The Aluminum Distribution and Translocation in Two Citrus Species Differing in Aluminum Tolerance

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Abstract

Background: Many citrus orchards of south China suffer from soil acidification, which induced aluminum (Al) toxicity. The Al-immobilization in vivo is crucial for Al detoxification. However, the distribution and translocation of excess Al in citrus species were not well illustrated.

Results: The seedlings of ‘Xuegan’ [Citrus sinensis (L.) Osbeck] and ‘Shatianyou’ [Citrus grandis (L.) Osbeck] that differed in Al tolerance were hydroponically treated with nutrient solution (Control) or supplemented by 1.0 mM Al³⁺ (Al toxicity) for 21 days after three months of pre-culture. The Al distribution at the tissue level of citrus species following the order: lateral roots > primary roots > leaves > stems. The fragmentation of fresh lateral roots revealed the ratio of Al distribution at the cell wall, cell organelle and cytoplasmic supernatant was about 8:2:1 of two citrus species under Al stress. Besides, the Al distribution at the lateral root cell wall components suggested the pectin is the most Al-accumulating site in citrus species. Compared to C. grandis, C. sinensis had a significantly higher Al concentration on the cell wall of lateral roots whereas remarkably lower Al levels on the leaves and stems. Furthermore, the Al translocation revealed by the absorption kinetics of the cell wall demonstrated that C. sinensis had a higher Al retention and stronger Al affinity on the root cell wall than C. grandis. According to the FTIR (Fourier transform infrared spectroscopy) analysis, the Al distribution and translocation might be affected by modifying the structure and components of the citrus lateral root cell wall.

Conclusions: A higher Al-retention, mainly targeted by the pectin of the root cell wall, and a lower Al translocation efficiency from roots to shoots contributed to a higher Al tolerance of C. sinensis than C. grandis.

Background

Arable soil acidification is increasing in China from 1980s to 2000s [1]. The soil acidification accelerates aluminum (Al) solubilization from minerals significantly when the soil pH is less than 5.0 [2]. The excess Al in the soil disturbs the nutrient and water balance of the rhizosphere, reducing crop yield [3], which represented one of the most limiting factors in tropical and subtropical regions [4]. For instance, it was reported that rice grain yield decreased by 28–62% [5], and wheat grain yield decreased by 23–100% [6] under Al toxicity.

The citrus orchards in south China frequently suffer from Al toxicity induced by soil acidification [7]. For instance, our investigation on the 319 soil samples from citrus orchards in Fujian province of China revealed the tested soil samples had an average pH of 4.34 and over 90% of which had a pH lesser than 5.0 [8]. In the field condition, excess Al inhibited the root development of Citrus aurantium L. significantly [9]. Accordingly, the citrus fruit yield decreased significantly under Al toxicity [10]. In the sandy culture, the biomass of citrus seedlings was depressed by Al toxicity, inducing the oxidative stress and the photosynthetic inhibition of citrus seedlings [11]. Likewise, in the hydroponic culture, high Al
concentration induced chlorotic and mottled leaves, thick root tips and less fibrous roots of citrus rootstocks [12].

The plant Al tolerance mainly relied on the rejection of Al uptake and the restriction of Al translocation [13]. The Al accumulated primarily in the roots of citrus seedlings [14, 15]. Furtherly, the Al partitioning at the cellular level revealed that the plant root cell wall is the primary target for Al-binding for most crops [16]. The cell wall was constituted mainly by polysaccharides, such as pectin, cellulose and hemicellulose. However, it is still debatable which cell wall component contributed most to the Al-binding under Al stress. For instance, Yang et al. [17] reported that hemicellulose is the main pool for Al accumulation in Arabidopsis. Differentially, Ye et al. [18] proposed that the cell wall pectin contributed mainly to Al binding in Panax notoginseng, a native plant adapted to acid soil. To our knowledge, the Al distribution pattern and the primary Al target site of citrus species are still less discussed. Besides, the potential mechanisms regarding the Al distribution and translocation in citrus species are not fully documented.

We have evaluated citrus species that differed in Al tolerance at bio-physiological [12, 19], transcriptional [20] and proteomic levels [21, 22]. Our prior studies indicated C. sinensis is much tolerant to Al toxicity than C. grandis. However, potential Al-tolerant mechanisms regarding Al distribution and translocation of citrus species are less discussed. In the present study, seedlings of C. sinensis (much Al-tolerant) and C. grandis (less Al-tolerant) were cultured by hydroponics using the nutrient solution without (as of Control) or with 1.0 mM Al$^{3+}$ (as Al toxicity). The Al distribution and translocation were investigated by cell wall fragmentation to explore the primary Al-binding site of citrus species. Besides, the study also discussed the kinetics analysis of Al adsorption and desorption, and the FTIR test of the root cell wall of two citrus species. The study will increase our understanding of the physiological mechanisms underlying the Al adaptation of citrus species.

**Results**

**The Al distribution at the tissue level of citrus species**

As shown in Fig. 1, 1.0 mM Al treatment significantly promoted Al content in lateral roots (Fig. 1a) and primary roots (Fig. 1b) compared to the Control in seedlings of two citrus species. A significant increase of Al content could also be found in the Al-treated leaves and the stems in C. grandis seedlings (Fig. 1c and 1d). However, no significant difference in Al content was observed in the Al-treated leaves and stems of C. sinensis seedlings compared to the Control. The comparison between citrus species demonstrated that the leaves and stems of C. grandis had remarkably higher Al content than C. sinensis under Al treatment. In contrast, C. sinensis lateral roots had a notably higher Al content than that of C. grandis under Al stress. Additionally, the Al content at tissue level was found to be lateral roots > primary roots > leaves > stems under Al stress.

The Al distribution at the subcellular level of citrus species
The cell wall, cell organelle and cytoplasmic supernatant of fresh lateral roots of citrus seedlings were further isolated. The Al content of each fraction was quantified respectively to investigate the Al distribution at the subcellular level (Fig. 2). The results indicated a significantly higher Al content in the Al-treated subcellular components of lateral roots than the Control in both citrus species. Strikingly, the cell wall had the most while the cytoplasmic supernatant had the most negligible Al content under Al stress of three cellular fragments. Regardless, there was no significant difference in Al content on the subcellular components of lateral roots in *C. sinensis* compared to *C. grandis* under Al stress.

The Al content of the root cell wall calculated by dry weight of citrus roots was presented in Fig. 3. The results indicated that under Al stress, the root cell wall of *C. sinensis* had a significantly higher Al than that of *C. grandis*.

**The Al distribution at cell wall fragments of citrus species**

The cell wall components of pectin, HC-I, HC-II and cellulose were isolated for Al quantification. The results of Fig. 4a indicated the Al content significantly increased in the pectin of the Al-treated cell wall of two citrus species. However, the Al content significantly decreased in HC-I and HC-II of *C. sinensis*. Differentially, there was no significant difference in Al content in HC-I and HC-II of *C. grandis* (Fig. 4b and 4c). In addition, the Al contents in the cellulose of two Citrus species had no remarkable difference under Al stress compared to control.

**The Al adsorption and desorption analysis of root cell wall of citrus species**

The cell wall of the lateral root of two citrus species was extracted for kinetics analysis of Al adsorption and desorption within 600 minutes. As shown in Fig. 5a, much Al uptake was found in the root cell wall of *C. sinensis* compared to *C. grandis*, indicating the root cell wall of *C. sinensis* had a higher capacity on Al binding than *C. grandis*. Moreover, the relative desorption rate in Fig. 5b showed that the lateral root cell wall of *C. sinensis* had a higher Al-binding rate than that of *C. grandis*. Differentially, *C. sinensis* had a lower Al desorption rate compared to *C. grandis* within 600 minutes.

**The Ftir Spectra Of Lateral Root Cell Wall**

The alterations of cell wall composition and structure by Al toxicity were revealed by FTIR analysis (Fig. 6). The wavenumber of spectra from two citrus species and related assignments were listed in Table 1. As found, the root cell wall of the Control from two citrus species had almost the same band position, indicating the similar composition of chemical groups of the root cell wall. However, band positions shifted under Al toxicity differentially in two citrus species. For instance, the vibration located at 3400 cm\(^{-1}\) was moved to 3396 cm\(^{-1}\) in *C. grandis* by Al toxicity. The vibration was shifted from 2856 cm\(^{-1}\) to 2858 cm\(^{-1}\) in *C. sinensis* and 2858 cm\(^{-1}\) to 2860 cm\(^{-1}\) in *C. grandis* by Al toxicity. It was also strikingly to find that most band positions from 1800 cm\(^{-1}\) to 800 cm\(^{-1}\), representing information of characteristic polysaccharides, amide and ester, were shifted in both of two citrus species by Al toxicity.
Apart from the band position shift, the relative absorbance at most band positions was also downregulated by Al toxicity in two citrus species compared to the Control (Fig. 6a and 6b). The digital subtraction spectra were generated by minus the Al-treated spectra from the Control spectra of the cell wall of two citrus species, respectively. As found in Fig. 6c, the band intensity of the *C. grandis* was stronger than that of *C. grandis* overall. The OPLS-DA on the relative absorbance also reflected a more apparent separation between the Control and Al-toxic cell wall of *C. grandis* than *C. sinensis* (Fig. 6d).

### Table 1

The infrared absorption frequencies of the cell wall and tentative assignment. Seedlings of *C. sinensis* and *C. grandis* were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al\(^{3+}\) (1.0 mM Al toxicity, pH 4.3). The lateral root cell wall samples were extracted for FTIR spectra analysis.

| Wavenumber (cm\(^{-1}\)) | Tentative Assignment | Reference |
|---------------------------|----------------------|-----------|
| **C. sinensis**           |                      |           |
| 3400                      | OH stretching        | [43]      |
| 2924                      | CH asymmetric stretching | [44]    |
| 2856                      | CH symmetric stretching | [44]    |
| 1740                      | C = O stretching of ester  | [44]    |
| 1649                      | C = O stretching of amide I band | [45]    |
| 1545                      | NH bending and CN stretching of amide II band | [45]    |
| 1514                      | NH bending and CN stretching of amide II band | [46]    |
| 1427                      | CH2 symmetric deformation | [47]    |
| 1375                      | COO- symmetric stretching and aliphatic group vibration | [48]    |
| 1329                      | C-O                  | [49]      |
| 1246                      | C = O stretching or NH bending of amide III bands | [47]    |
| 1155                      | phosphoryl group     | [44]      |
| 1103                      | C = O stretching, alchol hydroxyl, ether or ester base | [45]    |
| 1059                      | C-OH stretching of alcoholic groups and carboxylic acids | [50]    |
| 899                       | β-linkage between two glucose units | [47]    |
The citrus fruit trees are superbly adapted to the acid soil with potential high Al in south China [23]. Understanding of Al partition and mobilization in vivo is pivotal to reveal the Al tolerance of citrus species. Besides, it is of great significance to disclose the Al binding site of citrus species for Al toxicity mitigation. The present study addressed both challenges. The hydroponic culture had been widely carried out to explore the ion behavior of citrus species [24, 25]. Compared to our previous study in sandy culture with 1.0 mM Al treated for 18 weeks [20], the present 21 days’ hydroponic culture of citrus species resulted in almost the same Al level in leaves, indicating a reliable treatment of the study.

It has been evidenced that Al-induced phytotoxicity has many target sites from the apoplast to symplast in higher plants [26]. Accordingly, plant species varied in Al-tolerance have evolved different strategies to cope with Al toxicity based on Al distribution and translocation. Plant species native to acid soils are often found to retain excess Al in insensitive roots, protecting leaves from metabolic disruption [27]. For instance, Kopittke et al. [13] reported that the Al-tolerant wheat accumulated more than four times of Al in roots compared to the sensitive line. Similarly, the higher Al content on the root apex was also observed in Al-tolerant common bean compared to the Al-sensitive genotype [28]. The present results supported higher Al in roots under Al stress compared to shoots in citrus species (Fig. 1). Besides, C. sinensis had a significantly higher Al content in lateral roots but significantly lower Al content in shoots than C. grandis under Al stress. Thus, the less Al translocation from roots to shoots in C. sinensis might contribute to the mitigation of Al-toxicity in the shoot. Likewise, the higher Al translocation of C. grandis than C. sinensis was also reported in our previous study by 1.0 mM Al stress under 18 weeks’ sandy culture [12]. However, the Al stress within 21 days did not significantly differ in the biomass accumulation of two citrus species (data not shown). With the stress duration increased to 15 weeks, the C. sinensis seedlings had remarkably higher biomass accumulation than C. grandis in both leaves and roots (Fig. s1). Conclusively, the relatively higher Al tolerance of C. sinensis is related to a less Al translocation from the roots to shoots.

Plant root cell wall is the first defense conferring Al toxicity. Clarkson et al. [29] revealed that over 85% of Al accumulated on the cell wall of barley roots. For woody plants, more than 88% of total Al was localized in the root cell wall of the conifer [30]. In the study, the cell wall of citrus lateral roots accumulates higher Al content than the cell organelle and cytoplasmic supernatant. The ratio of Al concentration on the cell wall, cell organelle and cytoplasmic supernatant is about 8:2:1 (Fig. 2a) under 1.0 mM Al stress, indicating the prominent roles of the root cell wall in Al immobilization of citrus species. Interestingly, the ratio of cell wall-binding Al is very close to the finding of Al-treated tea (Camellia sinensis) roots [31]. Moreover, the Al distribution at cell wall fraction is in the order of pectin > HC-II > HC-I > cellulose, suggesting the pectin holds the most while the cellulose has the least Al. The contribution of pectin in Al sequestration was also reported in rice roots [32]. Li et al. [33] proposed that a high density of carboxylic groups on the pectin contributes to Al binding. Further studies regarding pectin content and related structural deformation under Al stress of citrus species are needed to reveal the role of pectin in Al detoxification.
Ma et al. [34] reported that the cell wall of Al sensitive wheat had higher Al retention than Al tolerant cultivar under 10 μM Al within 9 hours’ duration. By contrast, we observe that dry roots of *C. sinensis* had a remarkably higher Al content on the cell wall than that of *C. grandis* (Fig. 3), which is consistent with the higher Al content of lateral roots in *C. sinensis* than *C. grandis*. Therefore, we propose that the Al distribution pattern in higher plants depends on the toxic intensity, such as Al level and stress duration. For example, the Al tolerant cultivar might exclude Al encountering weak Al stress, which resulted in less Al accumulation. However, when the Al exclusion is not enough for Al detoxification, the mechanism of Al translocation *in vivo* was activated, such as Al stabilization on the roots or cell wall. For another, the Al distribution and translocation might reflect the flexible strategies for Al detoxification between woody and gramineous plants considering the different root structures and extreme variation of root biomass.

The adsorption and desorption kinetics demonstrated that the root cell wall of *C. sinensis*, an Al-tolerant species, had a higher Al affinity than *C. grandis* (Fig. 5). By contrast, the root cell wall of *C. grandis* exhibited a lower Al adsorption and a higher Al desorption, indicating less tight Al-binding on the root cell wall, which would facilitate higher Al translocation from apoplast to symplast. Therefore, we infer that Al-tolerant woody plants were prone to retain excess Al on the root cell wall to diminish Al translocation owing to their high capacity of the root systems. Besides, the Al binding firmly on roots is economical for Al resistance considering the energy cost during Al translocation. The findings of the present results also implied that organic material prepared from cell walls is promising in alleviating the Al toxicity of the citrus plants in acidic red soils.

The Al binding resulted in modification of the root cell wall, which could be assessed by FTIR analysis [35, 36]. In the study, the results that almost no new characteristic peak emerged indicated less effect of Al toxicity on the types of functional groups on the cell wall by Al toxicity overall in two citrus species. The modification of cell wall by Al stress might mainly be dependent on the abundance of chemical groups on the root cell wall of citrus species. For instance, the spectra at 3400 cm\(^{-1}\) (-OH stretching), was shifted to 3396 cm\(^{-1}\) under Al stress in *C. grandis*, suggesting the changed hydrogen-bonding mode and the destroyed connection of cell wall components by Al toxicity (Table 1). The results approved less Al tolerance of *C. grandis* than *C. sinensis* by considering the flexible deformation of hydrogen bonds between molecules [24]. Also, a study indicated the absorbance at 1740 and 1649 cm\(^{-1}\) represents the absorption of the esterified and non-esterified carboxyl groups of pectin, respectively [37]. The present results of downregulated relative absorbance at 1740 cm\(^{-1}\) and 1649 cm\(^{-1}\) were coincident with significantly higher Al accumulation in the pectin (Fig. 6c), suggesting the role of cell wall pectin in Al-binding under Al toxicity. Besides, it is also interesting to find that the vibrations from 1200 cm\(^{-1}\) to 900 cm\(^{-1}\) (Table 1), which belong to the polysaccharide fingerprint region [38], shifted and decreased under Al toxicity. The results indicated that the altered structure and content of cell wall polysaccharides under Al toxicity would affect the Al binding on the root of citrus species. Both of the digital subtraction spectra (Fig. 6c) and the OPLS-DA (Fig. 6c) of relative absorbance in two citrus species supported a much apparent alteration of the cell wall in *C. grandis* compared to *C. sinensis*, such as severer damage under Al toxicity. Similarly, a higher relative absorbance of the upper leaves corresponding to a much obvious
symptom of boron deficient orange seedlings compared to lower leaves was also reported based on the FTIR analysis [24]. Further studies based on isotope labeling of Al and pectin deformation and polysaccharides quantification of the citrus root cell wall are needed to disclose the Al spatial and temporal distribution.

**Conclusion**

The study indicated that Al toxicity altered the structure and components of the root cell wall in citrus seedlings. The pectin of the lateral root cell wall was most abundant in the Al-accumulation of citrus species. Compared to *C. grandis*, a less tolerant citrus species, *C. sinensis* had a higher Al retention on the root cell wall and a lower Al translocation efficiency from roots to shoots by 1.0 mM Al treated for 21 days in hydroponics. The novel mechanisms in Al partition and translocation could be crucial for citrus seedlings to cope with Al toxicity.

**Methods**

**Plant culture and treatments**

The citrus species ‘Xuegan’ [*Citrus sinensis* (L.) Osbeck] and ‘Suanyou’ [*Citrus grandis* (L.) Osbeck] used in the study were formally identified by Fujian Academy of Forestry Sciences (FAFS, Fuzhou, China). In December 2018, citrus fruits were harvested from the demonstration orchard of FAFS and stored in a fridge at 4 °C. For germination, the seeds of *C. sinensis* and *C. grandis* were sown in a plastic tray filled with clean river sand at the greenhouse in early April of 2019. Four weeks after germination, seedlings of uniform size (about 10 cm) were transferred to black tanks containing nutrient solution and aerated for 30 min every two hours. The nutrient solution contained 1 mM KNO$_3$, 1 mM Ca(NO$_3$)$_2$, 0.1 mM KH$_2$PO$_4$, 0.5 mM MgSO$_4$, 10 µM H$_3$BO$_3$, 2 µM MnCl$_2$, 2 µM ZnSO$_4$, 0.5 µM CuSO$_4$, 0.065 µM (NH$_4$)Mo$_7$O$_24$ and 20 µM Fe-EDTA. The pH of the nutrient solution was adjusted to 4.30 by 1 M HCl or NaOH and was replaced every two days. Three months after transplanting, the plants were subjected to the treatments with 0 (Control) or 1.0 mM Al (Al toxicity) in the nutrient solution described above (pH 4.30). The samples of citrus leaves, stems, primary roots and lateral roots were divided and collected 21 days after treatments when visible leaf chlorosis appeared on Al-treated *C. grandis* leaves.

**Quantification Of Al At The Tissue Level**

The leaves, stems, primary roots and lateral roots of citrus species were dried and digested in HNO$_3$/HClO$_4$ (5:1, v/v), and Al content was quantified according to Hsu [39].

**Quanification of Al on the cell wall, cell organelle and cytoplasmic supernatant of fresh lateral roots**

The cell wall, cell organelle and cytoplasmic supernatant were isolated according to Gao et al. [40]. In brief, 0.3 g of fresh citrus lateral roots was homogenized with buffer containing 0.25 M sucrose, 50 mM
Tris-HCl (pH 7.5) and 1 mM dithiothreitol at 4°C. The homogenate was then filtered through an 8-layer of cheesecloth to collect the filtrate 1. Then the residue left on the cheesecloth was washed twice by the buffer described above to get the filtrate 2. Both filtrate 1 and filtrate 2 were pooled and centrifuged at 300 g for 30 s. The resulting pellet after centrifugation and the residue on the cheesecloth were pooled as the cell wall debris. Meanwhile, the harvested supernatant was further centrifuged at 20,000 g for 45 min to obtain the cytoplasmic supernatant on the top and the cell organelle at the sediments. Finally, the samples of subcellular components were digested by 6 ml of HNO$_3$: HClO$_4$ (1:5, v/v) overnight and quantified by ICP-MS.

### Quantification of Al on cell wall fractions of citrus lateral roots

The crude cell wall of citrus lateral roots was extracted and fractioned according to Zhong and Lauchli [41] with modifications. Briefly, about 50 mg citrus lateral root was powdered and polled into a centrifuge tube with 5 ml ice-cold 75% ethanol for 20 min on ice. The samples were then centrifuged at 1000 g for 10 min. The supernatant was discarded, and the resulted pellets were centrifugated at 17,000 g for 10 min three times with 5 ml 80% ethanol, methanol-chloroform mixture (1:1, v/v) and acetone, respectively. The final pellets were pooled as crude cell wall after dried and weighted.

The dry crude cell wall from the lateral roots of citrus seedlings was added into ammonium oxalate (containing 0.1% NaBH$_4$, pH = 4.0) (5 mg cell wall/1 ml solution) in a boiling water bath for one hour and centrifuged at 17,000 g for 10 min for three times to isolate the pectin, hemicellulose 1 (HC-I), hemicellulose 2 (HC-II) and cellulose. After centrifugation, the supernatants of each were pooled and collected as cell wall pectin. The residue was washed by water twice then extracted by 4% KOH (containing 0.1% NaBH$_4$) or 24% KOH (containing 0.1% NaBH$_4$) under room temperature three times for 24 h in total subsequently. The obtained supernatant was HC-I and HC-II fractions, respectively. The debris left was pooled as cellulose. The volume of cell wall components was measured and stored at 4°C. The Al content of the cell wall and each cell wall component was quantified according to the method described above.

### Al Adsorption And Desorption Kinetics

The Al adsorption and desorption kinetics were performed according to Zheng et al. [42] with modifications. Briefly, the adsorption solution of 0.5 mM Al$^{3+}$ in 0.5 mM CaCl$_2$ (pH 4.30) was pumped by a peristaltic pump at 0.2 ml/min through a 2 ml column loaded with 10 mg root cell wall for Al adsorption. The solution after cell wall adsorption was then collected by a fraction collector at 20 min intervals until the Al content was equal to the adsorption solution. The residue Al$^{3+}$ left in the system was washed by 0.5 mM CaCl$_2$ (pH 4.5) at 0.6 ml/h for 1 h before Al desorption by 2.5 mM CaCl$_2$ (pH 4.30) at 0.2 ml/h until the Al concentration in the collector below detection limit. Finally, the Al content in the fraction collector was quantified, and the kinetics were analyzed within 600 min. The Al absorption and desorption kinetics were performed three times independently.
FTIR spectra analysis

2 mg dry cell wall of Citrus lateral root was mixed with 200 mg KBr and pressed into a disk by FW-5A Pressor. The IR spectra of cell walls ranging from 4000 – 400 cm$^{-1}$ were recorded using Vertex 70 spectrometer with a resolution of 4 cm$^{-1}$ and 32 scans per sample. The obtained spectra were normalized and baseline-corrected by OPUS management software before exported to Excel. The data of FTIR spectra were processed by Origin Pro 2020b (OriginLab Corporation, USA). The OPLS-DA (orthogonal partial least-squares discrimination analysis) was performed in SIMCA 14.1 (Umetrics AB, Umea, Sweden).

Data Analysis

Data analysis was performed by two-way analysis of variance, and significant differences (P < 0.05) among treatments were statistically evaluated by Two-Way ANOVA using Duncan's test, using the SPSS 16.0 (SPSS Corp., Chicago, IL, USA). All the values are presented as means ± SE. Figures except OPLS-DA were generated by using Sigmaplot 12.0.

Abbreviations

HC: hemicellulose;

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests

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Authors’ contributions

HZ wrote the manuscript; XL analyzed the experimental results; ML prepared the figures and tables; PH carried out the experiments; NL designed the experiment; ZH revised the drafts of the manuscript; LC reviewed drafts of the manuscript. All authors read and approved the final manuscript.

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The effects of Al toxicity on Al distribution in lateral roots (A), primary roots (B), leaves (C) and stems (D) of C. sinensis and C. grandis seedlings. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al3+ (1.0 mM Al toxicity, pH 4.3) for 21 days. The values represent mean ± SE (N = 5). Significant differences (p≤0.05) between treatments are indicated by different letters.
Figure 2

The Al distribution in the lateral root cell wall (A), cell organelle (B) and cytoplasmic supernatant (C) of C. sinensis and C. grandis seedlings. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al3+ (1.0 mM Al toxicity, pH 4.3) for 21 days. Fresh lateral roots were used for Al quantification at subcellular levels. The values represent mean ± SE (N = 5). Significant differences (p ≤ 0.05) between treatments are indicated by different letters.
Figure 3

The effects of Al toxicity on cell wall Al contents of the dry root of C. sinensis and C. grandis seedlings. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al3+ (1.0 mM Al toxicity, pH 4.3) for 21 days. Dry lateral roots were used for cell wall extraction and Al quantification. The values represent mean ± SE (N = 5). Significant differences (p ≤ 0.05) between treatments are indicated by different letters.
Figure 4

The Al contents on pectin (A), HC-I (B) HC-II (C) and cellulose (D) of lateral root cell wall of C. sinensis and C. grandis seedlings. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al³⁺ (1.0 mM Al toxicity, pH 4.3) for 21 days. The cell wall samples of lateral roots were fractionated for Al quantification. The values represent mean ± SE (N = 5). Significant differences (p ≤ 0.05) between treatments are indicated by different letters.
Figure 5

The Al Adsorption (A) and desorption (B) kinetics of lateral root cell wall of C. sinensis and C. grandis. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) for 21 days. The lateral root cell wall samples were extracted for Al adsorption and desorption kinetics analysis within 600 minutes.
Figure 6

The FTIR spectra of the lateral root cell wall in the region of 4000 – 500 cm⁻¹ (A), 1800 – 800 cm⁻¹ (B), digital subtraction spectra (C) and the OPLS-DA of relative absorbance (D) of two citrus species. Seedlings of C. sinensis and C. grandis were treated with nutrient solution (Control, pH 4.3) or supplemented by 1.0 mM Al³⁺ (1.0 mM Al toxicity, pH 4.3). The lateral root cell wall samples were extracted for FTIR spectra analysis. The digital spectra represent Control cell wall minus Al toxic-cell wall. The values represent mean ± SE (N = 5).

Supplementary Files

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