary root growth, whereas in the second the mitotic activity is highly increased (Fig. 5), resulting in a stimulation of primary root growth (Fig. 4).

In the case of U stress, the adjustment of cell-to-cell communication by callose deposition in the cell wall is likely a way to limit the diffusion of the radionuclide in the root tip, as reported for other toxic trace metals (reviewed in Krzesłowska, 2011). This strategy is probably sufficient per se to limit U spread at moderate doses (e.g. U20 condition). At a higher concentration of U, we observed that a second defense polymer, lignin, accumulated in the stele of root tip (Fig. 8B). Lignin deposition in the cell wall can contribute to an improved capacity of uranyl ion binding through hydroxyl or carboxyl moieties of the polymer. The preferential localization of U deposits in the cell wall of root cells supports this assumption (Misson et al., 2009; Laurette et al., 2012a). Thus, lignin can participate in forming a physical barrier to avoid U propagation to the whole plant, as reported for other trace elements (Lequeux et al., 2010; Liu et al., 2018).

In addition to ROS, NO is likely involved in the signaling of U stress in the root. Indeed, our results showed that ROS and NO both accumulate in the root tip as early as 6 hours of treatment in U30 (Fig. 7). This is in agreement with a previous report showing that NO is produced in the roots of Arabidopsis plants challenged with U and that NO inducers or inhibitors of NO synthesis modulate the toxicity of the radionuclide (Tewari et al., 2015). The role of NO as a signal molecule during U stress is also supported by the demonstration that (i) an excess of Fe reduces root tip growth through a NO-mediated mechanism (Zhang et al., 2018), (ii) lignin is synthesized in response to high NO levels (Böhm et al., 2010; Corti Monzón et al., 2014), and (iii) NO influences the cell cycle in a dose-dependent manner, from stimulation to inhibition (Novikova et al., 2017).

As explained above, the physiological and cellular effects of U depend, in part, on a cascade of events related to Pi deprivation signaling (Fe-dependent ROS production and callose deposition) (Fig. 9). The fact that U stress and Pi deprivation per se do not result in identical symptoms indicates that the radionuclide has direct toxic effects on cellular components and/or is able to amplify Pi deficiency (Fig. 9). Yet, the first hypothesis is not supported by experimental evidences since no molecular target of the uranyl ion has been identified in plants. However, the amplification of the Pi deprivation signaling cascade is likely to occur through the interaction of U with Fe homeostasis. Indeed, the competition between U and Fe for the formation of complexes with Pi (Berthet et al., 2018) and/or for protein ligands could release Fe$^{3+}$ and increase the production of ROS via the Fenton reaction (Fig. 9). Ferritin, which control Fe related oxidative stress in plants (Briat et al., 2010; Reyt et al., 2015), is an attractive, but not yet proven, protein candidate that could explain part of the toxicity of U in the root system. Indeed, U
has been shown to interact \textit{in vitro} with a human ferritin (Michon \textit{et al.}, 2010) and may actually replace Fe into ferritins in bacteria (Cvetkovic \textit{et al.}, 2010). Ferritin is not the only candidate for the binding of the radionuclide since uranyl has been shown to replace Fe$^{3+}$ in some proteins, including serum transferrin that is involved in Fe transport (Vidaud \textit{et al.}, 2007). The identification of plant proteins that are able to bind the uranyl ion is a future challenge to gain insight into the molecular mechanisms that mediate the direct toxic effects of U.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{model.png}
\caption{Tentative model showing the physiological and cellular responses of \textit{Arabidopsis} roots to U stress. This model illustrates the three processes that could contribute to changes in the root architecture of seedlings challenged with U. A- First, U accumulation induces Pi deprivation, which in turn triggers Fe redistribution in root tissues, ROS production, and callose deposition. These events are similar to the one described for the signaling cascade of Pi sensing in the root apex (Muller \textit{et al.}, 2015; Abel, 2017). B- Second, the competition between U and Fe for the formation of complexes with diverse ligands (Pi, organic acids, cell wall components, proteins...) could release free Fe$^{3+}$, trigger the production of ROS, and thus amplify the Pi sensing cascade. C- Third, U could directly affect root growth and architecture through toxic effects on specific targets, which are yet not known. NO is, in addition to ROS, a secondary messenger involved in U stress signaling in the root.}
\end{figure}
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CONTRIBUTIONS

NBCS, CA, JB, SR conceived and designed the study; NBCS, SR performed the experiments; NBCS, CA, JB, SR analyzed the data; NBCS, JB, SR wrote the paper. JB and SR should be considered joint senior author.
REFERENCES


Supporting information

Uncovering the physiological and cellular effects of uranium on the root system of Arabidopsis thaliana

Figure S1. Structure of the Arabidopsis root apex during U stress. The organization of the different zones of the root apex in our experimental conditions is shown. RC, root cap; M, meristem; EZ, elongation zone; DZ, differentiation zone. Observations were made using an optical microscope under bright light (magnification x200). Scale bar = 100 µm.
Figure S2. Phosphate and uranium availability in the root absorption zone of the culture media. Aliquots of the culture media were collected in the root absorption zone up to three days after the transfer of Col-0 seedlings to either No Phosphate medium (NP) or Low Phosphate medium (LP, 25 µM Pi) containing 20 or 30 µM uranyl nitrate (U20, U30). A- The amount of soluble Pi was measured spectrophotometrically using the molybdate/malachite green reagents. B- The amount of U was determined by ICP-MS. Curves represent mean ± SD with n = 6 to 16 samples per condition.