Root iTRAQ protein profile analysis of two Citrus species differing in aluminum-tolerance in response to long-term aluminum-toxicity

Huan-Xin Jiang1,2, Lin-Tong Yang1,3, Yi-Ping Qi4, Yi-Bin Lu1,3, Zeng-Rong Huang1,3 and Li-Song Chen1,3,5,6*

Abstract

Background: Limited information is available on aluminum (Al)-toxicity-responsive proteins in woody plant roots. Seedlings of ‘Xuegan’ (Citrus sinensis) and ‘Sour pummelo’ (Citrus grandis) were treated for 18 weeks with nutrient solution containing 0 (control) or 1.2 mM AlCl3 · 6H2O (+Al). Thereafter, we investigated Citrus root protein profiles using isobaric tags for relative and absolute quantification (iTRAQ). The aims of this work were to determine the molecular mechanisms of plants to deal with Al-toxicity and to identify differentially expressed proteins involved in Al-tolerance.

Results: C. sinensis was more tolerant to Al-toxicity than C. grandis. We isolated 347 differentially expressed proteins from +Al Citrus roots. Among these proteins, 202 (96) proteins only presented in C. sinensis (C. grandis), and 49 proteins were shared by the two species. Of the 49 overlapping proteins, 45 proteins were regulated in the same direction upon Al exposure in the both species. These proteins were classified into following categories: sulfur metabolism, stress and defense response, carbohydrate and energy metabolism, nucleic acid metabolism, protein metabolism, cell transport, biological regulation and signal transduction, cell wall and cytoskeleton metabolism, and jasmonic acid (JA) biosynthesis. The higher Al-tolerance of C. sinensis may be related to several factors, including: (a) activation of sulfur metabolism; (b) greatly improving the total ability of antioxidation and detoxification; (c) up-regulation of carbohydrate and energy metabolism; (d) enhancing cell transport; (e) decreased (increased) abundances of proteins involved in protein synthesis (proteolysis); (f) keeping a better balance between protein phosphorylation and dephosphorylation; and (g) increasing JA biosynthesis.

Conclusions: Our results demonstrated that metabolic flexibility was more remarkable in C. sinensis than in C. grandis roots, thus improving the Al-tolerance of C. sinensis. This provided the most integrated view of the adaptive responses occurring in Al-toxicity roots.

Keywords: Aluminum-toxicity, Citrus, iTRAQ, Proteomics, Root, Sulfur metabolism

* Correspondence: lisongchen2002@hotmail.com
1Institute of Plant Nutritional Physiology and Molecular Biology, Fujian Agriculture and Forestry University, Fuzhou 350002, China
3College of Resource and Environmental Science, Fujian Agriculture and Forestry University, Fuzhou 350002, China
Full list of author information is available at the end of the article

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Background
In many acidic soils through the tropics and subtropics, aluminum (Al) toxicity is a major factor limiting crop productivity. Approximately 30% of the world’s total land area is acidic [1]. Furthermore, the acidity of the soils is increasing due to the environmental problems including some farming practices and acid rain [2]. When soil pH is less than 5.0, toxic forms of Al (mainly Al$^{3+}$) are solubilized into the soil solution and accumulate to high concentration that rapidly inhibits plant root growth by damaging the roots functionally and structurally, subsequently decreasing nutrient and water uptake, eventually resulting in poor crop growth and productivity [3, 4]. A traditional strategy is to raise soil pH by application of lime, thus alleviating Al-toxicity; however, the approach is both costly and ecologically unsound from the long-term point of view. A more efficient strategy is to breed Al-tolerant crop cultivars. Fortunately, plants display wide variation in their ability to deal with Al-toxicity. Variation in Al-tolerance makes it possible to breed tolerant cultivars. The success of breeding programs relies on an understanding of the physiological, biochemical and molecular mechanisms that plants tolerate Al-toxicity. Since biological processes are ultimately controlled by proteins, identification and characterization of Al-tolerant proteins will not only increase our understanding of the molecular mechanisms on plant Al-tolerance, but also will provide new information that researchers will use to screen and breed crop cultivars suited for acidic soils with higher active Al.

During the long-term evolutionary process, higher plants have evolved two main strategies (i.e., external detoxification mechanisms and internal detoxification mechanisms) that enable them to tolerate high level of soil active Al [2, 3]. Although several mechanisms for the external detoxification (i.e., formation of non-toxic Al chelates with Al ligands released by roots, alkalization of the rhizosphere, modified cell wall and redistribution of Al) have been suggested, the Al-induced secretion of organic acid anions such as citrate, malate and oxalate from roots is the most-studied mechanism of Al-tolerance in higher plants [2, 5]. Genes involved in Al-induced root secretion of citrate and malate have been isolated from wheat (ALMT1), sorghum (SbMATE) and barley (HvMATE). Transgenic plants over-expressing these genes displayed increased Al-activated root secretion of malate and/or citrate and enhanced Al-tolerance [2, 5, 6]. Internal detoxification is mainly reached by both complexation and sequestration of Al [6].

Gene expression networks revealed by transcriptomics offer us the opportunity to understand the molecular mechanisms of plant Al-toxicity and -tolerance. Although Al-induced changes in gene expressions have been examined in some detail [7–9], limited data are available on Al-toxicity-responsive proteins in higher plants. Alteration of plant proteins is an inevitable process to deal with environmental stresses including Al-toxicity [4, 10, 11]. Proteomics is a very powerful tool for analyzing the functions of the plant proteins. Abundance of a protein is not only regulated at transcriptional, but also at translational and post-translational levels. Therefore, proteomic analysis may provide more accurate and comprehensive data than what genomic studies can provide. Recently, there were several studies reporting Al-induced changes in plant proteins using two-dimensional gel electrophoresis (2-DE) or isobaric tags for relative and absolute quantification (iTRAQ) technique. Study with Al-tolerant rice (Oryza sativa) cultivar showed that root abundances of copper/zinc superoxide dismutase (Cu/Zn SOD), S-adenosylmethionine synthetase, cysteine (Cys) synthase, 1-aminocyclopropane-1-carboxylate oxidase (ACC oxidase), glutathione S-transferase (GST) increased in response to Al-toxicity, and that Cys synthase might play a key role in Al-tolerance [4]. In Al-tolerant soybean cultivar roots, Al induced the production of heat shock proteins (HSPs), GST, chalcone-related synthetase, GTP-binding protein, and ATP Binding Cassette (ABC) transporter ATP-binding protein [12]. Duressa et al. [13] observed that Al resulted in a distinct protein profile changes in Al-tolerant and Al-sensitive soybean genotypes. In tomato roots, dehydroascorbate reductase (DHAR), glutathione reductase (GR), catalase (CAT), mitochondrial aldehyde dehydrogenase (ALDH), catechol oxidase, quinone reductase, and lactoylglutathione lyase involved in antioxidant and detoxification were induced by Al [14]. Dai et al. [15] isolated 35 proteins associated with Al-tolerance in Al-tolerant wild barley XZ16. There were 16 proteins, which were up-regulated in XZ16 roots, but down-regulated or unaltered in both Al-tolerant barley cultivars Dayton and Al-sensitive wild barley XZ61. Oh et al. [16] identified 19 up-regulated protein spots such as S-adenosylmethionine, oxalate oxidase (OXO), malate dehydrogenase (MDH), Cys synthase, ascorbate peroxidase (APX) and 28 down-regulated protein spots such as HSP 70, O-methytransferase 4, enolase and amylo- genin in + Al wheat roots. Wang et al. [11] identified 106 Al-toxicity-responsive proteins from Al-sensitive and -tolerant rice roots. They observed that glycolysis/gluconeogenesis was the most significantly up-regulated biochemical process in Al-toxic roots. To our knowledge, data available on the effects of Al-toxicity on root proteomics of woody plants are very limited.

Citrus spp. are cultivated mainly in acidic and strong acidic soils such as red soil, yellow soil or lateritic red soil. In 2011, Li et al. [17] investigated the pH of 319 ‘Guanximiyou’ pummelo (Citrus grandis) orchard soils
from Pinghe county, Zhangzhou, China. Up to 90% of soils had a lower pH than 5.0. Low pH and high Al are the factors contributing to poor Citrus growth and shortened lifespan of trees [18]. In this study, we compared Al-induced quantitative and qualitative changes in proteomes that occurred in roots of Al-tolerant Citrus sinensis and Al-sensitive C. grandis seedlings [19] using iTRAQ technique in order to identify differentially expressed proteins involved in Al-tolerance and to understand the molecular mechanisms of plants to deal with Al-toxicity.

Results
Plant growth and Al concentration in roots
As shown in Fig. 1a-c, 1.2 mM Al-treated (+Al) C. grandis seedlings displayed decreased whole plant and shoot dry weights (DWs) and unchanged root DW, compared with controls, while Al-toxicity did not significantly affect C. sinensis whole plant, shoot and root DWs. There was no significant difference in whole plant, shoot and root DWs between the two species at each given Al treatment. Root concentration of Al was higher in +Al seedlings than controls, while did not significantly differ between the two species at each given Al level (Fig. 1d).

It is worth mentioning that 1.2 mM Al treatment decreased C. grandis root DW, but had no influence on C. sinensis root DW and that Al-toxicity-induced decreases in whole plant and shoot DWs were more severe in C. grandis than in C. sinensis seedlings, when 20 μM H₃BO₃ in nutrient solution was replaced by 2.5 μM H₃BO₃. In addition, Al concentration was not lower in C. sinensis than in C. grandis roots at each given Al level (Additional file 1).

Primary data analysis and protein identification
A total of 333,528 spectra were produced from the iTRAQ experiment using control and Al-toxic C. sinensis and C. grandis roots as materials. By analyzing these spectra, we identified 42,532 known spectra, 38,369 unique spectra, 15,191 peptides, 14,266 unique peptides and 4160 proteins, respectively (Fig. 2a). The number of proteins in the genome is 46,147 (http://phytozome.jgi.doe.gov/pz/portal.html#info?alias=Org_Csinensis), due to the reasons of samples and technical restrain, 4160 identified proteins in current proteomic research is normal in plants. Protein number decreased with increased number of peptides that matched to proteins, but over 67% of the proteins (i.e., 2789 proteins) included at least two peptides (Fig. 2b).

Both protein mass distribution and distribution of sequence coverage were summarized in Fig. 2c and d. Proteins with 20–30 kDa and 30–40 kDa were at the most abundant, followed by proteins with 40–50 kDa and 50–60 kDa. Protein number decreased with increased sequence coverage.

Figure 2e shows the distribution of peptide length in Citrus roots. Less than 8.7% of all peptides are ≥20 amino acid residues and 80.6% are between 8 and 17 residues.

Differentially expressed proteins by iTRAQ
In this study, a protein was considered differentially expressed when the protein had both a log2 fold-change of more than 1.5 and a P-value of less than 0.05. Based on the two criteria, 251 differentially
expressed proteins were detected in + Al C. sinensis roots, 120 (47.8 %) of which displayed increased and 131 (52.2 %) displayed decreased abundance. These proteins were related to sulfur (S) metabolism, stress and defense response, carbohydrate and energy metabolism, nucleic acid metabolism, protein metabolism, cell transport, biological regulation and signal transduction, cell wall and cytoskeleton metabolism, jasmonic acid (JA) biosynthesis and others (Additional file 2 and Fig. 3a). In + Al C. grandis roots, we isolated 44 (30.3 %) up- and 101 (69.7 %) down-regulated proteins, which were grouped into following functional categories: S metabolism, stress and defense response, carbohydrate and energy metabolism, nucleic acid metabolism, protein metabolism, cell transport, biological regulation and signal transduction, cell wall and cytoskeleton metabolism, and others (Additional file 2 and Fig. 3b). Figures 4 and 5 showed Al-induced alterations of proteins involved in S metabolism in C. sinensis and C. grandis roots and JA biosynthesis in C. sinensis roots, respectively.

As shown in Fig. 3c, a total of 347 differentially expressed proteins were detected from + Al C. grandis and C. sinensis roots. Among these proteins, 202 (96) proteins only presented in C. sinensis (C. grandis) and 49 proteins were shared by the two species. Of the 49 overlapping proteins, 45 proteins were regulated in the same direction upon Al exposure in the both species, the remaining four proteins were regulated in the opposite direction.

qRT-PCR analysis of genes for some differentially expressed proteins

To evaluate the correlation between mRNA and protein levels, we assayed the expression levels of genes for 10 differentially expressed proteins (i.e., gi|281426908, gi|75154467, gi|1171937, gi|34099833, gi|378724814, gi|21264375, gi|301341860, gi|380863042, gi|110007377 and gi|3913733) in C. sinensis and C. grandis roots (Fig. 6).

For C. grandis roots, the expression levels of nine genes matched well with our iTRAQ data, while only four genes matched well with the iTRAQ data for C. sinensis roots (Additional file 2). Six genes [i.e., gi|1171937, gi|34099833, gi|21264375, gi|301341860, gi|380863042 and gi|3913733] were regulated differentially at transcript and protein levels in the both species. Thus, it is

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**Fig. 2** Spectra, peptides and proteins identified from iTRAQ proteomics by searching against C. sinensis database (a), number of peptides that match to proteins using MASCOT (b), protein mass distribution (c), distribution of protein sequence coverage (d), and distribution of peptide length (e)
reasonable to assume that the discrepancy between the expression levels of genes and the abundances of the corresponding protein might be caused by post-translational modifications (PTMs).

Discussion

*C. sinensis* is more tolerant to Al-toxicity than *C. grandis*

Our results showed that Al-toxicity inhibited *C. grandis* shoot growth, but had no significant influence on *C. sinensis* growth, and that root Al concentration did not differ between the two species (Fig. 1). When 20 μM H₃BO₃ in nutrient solution was replaced by 2.5 μM H₃BO₃, 1.2 mM Al inhibited *C. grandis* root and shoot growth, but only significantly inhibited *C. sinensis* shoot growth even though Al concentration was higher in Al-toxic *C. sinensis* roots than in Al-toxic *C. grandis* ones. Furthermore, Al-toxicity-induced inhibition of shoot growth was less severe in *C. sinensis* seedlings than in *C. grandis* ones (Additional file 1). To conclude, the present work, like that of previous workers [19, 20], demonstrates that *C. sinensis* is more tolerant to Al-toxicity than *C. grandis*.

As shown in Additional file 2, we identified 251 differentially expressed proteins from +Al *C. sinensis* roots, while only 145 differentially expressed proteins...
were identified from +Al C. grandis roots. This demonstrated that the metabolic flexibility was more remarkable in C. sinensis roots than in C. grandis ones in response to Al, which may contribute to the higher Al-tolerance of C. sinensis.

S metabolism was up-regulated in Al-treated roots
S metabolism is a core pathway for the synthesis of molecules required for some environmental stresses including Al [4, 10]. Transgenic plants over-expressing S metabolism-related genes such as GST and glutathione peroxidase (GPX) displayed enhanced tolerance to Al-toxicity and/or to oxidative stress [21–23]. The up-regulation of ATP sulfurylase 1 (ATPS 1) and probable GPX 4 in C. sinensis roots and Cys synthase and GST in C. sinensis and C. grandis roots in response to Al (Additional file 2 and Fig. 4) indicated that S metabolism was enhanced in Al-treated roots. This agrees with the reports that S metabolism-related enzymes (i.e. GST, GPX, ATPS and Cys synthase) were induced by various stresses including Al [1, 4, 16, 24], and that Al-toxicity increased rice root concentrations of reduced glutathione (GSH) + oxidized glutathione (GSSG) and GSH [4] and Citrus leaf concentrations of GSH and GSSG [25]. Thus, S metabolism may play a role in response to Al-toxicity by the synthesis of molecules required for Al detoxification and confer Al-tolerance in both C. grandis and C. sinensis, particularly in the latter.

Proteins involved in stress and defense response
Increased reactive oxygen species (ROS) level is commonly presumed to be a major component of Al-toxicity. Here, the abundances of six antioxidant enzymes (i.e., gi|262192812, gi|378724814, gi|255587906, gi|218138216, gi|110007377 and gi|195548074) increased in +Al C. sinensis roots (Additional file 2). Al-induced up-regulation of antioxidant enzymes has also been observed in Al-toxic roots of tomato [14] and rice [11]. However, we only identified one up-regulated (i.e., gi|255549391) and two down-regulated (gi|225626263 and gi|89276748) antioxidant enzymes from +Al C. grandis ones (Additional file 2). Germin and germin-like proteins (GLPs), which have different enzyme functions such as OXO and SOD, play a role in various abiotic stresses [26, 27]. Study showed that Al-induced OXO expression in wheat roots could be involved in detoxification mechanism...
Thus, Al-induced increase in the levels of OXO 2 and seven GLPs in *C. sinensis* roots might play a role in Al-tolerance of *C. sinensis*. However, we only isolated an up-regulated OXO 2 in + Al *C. grandis* roots (Additional file 2). Also, we isolated four up-regulated proteins [i.e., blue copper protein (BCP), CBS domain-containing protein CBSX3, thioredoxin m (mitochondrial)-type, and plastid-lipid associated protein] involved in ROS scavenging from + Al *C. sinensis* roots, while only one up-regulated BCP (gi|21264375) and one down-regulated BCP (gi|2493318) were isolated from + Al *C. grandis* roots (Additional file 2). To sum up, the abundances of proteins involved in ROS detoxification were greatly enhanced in + Al *C. sinensis* roots, but were less affected in + Al *C. grandis* ones.

In addition to ROS formation, Al-toxicity also results in overproduction of aldehydes and methylglyoxal (MG) through the activation of various enzymatic and non-enzymatic reactions. As expected, the abundances of ALDH 7b involved in the detoxification of the aldehydes, alcohol dehydrogenase (ADH)-like 2, which catalyzes the inter-conversions between alcohols and aldehydes, and glyoxalase (Gly) I and Gly II involved in the detoxification of MG were enhanced in Al-toxic *C. sinensis* roots, while the abundance of the probable aldo-keto reductase (AKR) 1 involved in the detoxification of
lipid peroxidation- and/or glycolysis-derived reactive carbonyls [i.e., malondialdehyde (MDA), 4-hydroxynonenal and MG] was down-regulated in + Al C. grandis roots (Additional file 2). Thus, the ability of aldehydes and MG detoxification might be enhanced in + Al C. sinensis roots, thus contributing its higher Al-tolerance, but down-regulated in + Al C. grandis roots.

The up-regulation of two HSPs (i.e., gi|460411113 and gi|211906498) in + Al C. sinensis roots (Additional file 2) agrees with the report that two protein spots of dnaK-type molecular chaperone hsc70.1 (At5g02500) in Al-tolerant Arabidopsis ecotype increased in response to Al-toxicity [29], and that the abundance of HSP (LOC_Os06g50300) increased only in Al-tolerant rice roots [11].

Dehydrins, or group 2 LEA (late embryogenesis abundant) proteins, can function as antioxidants, ion sequestrans, or metal ion transporters in plant phloem sap [30]. Al-induced up-regulation of desiccation-related protein PCC13-62 in + Al C. sinensis roots (Additional file 2) agrees with the report that root expression level of gene encoding desiccated-related protein under Al-stress was higher in Al-tolerant soybean genotype than in Al-sensitive one [31]. However, Al-toxicity decreased the levels of LEA14-A in C. sinensis roots (Additional file 2).

We isolated seven up-regulated (i.e., gi|131026, gi|25587430, gi|21542143, gi|75163188, gi|147721660, gi|7388028 and gi|75115690) and one down-regulated (i.e., gi|190613877) pathogenesis-related (PR) proteins from + Al C. sinensis roots, four down-regulated (i.e., gi|190613877, gi|288557882, gi|21542144 and gi|332319679) and one up-regulated (i.e., gi|75115690) PR proteins from + Al C. grandis ones. Al-induced increase in S-norcoclaurine synthase (gi|75115690) was higher in C. sinensis than in C. grandis roots. Generally speaking, Al-toxicity increased the abundances of PR proteins in C. sinensis roots and decreased their levels in C. grandis roots (Additional file 2). This agrees with the reports that PR-10 protein was highly up-regulated at the transcriptional and translational levels in Al-tolerant soybean cultivar roots [12], that root PR Bet v I family protein (LOC_Os03g18850) were expressed at higher level in the tolerant rice cultivar than in the sensitive one [11], and that the expression levels of several PR genes were higher in Al-tolerant soybean genotype roots than in Al-sensitive ones [31]. Thus, the up-regulation of PR proteins in + Al C. sinensis roots might contribute to its Al-tolerance.

Harpin proteins can lead to multiple responses in plants, such as systemic acquired resistance, hypersensitive response, and enhancement of growth and drought tolerance. Over-expression of a harpin-encoding gene in rice increased drought tolerance through ABA signaling [32]. Activated harpin binding protein-1 (HrBP1) can act upstream of the salicylic acid (SA), JA, and ethylene signaling pathways in plant cells [33]. The up-regulation of HrBP1 in + Al C. sinensis roots might play a role in Al-tolerance through acting those signaling pathways.

To conclude, the total ability of antioxidation and detoxification was greatly enhanced in + Al C. sinensis roots, thus improving the Al-tolerance of C. sinensis.

**Proteins involved in carbohydrate and energy metabolism**

Rapid turnover rate of fine roots requires systematic rebuilding of tissues, which is associated with a high reduced carbon cost. Building new root biomass not only requires carbon skeletons to produce cellulose, lignin and other structural compounds, it also requires metabolic energy (growth respiration). Under stress conditions, the demand may increase owing to initiation of response mechanisms and secondary metabolism [34]. Thus, carbohydrate and energy metabolism might be enhanced in Al-treated roots to maintain the primary metabolic pathways and the carbohydrate balance. As expected, all the 18 differentially expressed proteins involved in carbohydrate metabolism were up-regulated in + Al C. sinensis roots except for two proteins (i.e., gi|75191271 and gi|1351359; Additional file 2). Wang et al. [11] observed that glycolysis/gluconeogenesis was the most significantly up-regulated biochemical process in rice roots in response to Al, suggesting that the enhancement of root glycolytic/gluconeogenetic pathway might promote Al-tolerance by balancing the level of available energy to prevent intracellular shortage. Based on these results, we concluded that the up-regulation of carbohydrate and energy metabolism in + Al C. sinensis roots was an adaptive response to meet the increased requirement for carbon skeletons and energy. By contrast, we identified four down- and nine up-regulated proteins involved in carbohydrate and energy metabolism from + Al C. grandis roots (Additional file 2). Obviously, Al-induced adaptive responses in carbohydrate and energy metabolism were less pronounced in C. grandis than in C. sinensis roots.

**AI-induced inhibition of nucleic acid biosynthesis**

We found that all these differentially expressed proteins related to nucleic acid metabolism in C. sinensis and C. grandis roots except for protein RAD-like 3 involved in regulation of transcription in the two species and ribonuclease, which catalyzes the degradation of RNA into smaller components in C. grandis, were down-regulated in response to Al-toxicity (Additional file 2), demonstrating that Al-toxicity might impair nucleic acid biosynthesis in C. sinensis and C. grandis roots and increased RNA degradation in C. grandis roots. This
agrees with the report that Al-toxicity decreased the concentrations of RNA and DNA, and increased the activities of deoxyribonuclease and ribonuclease in longan (Dimocarpus longan) leaves [35].

**Al-induced alteration in protein metabolism**

As shown in Additional file 2, all the 39 differentially expressed ribosomal proteins involved in mature ribosome assembly and translation processes were down-regulated in + Al *C. sinensis* (26) and *C. grandis* (13) roots. In addition, the abundances of two proteins (i.e., casein kinase II regulatory subunit and ribosome biogenesis regulatory protein homolog) involved in translation processes, eukaryotic translation initiation factor 5B-like involved in the initiation phase of eukaryotic translation, and peptidyl-prolyl cis-trans isomerase FKBP15-3 involved in protein folding in *C. sinensis* roots, and of two proteins (i.e., probable prefoldin subunit 1 and prefoldin subunit, putative) involved in protein folding decreased in response to Al-stress except for peptidyl-prolyl cis-trans isomerase in *C. sinensis* roots. Therefore, we concluded that Al-toxicity impaired protein biosynthesis in roots. Similar results have been obtained on Al-stressed rice roots [11]. The Al-induced down-regulation of many proteins involved in protein biosynthesis and processing, which are the major consumers of ATP and nutrients, indicated that the translation was reasonably regulated to save energy and nutrients in roots, especially in the Al-tolerant species. Similar results have been obtained on Al-stressed *Arabidopsis* [36] and rice [11] roots. In addition, the widespread down-regulation of root ribosomal proteins, especially in *C. sinensis* roots, might indicate a redeployment of resources to meet the greater demands for amino acids in non-ribosomal peptide synthesis of glutathione and phytochelation for Al complexation. This agrees with the above inference that S metabolism was up-regulated in Al-treated roots, especially in *C. sinensis* roots, and with the reports that Al-toxicity increased rice root concentrations of GSH + GSSG and GSH [4] and *Citrus* leaf concentrations of GSH and GSSG [25].

Protein degradation not only provides molecular substrates for plant respiration but also initiates adaptive responses to (a)biotic stresses by reallocating nutrients from non-essential areas of metabolism to vital cellular activities [37]. As expected, we isolated 13 up-regulated (i.e., two gi|14549156 and gi|14549156) proteins in proteolysis from + Al in *C. grandis* roots (Additional file 2). Wang et al. [11] found that the abundance of Cys protease (LOC_Os09g39970) and mitochondrial-processing peptidase subunit (LOC_Os03g11410) were increased in + Al rice roots, suggesting that Cys protease might play a role in resisting against toxic Al because root Cys protease was expressed at a higher level in the tolerant than in the sensitive cultivar when exposed to Al-toxicity. The more widespread up-regulation of protein in proteolysis in *C. sinensis* than in *C. grandis* roots might contribute to higher Al-tolerance of *C. sinensis*.

Inactive proteins (i.e., incorrect folding) and proteins, which are no longer required for cell, are tagged by ubiquitin for proteolysis. Unlike the above protease, the abundances of all the 11 differentially expressed proteins involved in ubiquitination in *C. sinensis* (6) and *C. grandis* (5) roots decreased in response to Al-toxicity (Additional file 2), meaning that the ubiquitination of some proteins might be impaired in Al-stressed roots.

Six proteins related to amino acid metabolism were altered in + Al *C. sinensis* and *C. grandis* roots (Additional file 2). Al-induced decrease in the abundance of aspartate transaminase in *C. sinensis* roots implied that Al-toxicity might decrease N metabolism flux, because it is the major enzyme controlling aspartate that is used to transport N from sources to sinks [38]. This disagrees with the report that Al-toxicity increased aspartate transaminase level in rice roots, especially in Al-tolerant cultivar [11]. In plants, sarcosine oxidase metabolizes both sarcosine and pipicolate with preferential utilization of the latter as an endogenous substrate [39]. Here, we first isolated up-regulated sarcosine oxidase in + Al *C. sinensis* roots. Nitrite reductase (NiR) is considered to be a controlling enzyme in plant NO\_\textsubscript{2} assimilation. The up-regulation of NiR in + Al *C. sinensis* roots agrees with the report that combined treatment of acidity and Pb\textsuperscript{2+} led to an increase in NiR activity of soybean roots [40]. Phenylalanine-ammonia-lyase (PAL) catalyzes the non-oxidative deamination of L-phenylalanine to form trans-cinnamic acid and a free ammonium ion. We observed that Al-toxicity only affected PAL level in Al-intolerant *C. grandis* roots (Additional file 2). However, Wang et al. [11] showed that Al-toxicity decreased PAL abundance in rice roots, especially in Al-tolerant cultivar. Glutamate decarboxylase (GAD) catalyzes the conversion of glutamate to \( \gamma \)-aminobutyrate (GABA). Root-specific GAD (GADI) is essential for sustaining GABA level in *Arabidopsis* [41]. Therefore, GABA level might be elevated in Al-treated *C. grandis* roots due to enhanced level of putative GAD (Additional file 2). The observed higher level of putative GAD also agrees with the report that rice root GAD was expressed primarily under phosphate deprivation [42].
Proteins involved in cell transport

As shown in Additional file 2, we obtained seven up- and three down-regulated proteins in cell transport from + Al C. sinensis roots, and three up- and three down-regulated proteins from + Al C. grandis roots. ABC transporter facilitates the movement of S-glutathionylated toxins formed in GST-catalyzed conjugation of GSH to toxins and other substrates across biological membranes [43]. The up-regulation of ABC transporter I family member 17 in + Al C. sinensis roots agrees with the increased requirement for the transport of S-glutathionylated toxins due to increased GST level. Al-induced increase in ABC transporter has been observed in soybean roots [12]. In Arabidopsis, a putative ABC transporter-like protein is required for Al-tolerance [44]. The induction of aquaporin in + Al C. sinensis roots might be involved in Al-tolerance of C. sinensis by regulating the transport of water or micronutrients across cell membranes [45]. However, Wang et al. [11] reported that the abundance of aquaporin PIP2 was down-regulated by ca. 50 % only in Al-tolerant rice cultivar, and that NIP2 in Al-tolerant and -sensitive roots was up-regulated to a similar extent by Al. The up-regulation of porin/voltage-dependent anion-selective channel protein in + Al C. sinensis agrees with the report that the abundance of root mitochondrial outer membrane porin 2 (LOC_Os05g45950) increased in Al-tolerant rice cultivar when exposed to Al-toxicity, and decreased in Al-sensitive cultivar [11]. Plasma membrane H+-ATPase has an important role in plant response to nutrient and environmental stresses, such as salt stress, Al-stress, P and K deficiencies [46]. Mitochondrial ADP/ATP carrier plays a central role in aerobic cell energetics by providing the ATP generated by oxidative phosphorylation to the cytosol [47]. Sec61, an endoplasmic reticulum (ER) membrane protein translocator, participates in the translocation of newly-synthesized proteins into the lumen of the ER. The hydrolysis of the signal peptide for newly-synthesized proteins is carried out by specific proteases called signal peptidases located on the luminal side of ER. The removal of the signal peptide is crucial for the maintenance of ER homeostasis [48]. The induction of protein transport protein Sec61 subunit β in + Al C. sinensis roots might provide an adaptive response by maintaining ER homeostasis. However, its abundance decreased in + Al C. grandis roots. By contrast, three proteins (i.e., gi|28380129, gi|289584365 and gi|225463791) in protein transport decreased in + Al C. sinensis roots. Besides its role in Fe storage, ferritin also plays a role in preventing oxidative damage by storing free Fe in a safe form [49]. Our finding that + Al C. sinensis roots had higher abundance of ferritin 1 agrees with the report that there was an increase in the expression of the ferritin Fe storage gene, AtFER1 in P-deficient Arabidopsis leaves [50]. Nucleobase-ascorbate transporters (NATs) have been identified in prokaryotes, fungi, plants and mammals. In plants, maize leaf permease1 (ZmLP1E1) has been characterized by functional complementation of a pure transport-deficient Aspergillus nidulans strain and is necessary for proper chloroplast development in maize. Interestingly, leaf permease1 protein does not contain a plastidic transit peptide and is expressed only in non-photosynthetic tissues such as roots and etiolated leaves [51]. The up-regulation of putative permease 1 in + Al C. sinensis roots might be involved in Al-tolerance through maintaining chloroplast morphology. Plant lipid transfer proteins (LTPs) are responsible for the shuttling of phospholipids and other fatty acid groups between cell membranes. Some LTPs with broad specificity are termed non-specific LTPs (nsLTPs). We found that the abundances of nsLTP2 and nsLTP-like protein decreased in + Al C. grandis roots. This agrees with the reports that nsLTP (E30131) was up-regulated in Al-tolerant cultivar rice roots, and down-regulated in Al-sensitive ones [52], that the expression levels of two genes encoding LTP-like proteins (DT045053 and DT045054) in wheat roots were higher in Al-tolerant than in Al-sensitive near isogenic line [53], and that mRNA levels of LTPs were higher in Al-tolerant than in Al-sensitive soybean genotype [31]. Similarly, putative phosphatidylglycerol/phosphatidilinositol transfer protein DDB_G0282179-like involved in lipid transport was increased in + Al C. sinensis roots. However, Wang et al. [11] showed that Al-toxicity decreased the abundance of LTP in Al-tolerance rice roots, but did not significantly affect its level in Al-sensitive ones. Taken all together, the ability of cell transport was enhanced in + Al roots, especially in C. sinensis, which might contribute to its higher Al-tolerance.

Proteins involved in biological regulation and signal transduction

Perception and transmission of stress signals by the cell play crucial roles in plant response to abiotic stresses. Protein kinases and phosphatases are the key players in cell signals. Here, we isolated five up-regulated proteins in phosphorylation (i.e., gi|1174718, gi|388325711, gi|25572716 and gi|255570037) and dephosphorylation (i.e., gi|75248508) in + Al C. sinensis roots (Additional file 2), demonstrating that Al-toxicity triggered phosphorylation-dependent signal transduction pathway in C. sinensis roots, which might be involved in its Al-tolerance. By contrast, two down-regulated (i.e., gi|6016428 and gi|52077942) and two up-regulated (i.e., gi|33772201 and gi|255558866) proteins in phosphorylation and one up-regulated protein in dephosphorylation (i.e., gi|75248508) were detected in + Al C. grandis roots (Additional file 2). This suggested that the balance between phosphorylation and dephosphorylation might be upset and phosphorylation of some proteins might be impaired in + Al C. grandis roots. Obviously, C. sinensis roots...
had a higher capacity to keep a better balance between phosphorylation and dephosphorylation than C. grandis ones under Al-toxicity, which might contribute to C. sinensis Al-tolerance.

Purple acid phosphatases (PAPs) play important roles in P acquisition and recycling in plants [54]. Wang et al. [11] observed that PAP1 expressed at a higher level in Al-tolerant than in Al-sensitive rice root, suggesting that Al-tolerant cultivar might relieve the Al-toxicity through improving the P acquisition. The higher up-regulation of PAP8 in + Al C. sinensis than in C. grandis roots agrees with our report that the former might more effectively acquire P than the latter with or without Al-stress [19].

Plasma membrane-associated cation-binding protein 1 (PCaP1) is involved in intracellular signals through interaction with phosphatidylinositol phosphates and calmodulin [55]. Our finding that + Al C. sinensis roots displayed higher abundance of PCaP1 (Additional file 2) agrees with the report that Cu-stress indeed the expression of PCaP1 in Arabidopsis [56].

Our finding that the level of putative G3BP-like protein-like, a protein involved in the Ras signal transduction pathway, was decreased only in + Al C. grandis roots (Additional file 2). This agrees with the report that the abundance of G3BP was up-regulated in wild Al-tolerant barley (XZ16) and unchanged in Al-tolerant barley cultivar (Dayton), and down-regulated in Al-sensitive wild barley (XZ61) when exposed to Al-toxicity [15]. These results indicated that the down-regulation of G3BP might be associated with the lower Al-tolerance of C. grandis.

Proteins involved in cell wall and cytoskeleton metabolism

Plant cytoskeleton is highly dynamic networks of proteinaceous components consisting mainly of microtubules and microfilaments. Al-induced growth inhibition and swelling of roots demonstrated that plant cytoskeleton could be a target of Al-toxicity [57]. Studies showed that root microtubules and microfilaments were altered by Al-toxicity [57, 58]. In wheat, Al led to disorganization of actin filaments and formation of actin deposits [59]. As expected, we identified five down-regulated (i.e., gi|14423860, gi|135444, gi|75250086, gi|241865168 and gi|195976596) and two down-regulated (i.e., gi|297600120 and gi|59799374) proteins related to cytoskeleton metabolism in C. sinensis and C. grandis roots, respectively. Apart from cytoskeleton proteins, we isolated two down-regulated proteins (i.e., gi|449454512 and gi|224129194) in cell wall metabolism in + Al C. sinensis roots (Additional file 2). To conclude, Al-induced alterations in root cell wall and cytoskeleton metabolism differed between the both species.

JA biosynthesis

Phytohormones play a key role in plant response to (abiotic stresses. Among these, one of the most important signal molecules is JA [60]. Here, we obtained four up-regulated proteins [i.e., two lipoxygenases (LOXs), long-chain acyl-CoA oxidase (ACX) and 12-oxophytodienoate reductase 2 (OPR2)] involved in JA biosynthesis (Additional file 2; Fig. 5). LOX is one of the key enzymes responsible for the biosynthesis of JA. Al-induced up-regulation of LOX has also been reported in soybean [61] and Cassia tora [62, 63] roots. In Arabidopsis and rice, OPR3 and OPR 7 are responsible for JA production, respectively [64, 65]. ACX catalyze the first step in the peroxisomal β-oxidation stage of JA biosynthesis. Wound-induced JA accumulation was reduced in a mutant that was defective in ACX1 and was abolished in a mutant that was impaired in both ACX1 and its closely related paralog, ACX5 [66]. Thus, both JA biosynthesis and level might be increased in + Al C. sinensis roots, thus enhancing plant Al-tolerance. Because LOX may increase the formation of oxidation products, Al-induced up-regulation of LOX has been suggested to be a response to Al-toxicity [61].

Conclusions

Our results demonstrate that C. sinensis is more tolerant to Al-toxicity than C. grandis. Using iTRAQ technique, we isolated 347 differentially expressed proteins from + Al C. sinensis and C. grandis roots. Among these proteins, 202 proteins only presented in C. sinensis, 96 proteins only presented in C. grandis, and 49 proteins were shared by the two species. In the 49 overlapping proteins, 45 (4) proteins were regulated in the same (opposite) direction upon Al exposure. This indicated more remarkable metabolic flexibility in C. sinensis than in C. grandis roots, thus improving the Al-tolerance of C. sinensis. As shown in Fig. 7, the higher Al-tolerance of C. sinensis might be related to following several factors, including: (a) activation of S metabolism; (b) greatly improving the total ability of antioxidation and detoxification; (c) up-regulation of carbohydrate and energy metabolism; (d) enhancing cell transport; (e) decreasing (increasing) proteins related to protein synthesis (proteolysis); (f) maintaining a better balance between protein phosphorylation and dephosphorylation; and (g) increasing JA biosynthesis. To sum up, we identified more differentially expressed proteins than those of previous studies in other plant roots and provided the most integrated view of the adaptive responses occurring in + Al roots. Therefore, our analysis of root Al-toxicity-responsive proteins will increase our understanding of Al-tolerant mechanisms in higher plants.
Methods

Plant culture and Al treatments

This study was conducted from February to December, 2012 at Fujian Agriculture and Forestry University, Fuzhou, China. Both plant culture and Al treatments were performed according to Jiang et al. [67] and Yang et al. [19]. Briefly, five-week-old uniform seedlings of ‘Sour pummelo’ [Citrus grandis (L.) Osbeck] and ‘Xuegan’ [Citrus sinensis (L.) Osbeck] with a single stem were selected and transplanted to 6 L pots containing fine river sand. Seedlings, two to a pot, were grown in a greenhouse under natural photoperiod. Each pot was supplied with 500 mL of nutrient solution every 2 days. The nutrient solution contained the following macronutrients (in mM): KNO$_3$, 1; Ca(NO$_3$_2)$_2$, 1; KH$_2$PO$_4$, 0.1; and MgSO$_4$, 0.5; and micronutrients (in μM): H$_2$BO$_3$, 20; MnCl$_2$, 2; ZnSO$_4$, 2; CuSO$_4$, 0.5; (NH$_4$)$_2$MoO$_4$, 0.065; and Fe-EDTA, 20. Six weeks after transplanting, each pot was supplied daily with a nutrient solution containing 0 (control) or 1.2 mM AlCl$_3$·6H$_2$O (+Al) until the sand was saturated. The pH of the nutrient solutions was adjusted to 4.1 - 4.2 using HCl or NaOH. Eighteen weeks after the beginning of Al treatments, approx. 5-mm-long root apices from new white fibrous roots were excised and immediately frozen in liquid N$_2$. Samples were stored at −80 °C until extraction. The remaining seedlings that were not sampled were used to measure whole plant, root and shoot DWs and root Al concentration.

Plant DWs and Al concentration in roots

At the end of the experiment, 10 plants per treatment from different pots were harvested and divided into shoots (leaves + stems) and roots. Their DWs were measured after being dried at 70 °C for 48 h. Root Al concentration was determined colorimetrically by the aluminon after being digested in a
mixture of HNO₃ : HClO₄ [68]. There were six replicates per treatment.

Protein extraction
Proteins were extracted from frozen roots using a phenol extraction procedure according to Yang et al. [69]. Briefly, equal amounts of frozen root tips from six plants (one plant per pot) were mixed as a biological replicate. There was one biological replicates for each treatment. About 1 g frozen mixed samples were well ground in liquid N₂ with a mortar and pestle. Four milliliter of ice-cooled buffer containing 100 mM Tris–HCl pH 7.8, 100 mM KCl, 50 mM L-ascorbic acid, 1 % (v/v) Triton X-100, 1 % (v/v) β-mercaptoethanol and 1 mM phenylmethanesulfonyl fluoride (PMSF) was added to the powder and gently pulverized. The mixture was allowed to thaw slowly on ice. The resulting suspension was transferred to a 10 mL tube, then an equal volume of Tris-phenol (pH 8.0) was added. The mixture was thoroughly vortexed before centrifuging at 13,000 g for 15 min at 4 °C. The upper phenolic phase was transferred to a 50 mL tube, and then five volumes of 100 mM ammonium acetate/methanol were added. After being mixed carefully, the mixture was stored at −20 °C overnight. The supernatant was removed carefully after centrifugation at 13,000 g for 15 min at 4 °C, then the protein pellets were resuspended in 25 mL of ice-cooled methanol for 2 h at −20 °C. Protein pellets were collected by centrifugation at 13,000 g for 15 min at 4 °C, and then were resuspended in 25 mL of ice-cooled acetone containing 0.1 % β-mercaptoethanol and kept at −20 °C for 2 h. After centrifugation at 13,000 g for 15 min at 4 °C, the pellets were washed twice with 25 mL of ice-cooled acetone, and then dried by lyophilization and finally stored at −80 °C until use. Lyophilized protein samples were grinded to fine powder with pestle and liquid N₂, then transferred to clean tubes and dissolved in lysis buffer [7 M urea, 2 M thiourea, 4 % 3-((3-cholamidopropyl)dimethylammonio)-1-propanesulfonate (CHAPS), 40 mM Tris–HCl, pH 8.5, 1 mM PMSF and 2 mM EDTA]. After 5 min, 10 mM dithiothreitol (DTT) was added to the samples. The suspension was sonicated at 200 W for 15 min and then centrifuged at 4 °C, 30,000 g for 15 min. The supernatant was transferred to another tube. To reduce disulfide bonds in proteins of the supernatant, 10 mM DTT was added and incubated at 56 °C for 1 h. Subsequently, 55 mM iodoacetamide (IAM) was added to block the cysteines, incubated for 1 h in the dark room. The supernatant was mixed well with 55 volumes of chilled acetone for 2 h at −20 °C to precipitate proteins. After centrifugation at 4 °C, 30,000 g, the supernatant was discarded, and the pellet was air-dried for 5 min, dissolved in 500 µL 0.5 M tetraethylammonium bromide (TEAB), and sonicated at 200 W for 15 min. Finally, samples were centrifuged at 4 °C, 30,000 g for 15 min. The supernatant was transferred to a new tube and quantified using a Bio-Rad Protein Assay kit based on the Bradford method using bovine serum albumin as a standard. The proteins in the supernatant were kept at −80 °C for further analysis.

iTRAQ analysis and bioinformatic analysis of proteins
iTRAQ analysis was implemented at Beijing Genomics Institute (BGI, Shenzhen, China).

One-hundred µg protein was taken out of Al-toxic and control sample solution and then the protein was digested with Trypsin Gold (Promega, Madison, WI, USA) with the ratio of protein : trypsin = 30 : 1 at 37 °C for 16 h. After trypsin digestion, peptides were dried by vacuum centrifugation. Peptides were reconstituted in 0.5 M TEAB and processed according to the manufacture’s protocol for 8-plex iTRAQ reagent (AB Sciex Inc., MA, USA). Briefly, one unit of iTRAQ reagent was thawed and reconstituted in 24 µL isopropyl-nanol. Al-toxic and control samples for C. sinensis (C. grandis) were labeled with 118 and 119 (115 and 117) tags, respectively. The peptides labeled with the isobaric tags were incubated at room temperature for 2 h. The labeled peptide mixtures were then pooled and dried by vacuum centrifugation.

For strong cationic exchange (SCX) chromatography using a Shimadzu LC-20AB HPLC Pump system (Shimadzu Co. Kyoto, Japan), the iTRAQ-labeled peptide mixtures were redissolved in 4 mL of buffer A [25 mM NaH₂PO₄ in 25 % acetonitrile (CAN), pH 2.7] and loaded onto a 4.6 × 250 mm Ultremex SCX column. The peptides were eluted at a flow rate of 1 mL min⁻¹ with a linear gradient of 5 % buffer B (25 mM NaH₂PO₄ and 1 M KCl in 25 % ACN, pH 2.7) for 7 min, 5–60 % buffer B for 20 min and 60–100 % buffer B for 2 min and maintained in 100 % buffer B for 1 min before equilibrating with buffer B for 10 min prior to the next injection. Elution was monitored by measuring the absorbance at 214 nm. The eluted peptides were pooled as 20 fractions, desalted by C-18 coloum and vacuum dried.
Each fraction was resuspended in a certain volume of buffer C (5% ACN, 0.1% formic acid) and centrifuged at 20,000g for 10 min. In each fraction, the final concentration of peptide was ca. 0.5 μg mL⁻¹. A total of 5 μL of supernatant was loaded onto a Shimadzu LC-20 AD nanoHPLC (Shimadzu Co. Kyoto, Japan) by the autosampler onto a 2 cm C18 trap column (inner diameter, 200 μm), and the peptides were eluted onto a resolving 10 cm analytical C18 column (inner diameter, 75 μm). The samples were loaded at 8 μL min⁻¹ for 4 min; then eluted at 300 nL min⁻¹ with a linear gradient of 5% buffer D (95% ACN, 0.1% formic acid) for 5 min, 3–35% buffer D for 35 min, 35–60% buffer D for 5 min, 60–80% buffer D for 2 min and maintained in 80% buffer D for 2 min, finally returned to 5% buffer D within 1 min and maintained in 5% buffer D for 10 min.

For the TripleTOF analysis, a TripleTOF 5600 system 5600 (AB SCIEX, Concord, ON, Canada) fitted with a Nanospray III source and a pulled quartz tip as the emitter was used. Data were acquired using an ion spray voltage of 2.5 kV, N gas of 30 psi, nebulizer gas of 15 psi, and an interface heater temperature of 150 °C. The MS was operated with an RP of ≥30,000 FWHM for TOF-MS scans. For information dependent acquisition (IDA), survey scans were acquired in 250 ms and as many as 30 product ion scans were collected if they exceeded a threshold of 120 cps with a 2⁺ to 5⁺ charge state. The total cycle time was fixed to 3.3 s and the Q2 transmission window was 100 Da for 100%. Four time bins were summed for each scan at a pulser frequency value of 11 kHz through monitoring of the 40 GHz multi-channel TDC detector with four-anode/channel detection. A sweeping collision energy setting of 35 ± 5 eV coupled with iTRAQ adjust rolling collision energy was applied to all precursor ions for collision-induced dissociation. Dynamic exclusion was set for 1/2 of peak width (15 s), and then the precursor was refreshed off the exclusion list.

Raw data files acquired from the TripleTOF were converted into MGF files using Proteome Discoverer 1.2 and the MGF files were searched. Proteins identification was performed by using Mascot search engine (Version 2.3.02; Matrix Science, London, UK) against C. sinensis database (http://www.phytozome.net/cgi-bin/gbrowse/citrus/). For protein identification, a mass tolerance of ±0.05 Da was permitted for intact peptide masses and ±0.1 Da for fragmented ions, with allowance for one missed cleavages in the trypsin digests. Pyrophosphorylation of glutamine and variable oxidation of methionine and iTRAQ labeling of tyrosine were set as variable modification; carbamidomethylation of Cys and iTRAQ labeling of lysine the N-terminal amino group of peptides were set as fixed modification. The peptide charge was set as Mr, and monoisotopic mass was chosen. To decrease the probability of false peptide identification, only peptides with significance scores (≥20) at the 99% confidence interval by a Mascot probability analysis greater than “identity” were counted as identified. Each confident protein identification included at least one unique peptide. An automatic decoy database search strategy was used to estimate the false discovery rate (FDR), which was calculated as the false positive matches divided by the total matches. In the final search results, the FDR was less than 1.5%. The iTRAQ 8-plex was chosen for quantification during the search.

The search results were filtered before data exportation. The filters were used for protein identification with these options: significance threshold $P<0.05$ (with 95% confidence) and ion score or expected cut-off less than 0.05 (with 95% confidence). For protein quantitation, the filters were set as follows: (a) “median” was chosen for the protein ratio type; (b) the minimum precursor charge was set to 2 and minimum peptides were set to 2, and only unique peptides were used for quantitation; and (c) normalization by median intensities, and outliers were removed automatically. The peptide threshold was set as above for identity. A 1.5 log2-fold change was set to identify significant differentially expressed proteins in addition with a $P$-value of less than 0.05. Distribution fitting of protein ratios data was performed using SPSS software. The two protein datasets were fitted into Normal distribution models and the threshold for significance of $+/−$ 1.5 was sufficient to separate differentially expressed proteins.

Bioinformatic analysis of proteins was performed according to Yang et al. [69].

Quantitative RT-PCR (qRT-PCR) analysis of gene expression

Root tips of six plants from different pots were mixed as a biological replicate. Equal amounts of root tips were collected from each plant. There were three biological replicates for each treatment. Total RNAs were independently extracted three times from the frozen roots of Al-toxic and control plants using Recalciertant Plant Total RNA Extraction Kit (Centrifugal column type, Biotek Corporation, China) according to manufacturer’s instructions. Gene-specific primers were designed using Primer Software Version 5.0 (PREMIER Biosoft International, CA, USA) according to the corresponding sequences of selected proteins in Citrus genome (http://www.phytozome.net/cgi-bin/gbrowse/citrus/). The sequences of the F and R primers used are given in Additional file 3. qRT-PCR analysis was performed according to Zhou et al. [70]. For the normalization of gene expression, β-tubulin (JN580571) gene was used as an internal standard and the roots from control plants were used as reference sample, which was set to 1.
Experimental design and statistical analysis
There were 40 seedlings (20 pots) in a completely randomized design. Experiments were performed with 3–10 replicates except for iTRAQ analysis (i.e., one biological replicate). The replicates represented material from individual plant except for iTRAQ and qRT-PCR analysis (i.e., each biological replicate was created by pooling equal roots from six different plants (one plant per pot)). Differences among four treatment combinations were analyzed by 2 (species) × 2 (AI levels) ANOVA. The unpaired t-test was applied for comparison between two means (i.e., qRT-PCR data). Power analysis was performed by using pwr.t.test functions of pwr.package in R environment (Version 3.2.0). Under empirical μ and σ, the minimum sample size of 8 and 3 could generate a power value of more than 0.8 in plant DW measurement and the other experiments, respectively.

Availability of supporting data
The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD002301.

Additional files

Additional file 1: Effects of Al-toxicity on plant growth and root AI concentration in Citrus sinensis and C. grandis seedlings. (DOC 184 kb)

Additional file 2: List of differentially expressed proteins in Al-toxicity Citrus sinensis (CS) and C. grandis (CG) roots. (DOC 418 kb)

Additional file 3: Specific primer pairs used for qRT-PCR expression analysis. (DOC 40 kb)

Abbreviations
12,13-EDOT: 12,13-epoxyoctadecatrienoic acid; 13-HPOJ: 13S-hydroperoxyoctadecatrienoic acid; 2-DE: Two-dimensional gel electrophoresis; ABC: ATP Binding Cassette; ACC: 1-aminoacyclopropane-1-carboxylate; ACX: Acai-CoA-oxidase; ADH: Alcohol dehydrogenase; ALR: Aldo-keto reductase; AI: Aluminum; ALDH: Aldehyde dehydrogenase; AOC: Alene oxide cyclase; ADG: Alene oxide synthase; APX: Ascorbate peroxidase; ATPS: ATP sulfurylase; BCP: Blue copper protein; CAT: Catalase; cis(-)-OPDA: cis(-)-12-oxophytodienoic acid; Cu/Zn SOD: Copper/Zinc superoxide dismutase; DHAR: dehydroascorbate reductase; DW: Dry weight; GLP: Germin-like protein; Gm: Glyoxalase; GPX: Glutathione peroxidase; GR: Glutathione reductase; GSH: Reduced glutathione; GSSG: Oxidized glutathione; GST: Glutathione S-transferase; HSP: Heat shock protein; iTRAQ: Isobaric tags for relative and absolute quantification; JA: Jasmonic acid; LEA: Late embryogenesis abundant protein; LOP: Lipoygenase; LTP: Lipid transfer protein; MDA: Malondialdehyde; MDH: Malate dehydrogenase; MG: Methylglyoxal; MLB: Major latent protein; NAT: Nucleobase-acorbate transporter; NIR: Nitrite reductase; nLTP: Non-specific LTP; OPR-8-3-oxo-2-(3Z)-pentenyl)cyclopentane-1-octanoic acid; ORP: Oxophytodienoic acid reductase; OXO: Oxalate oxidase; PAL: Phenylalanine ammonia-lyase; PAP: Purple acid phosphatase; POD: Peroxidase; PTM: Post-translational modification; ROS: reactive oxygen species; S: sulfur; SA: Salicylic acid.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
H.X.J. and L.T.Y. contributed equally as first author. H.X.J carried out bioinformatic analysis of proteins and drafted the manuscript. L.T.Y. extracted the proteins, carried out the qRT-PCR analysis, and participated in the design of the study. Y.P.Q. participated in the design of the study. Y.B.L. carried out the cultivation of seedlings. Z.R.H. carried out the analysis of AI L.S.C designed and directed the study and revised the manuscript. All authors have read and approved the final manuscript.

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Author details
1Institute of Plant Nutritional Physiology and Molecular Biology, Fujian Agriculture and Forestry University, Fuzhou 350002, China. 2College of Life Science, Fujian Agriculture and Forestry University, Fuzhou 350002, China. 3College of Resource and Environmental Science, Fujian Agriculture and Forestry University, Fuzhou 350002, China. 4Institute of Materia Medica, Fujian Academy of Medical Sciences, Fuzhou 350001, China. 5The Higher Educational Key Laboratory of Fujian Province for Soil Ecosystem Health and Regulation, Fujian Agriculture and Forestry University, Fuzhou 350002, China. 6Fujian Key Laboratory for Plant Molecular and Cell Biology, Fujian Agriculture and Forestry University, Fuzhou 350002, China.

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References
1. von Uexküll HR, Mutert E. Global extent, development and economic impact of acid soils. In: Date RA, Grundon NJ, Raynet GE, editors. Plant-Soil Interactions at Low pH: Principles and Management. Dordrecht: Kluwer Academic Publishers; 1995. p. 5–19.
2. Yang LT, Qi YP, Jiang HK, Chen LS. Roles of organic acid anion secretion in aluminium tolerance of higher plants. BiolMed Res Int. 2013;2013:173682.
3. Kochian LV, Hoekenga OA, Piñeros MA. How do crop plants tolerate acid soils? Mechanisms of aluminium tolerance and phosphorus efficiency. Ann Rev Plant Biol. 2004;55:459–93.
4. Yang Q, Wang Y, Zhang J, Shi W, Qian C, Peng X. Identification of aluminium-responsive proteins in rice roots by a proteomic approach: cysteine synthase as a key player in Al resistance. Proteomics. 2007;7:737–49.
5. Ma JF. Syndrome of aluminium toxicity and diversity of aluminium resistance in higher plants. Int Rev Cytol. 2007;264:225–52.
6. Liu J, Piñeros MA, Kochian LV. The role of aluminum sensing and signaling in plant aluminum resistance. J Integr Plant Biol. 2014;56:221–30.
7. Chandran D, Sharopova N, Iwashuta S, Gant JS, Vandenbosch KA, Samac DA. Transcriptome profiling identified novel genes associated with aluminum toxicity, resistance and tolerance in Medicago truncatula. Planta. 2008;228:151–66.
8. Fan W, Lou HQ, Geng YL, Liu MY, Wang ZQ, Yang LG, et al. Identification of early Al-responsive genes in rice bean (Vigna umbellata) roots provides new clues to molecular mechanisms of Al toxicity and tolerance. Plant Cell Environ. 2014;37:1586–97.
9. Zhou XX, Yang LT, Qi YP, Guo P, Chen LS. Mechanisms on boron-induced alleviation of aluminium-toxicity in Citrus grandis seedlings at a transcriptional level revealed by cDNA-AFLP analysis. PLoS ONE. 2015;10:e0115485.
10. Hostian Z, Komatsu S. Contribution of proteomic studies towards understanding plant heavy metal stress response. Front Plant Sci. 2013;3:310.
11. Wang ZQ, Xu Y, Geng QQ, Xie C, Fan W, Yang JL, et al. Root proteome of rice studied by iTRAQ provides integrated insight into aluminium stress tolerance mechanisms in plants. J Proteomics. 2014;98:189–205.
12. Chen Y, Qi JL, Wang SS, Su J, Xu GH, Zhang MS, et al. Comparative proteome analysis of differentially expressed proteins induced by Al toxicity in soybean. Physiol Plant. 2007;131:542–54.
13. Duressa D, Soliman K, Taylor R, Senwo Z. Proteomic analysis of soybean roots under aluminium stress. Int J Plant Genomics. 2011;2011:282531.
14. Zhou S, Sauvé R, Thannhauser TW. Proteome changes induced by aluminium stress in tomato roots. J Exp Bot. 2009;60:1849–57.
Science. 1992;257:85.

glutathione peroxidase in chloroplasts or role for isoenzymes of aspartate aminotransferase in plant nitrogen.

Arabidopsis

Nicotiana tabacum

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L.) in a

Arabidopsis

Citrus

Tamarix hispida

Triticum turgidum

pummelo leaf mineral nutrient status

mutants define an

ALS3

Arabidopsis thaliana

roots. Mol Biol

Trichoderma virens

in

Dimocarpus

31. Duressa D, Soliman KM, Taylor RW, Chen DQ. Gene expression profiling in

30. Hanin M, Brini F, Ebel C, Toda Y, Takeda S, Masmoudi K. Plant dehydrins and

21. Ezaki B, Gardner RC, Ezaki Y, Matsumoto H. Expression of aluminum-induced

20. Yang LT, Jiang HX, Qi YP, Chen LS. Differential expression of genes involved

19. Yang LT, Jiang HX, Tang N, Chen LS. Mechanisms of aluminum-tolerance in

15. Dai H, Cao F, Chen X, Zhang M, Ahmed IM, Chen ZH, et al. Comparative proteomic analysis of aluminum tolerance in Tibetan wild and cultivated barleys. PLoS ONE. 2013;8, e63428.

16. Oh MW, Roy SK, Kamal AH, Cho K, Cho SW, Park CS, et al. Proteome analysis of roots of wheat seedlings under aluminum stress. Mol Biol Rep. 2014;41:671–81.

17. Li Y, Han MQ, Lin F, Ten Y, Lin J, Zhu DH, Guo P, Weng YB, Chen LS. Soil chemical properties, Guaniniviruses' pumello leaf mineral nutrient status and fruit quality in the southern region of Fujian province. China. J Soil Sci Plant Nutr. 2015;15:615-28.

18. Lin Z, Myhre DL. Citrus root growth as affected by soil aluminum level under field conditions. Soil Sci Soc Amer J. 1990;54:1340–4.

19. Yang LT, Jiang HX, Tang N, Chen LS. Mechanisms of aluminum-tolerance in two species of Citrus: secretion of organic acid anions and immobilization of aluminum by phosphorus in roots. Plant Sci. 2011;180:251–38.

20. Yang LT, Jiang HX, Qi YP, Chen LS. Differential expression of genes involved in alternative glycolytic pathways, phosphorus scavenging and recycling in response to aluminum and phosphorus interactions in Citrus roots. Mol Biol Rep. 2012;39:355–66.

21. Ezaki B, Gardner RC, Ezaki Y, Matsumoto H. Expression of aluminum-induced genes in transgenic Arabidopsis plants can ameliorate aluminum stress and/or oxidative stress. Plant Physiol. 2000;127:918–27.

22. Dixit P, Mukherjee PK, Ramachandran V, Eapen S. Glutathione transferase from Trichodesma virness enhances cadmium tolerance without enhancing its accumulation in transgenic Nicotiana tabacum. PLoS ONE. 2011;6(11), e16360.

23. Yoshimura K, Miyao K, Gaber A, Takeda T, Kanaboshi H, Miyasaka H, et al. Enhancement of stress tolerance in transgenic tobacco plants overexpressing Chlamydomonas glutathione peroxidase in chloroplasts or cytosol. Plant J. 2004;37:21–33.

24. Navrot N, Collin V, Gualberto J, Gelhaye E, Hirasewa M, Rey P, et al. Plant glutathione peroxidases are functional peroxiredoxins distributed in several subcellular compartments and regulated during biotic and abiotic stress. Plant Physiol. 2006;142:1634–79.

25. Chen LS, Qi YP, Li YH. Effects of aluminum on light energy utilization and photoprotective systems in Citrus leaves. Ann Bot. 2005;96:345–51.

26. Dunwell JM, Gibbings JG, Mahmood T, Naqui SMS. Germin and germin-like proteins: evolution, structure, and function. Crit Rev Plant Sci. 2008;27:342–75.

27. Li HY, Jiang J, Wang S, Liu FF. Expression analysis of ThrGP, a new germin-like protein gene, in Tamarix hispida J For Res. 2010;21:323–30.

28. Delisle G, Champoux M, Houde M. Characterization of oxalate oxidase and cell death in Al-sensitive and tolerant wheat roots. Plant Cell Physiol. 2001;42:324–33.

29. Karuppanapandian T, Rhee SJ, Kim EM, Han BK, Hoekenga OA, Lee GP. Proteomic analysis of differentially expressed proteins in the roots of Columbia-0 and Landsberg erecta ectotypes of Arabidopsis thaliana in response to aluminum-toxicity. Can J Plant Sci. 2012;92:1267–82.

30. Hanin M, Brini F, Ebel C, Toda Y, Takeda S, Masmoudi K. Plant dehydrins and stress tolerance: versatile proteins for complex mechanisms. Plant Signal Behav. 2011;6:1503–9.

31. Duressa D, Soliman KM, Taylor RW, Chen DQ. Gene expression profiling in soybean under aluminum stress: genes differentially expressed between Al-tolerant and Al-sensitive genotypes. Am J Mol Biol. 2011;4:156–47.

32. Wei ZM, Laby RJ, Zumoff CH, Bauer DW, He SY, Collmer A, et al. Harpin, a wall-associated box protein, alters plasma membrane PtdInsP signals on plasma membrane. Plant Signal Behav. 2011;6:1503–9.

33. Wei ZM, Laby RJ, Zumoff CH, Bauer DW, He SY, Collmer A, et al. Harpin, a wall-associated box protein, alters plasma membrane PtdInsP signals on plasma membrane. Plant Signal Behav. 2011;6:1503–9.

34. Stobrawa K, Lorenc-Plucińska G. Changes in carbohydrate metabolism in

35. Oh SH, Choi WJ, Lee IT, Yun SJ. Cloning and characterization of a rice cDNA encoding glutathione deracoxabase. J Biochem Mol Biol. 2005;38:595–601.

36. Rea P. MRP subfamily ABC transporters from plants and yeast. J Exp Bot. 1999;50:895–103.

37. Larsen PB, Geisler MJB, Jones CA, Williams KM, Cancel JD. ALS3 encodes a phloem-localized ABC transporter-like protein that is required for aluminum tolerance in Arabidopsis. Plant Cell. 2005;17:533–40.

38. Maurel C, Santoni V, Lau TT, Wudack MM, Verducz J. The cellular dynamics of plant aquaporin expression and functions. Curr Opin Plant Biol. 2009;12:690–8.

39. Shen H, Chen J, Wang Z, Yang C, Sasaki T, Yamamoto Y, et al. Root plasma membrane H+-ATPase is involved in the adaptation of soybean to phosphorus starvation. J Exp Bot. 2006;57:1553–62.

40. Minka N, Weber AP. Intracellular metabolite transporters in plants. Mol Plant. 2010;3:21–53.

41. Coleman CE, Lopes MA, Gillikin JW, Boston RS, Larksins BA. A defective signal peptide in the maize high-lysine mutant floury 2. Proc Natl Acad Sci U S A. 1995;92:6828–31.

42. Briat JF, Raveret K, Arnaud N, Duc C, Bouchereau J, Touranche B. New insights into ferritin synthesis and function highlight a link between iron homeostasis and oxidative stress in plants. Ann Bot. 2010;105:811–22.

43. Misson J, Raghotthama KG, Jain A, Jouhet J, Block MA, Bilgry R, et al. A genome-wide transcriptional analysis using Arabidopsis thaliana Affymetrix gene chips defined plant responses to phosphate deprivation. Proc Natl Acad Sci U S A. 2005;102:11954–9.

44. Schultz NP, Bruntell TP, Allen A, Dellaporta SL, Nelson T, Chen J. Leaf permease1 gene of maize is required for chloroplast development. Plant Cell. 1996;8:643–70.

45. Mao CZ, Yang L, Zheng BS, Wu YR, Liu FY, Yi KY, et al. Comparative mapping of QTLs for Al tolerance in rice and identification of positional Al-induced genes. J Zhejiang Univ (Sci). 2004;5:634–43.

46. Guo P, Bai G, Carver B, Li R, Bernarda A, Baum M. Transcriptional analysis between two wheat near-isogenic lines contrasting in aluminum tolerance under aluminum stress. Mol Genet Genomics. 2007;277:1–12.

47. Kaida R, Serada S, Norioka N, Norioka S, Neumetzler L, Pauly M, et al. Potential role for purple acid phosphatase in the dephosphorylation of wall proteins in tobacco cells. Plant Physiol. 2010;153:603–10.

48. Kato M, Nagasaka-Takeuchi N, Ide Y, Tomikawa R, Maeshima M. PCaPs, possible regulators of PtdInsP signals on plasma membrane. Plant Signal Behav. 2010;5:849–50.

49. Ide Y, Nagasaki N, Tomikawa R, Suito M, Kamiya T, Maeshima M. Molecular properties of a novel, hydrophilic cation-binding protein associated with the plasma membrane. J Exp Bot. 2007;58:1173–83.

50. Biancadoro EB, Jones DL, Gilroy S. Alterations in the cytoskeleton accompany aluminum-induced growth inhibition and morphological changes in primary roots of maize. Plant Physiol. 1998;118:159–72.

51. Améndos M, Corrales I, Poschenrieder C, Illéssy S, Bollo JI, Bialek F, et al. Different effects of aluminum on the cytoskeleton of maize root tip cells of Triticum turgidum. Protoplasma. 2005;225:129–40.

52. Santino A, Taurino M, De Domenico S, Bonsegna S, Poltronieri P, Pastor V, et al. Jasmone signaling in plant development and defense response to multiple (al)biotic stresses. Plant Cell Rep. 2013;32:1085–98.

53. You J, Zhang H, Liu N, Gao L, Kong L, Yang Z. Transcriptomic responses to aluminum stress in soybean roots. Genome. 2011;54:923–33.
62. Wang YS, Yang ZM. Nitric oxide reduces aluminum toxicity by preventing oxidative stress in the roots of Cassia tora L. Plant Cell Physiol. 2005;46:1915–23.

63. Xue YJ, Tao L, Yang ZM. Aluminum-induced cell wall peroxidase activity and lignin synthesis are differentially regulated by jasmonate and nitric oxide. J Agric Food Chem. 2008;56:9676–84.

64. Schaller F, Biesgen C, Müsing C, Altmann T, Weiler EW. 12-Oxophytodienoate reductase 3 (OPR3) is the isoenzyme involved in jasmonate biosynthesis. Planta. 2002;210:979–84.

65. Tani T, Sobajima H, Okada K, Chujo T, Arimura S, Tsutsuji N, et al. Identification of the OsOPR7 gene encoding 12-oxophytodienoate reductase involved in the biosynthesis of jasmonic acid in rice. Planta. 2008;227:517–26.

66. Schilmiller AL, Koo AJ, Howe GA. Functional diversification of acyl-coenzyme A oxidases in jasmonic acid biosynthesis and action. Plant Physiol. 2007;143:812–24.

67. Jiang HX, Tang N, Zheng XS, Li Y, Chen LS. Phosphorus alleviates aluminum-induced inhibition of growth and photosynthesis in Citrus grandis seedlings. Physiol Plant. 2009;137:298–311.

68. Hsu PH. Effect of initial pH, phosphate, and silicate on the determination of aluminum with aluminon. Soil Sci. 1963;96:230–8.

69. Yang LT, Qi YP, Lu YB, Gu P, Sang W, Feng H, et al. iTRAQ protein profile analysis of Citrus sinensis roots in response to long-term boron deficiency. J Proteomics. 2013;93:179–206.

70. Zhou CP, Qi YP, You X, Yang LT, Guo P, Ye X, et al. Leaf cDNA-AFLP analysis of two Citrus species differing in manganese tolerance in response to long-term manganese-toxicity. BMC Genomics. 2013;14:621.