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ABSTRACT

 Aluminum (Al)-induced decrease in leaf hydration has been associated with low gas exchange, especially stomatal conductance (*gs*). However, the mechanisms explaining these responses are 28 unclear. *Citrus limonia* was exposed to 0 and 1480 μ M Al in nutrient solution for 90 days to test whether the low *gs* and leaf hydration in plants exposed to Al is associated with increased 9-*cis*-epoxycarotenoid dioxygenase (NCED) gene expression and abscisic acid (ABA) 31 biosynthesis. Relative leaf water content (RWC), water potential (Ψ w) and gas exchange in the leaves, as well as leaf and root *ClNCED3*, *ClNCED1* and *ClNCED5* expression and accumulation of ABA and its metabolites (phaseic acid, dihydrophaseic acid, (+)-7′-hydroxy- ABA and ABA-β-D-glucosyl ester) were measured. Al up-regulated *ClNCED3* and induced 35 ABA accumulation in the roots before impairments in leaf water status (low Ψw , RWC and *gs*) could be observed. Leaf ABA concentration increased from 7 to 90 days and this could be partially explained by the up-regulation of *ClNCED3*, *ClNCED1* and *ClNCED5* in this organ. Stomatal closure occurred concomitantly with the increase of ABA concentration, and this result provides further evidence of the role of ABA modulation of plant hydration under Al stress.

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- **Keywords:** ABA, aluminum, *Citrus*, plant signalling, water relations, NCED genes
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1. Introduction

 Several studies have shown low stomatal conductance (*gs*) in plants under aluminum (Al) toxicity. In comparison to plants not exposed to Al, *gs* values may reduce by 80% in *Solanum lycopersicum* (Simon et al., 1994) and *Secale cereale* (Silva et al., 2012), 44% in *Zea mays* (Anjum et al., 2016), 38% in *Hordeum vulgare* (Ali et al., 2011), 30% in *Citrus reshni* ('Cleopatra' tangerine) (Chen et al., 2005b), 40% in *C. grandis* ('Sour Pummelo') (Jiang et al., 2008) and 50% in *C. limonia* ('Rangpur' lime) (Silva et al., 2018).

 As *gs* is controlled tightly by plant water status (Dodd et al., 2003; Huber et al., 2019), one explanation for the low *gs* in plants exposed to Al could be the inhibition of root growth (Kopittke et al., 2008; Singh et al., 2017; Silva et al., 2019). Al-induced reduction in *gs* is considered an indirect (long distance) effect of Al because it is nearly all retained in negatively charged pectin nets of root cells (Kopittke et al., 2015). Thus, Al toxicity results in lower root surface area (Panda et al., 2009; [Szatanik-Kloc,](http://www.international-agrophysics.org/Author-Alicja-Szatanik-Kloc/111503) 2016), reduction in water uptake, water deficiency in the shoot (Tamás et al., 2006; Yang et al., 2013) and low *gs* (Vitorello et al., 2005). However, most studies in which Al induced low *gs* (Simon et al., 1994; Jiang et al., 2008; Ali et al., 2011; Silva et al., 2012; Banhos et al., 2016; Silva et al., 2018; Cavalheiro et al., 2020) were conducted with plants growing directly on nutrient solution, where water is constantly available. In addition, fibrous xylem vessels (Banhos et al., 2016), more lignin deposition (Silva et al., 2019) and structural damage (Batista et al., 2013) to the root vascular cylinder have been observed in plants under Al toxicity. Aluminum also decreases root hydraulic conductivity in maize plants maintained in nutrient solution (Gunsé et al., 1997), although this study did not measure *gs* and neither associated both variables. Furthermore, low 67 leaf water potential (Ψ w) and decreased relative leaf water content (RWC) can be observed in plants exposed to Al, even when the plants are grown directly on nutrient solution (Banhos et al., 2016; Silva et al., 2018; Cavalheiro et al., 2020).

 Besides low root growth and compromised plant hydraulics, root-to-shoot chemical signalling could also explain low *gs* in plants exposed to Al in nutrient solution. For example, abscisic acid (ABA) signalling controls *gs* when roots are exposed to Al, but only few studies have examined the role of ABA in plants exposed to Al, and these studies have focused on the role of ABA in root Al resistance, without measuring *g^s* (Shen et al., 2004; Hou et al., 2010; Reyna-Llorens et al., 2015; Kopittke, 2016). Recent evidence showed that decrease in root hydraulic conductance and increase in ABA could explain Al-induced stomatal closure in tomato plants (Gavassi et al., 2020). ABA strongly controls stomatal movement (Zhang and Davies, 1989; Merilo et al., 2015), and stomatal closure is one of the most studied roles of ABA in response to drought, high temperature and salt stress (Xiong and Zhu, 2003; Mehrotra et al., 2014). In plants under drought, ABA rapidly accumulates causing low *gs* to reduce transpiration (Zhang et al., 2008; Estrada-Melo et al., 2015). Cellular ABA concentration 82 continuously fluctuates, enabling plants to grow while coping with stressful conditions (Ma et 83 al., 2018). ABA concentration is regulated by its biosynthesis (Ng et al., 2014), which 84 originates from the cleavage of xanthophyll precursors, and also its degradation (Xu et al., 85 2013). The main oxidative route of ABA catabolism is the 8' hydroxylation, which produces 86 8'-hydroxy-ABA (Cutler and Krochko, 1999). This compound isomerizes to phaseic acid (PA), which may be reduced to dihydrophaseic acid (DPA) (Okamoto et al., 2009). A minor 88 oxidative route produces (+)-7'-hydroxy-ABA (7'OHABA), whereas a minor reductive 89 pathway produces an unstable 1',4'-diol ABA. ABA and its metabolites may also be conjugated with glucose to form ABA-β-D-glucosyl ester (ABA-GE) (Zaharia et al., 2005).

 The enzyme 9-*cis*-epoxycarotenoid dioxygenase (NCED) catalyzes the rate-limiting step in ABA biosynthesis (Thompson et al., 2000). This gene encoding NCED form a small multigene family containing five members (*NCED2*, *3*, *5*, *6*, *9*) in *Arabidopsis thaliana* (Tan et al., 2003), and *NCED3* is mainly responsible for ABA accumulation under drought stress in *Arabidopsis* (Iuchi et al., 2001). *NCED3* has been demonstrated to act with *NCED5* against drought stress (Frey et al., 2012). Over-expression of *NCED* leading to ABA over- accumulation was first achieved in tomato and tobacco (Thompson et al., 2000); in tomato this increased root hydraulic conductivity and lowered *gs*, resulting in higher water use efficiency (Thompson et al., 2007). In *Citrus*, when an *NCED3* ortholog, *CrNCED1*, was isolated from 'Cleopatra' mandarin (*Citrus reshni*) and overexpressed in transgenic tobacco, the plants showed higher levels of ABA and enhanced tolerance against drought, salt, and oxidative stresses when compared with WT (Xian et al., 2014). Although *NCED* genes have been well characterized in model plants under water deficiency (Xian et al., 2014), their responses in plants exposed to Al remain unknown.

 As Al reduces *gs* by a mechanism not yet elucidated, it is possible that Al toxicity alters *NCED* expression and the plant accumulates more ABA when compared to those not exposed to Al. Here we tested whether 1480 μ M Al (40 mg L⁻¹), an Al concentration that reduces g_s in *Citrus* plants (Banhos et al., 2016; Cavalheiro et al., 2020), also up-regulates *NCED1*, *NCED3* and *NCED5*, and promotes ABA accumulation in roots and leaves of *C. limonia*. Furthermore, we sought to elucidate if ABA biosynthesis is induced before or after the reduction in leaf 111 hydration, evidenced by low Ψ_w , RWC and g_s .

2. Material and methods

2.1. Plant material and experimental conditions

 Seventy-two three-month-old and 15 ± 1 cm-high 'Rangpur' lime plants (*Citrus limonia* L.) were used for studying the plant hydration capacity when subjected to Al within a 90-day period. This species is an important rootstock for rain-fed *Citrus* plantations due to its high drought resistance (Banhos et al., 2016). At the beginning of the study, the plants had five leaves and were grown directly on an aerated nutrient solution inside opaque plastic boxes (50 cm in length x 30 cm in width x 15 cm in height; 20 L), with six plants per box, in a greenhouse.

121 The nutrient solution used was based on the solution proposed by Clark (1975), and it has been used to test Al tolerance in *C. limonia* (Banhos et al., 2016; Silva et al., 2018; Silva 123 et al., 2019; Cavalheiro et al., 2020). It contained 1372.8 μ M Ca(NO₃)₂, 507.0 μ M NH₄NO₃, 124 224.4 μM KCl, 227.2 μM K₂SO₄, 218.6 μM KNO₃, 483.2 μM Mg(NO₃)₂, 12.9 μM KH₂PO₄, 125 26.0 μ M FeSO₄, 23.8 μ M NaEDTA, 3.5 μ M MnCl₂, 9.9 μ M H₃BO₃, 0.9 μ M ZnSO₄, 0.2 μ M 126 CuSO₄ and 0.4 μ M NaMoO₂. In previous studies we noted an Al-induced decrease in gas exchange rates when *C. limonia* was exposed to 1480 μ M Al (40 mg L⁻¹) (Banhos et al., 2016; Silva et al., 2018; Cavalheiro et al., 2020). Therefore, the solution contained the aforementioned macro and micronutrients, as well as 0 and 1480 µM Al provided through AlCl3. The nominal chemical composition of this solution was also tested on Geochem-EZ 131 software (Shaff et al., 2010), resulting in more than 85% free Al^{3+} available. The pH of the 132 solution was measured daily and maintained at 4.0 ± 0.1 (using NaOH and/or HCl) to guarantee Al solubility. The solution was totally replaced every 15 days, and the treatment with no added Al contained only trace amounts of Al.

 Expanded polystyrene (Isopor®) 50 x 30 cm plates (2cm thick), with six holes (2.5 cm in diameter) each, were floated on the nutrient solution in the boxes, and the plants were fixed in these holes with polyurethane foam strips that were placed around the plant collar. The boxes with six plants each were randomly arranged on benches (80 cm above the ground) inside a 139 greenhouse with semi-controlled conditions (air temperature $28.0 \pm 2^{\circ}$ C; relative humidity 65.3 $\pm 2.5\%$ — between 9h and 11:30h). The photoperiod of 13h of natural sunlight measured inside 141 the greenhouse provided a photosynthetic photon flux density (PPFD) of $862.7 \pm 184.4 \mu$ mol 142 photons $m^{-2} s^{-1}$, between 9h and 11:30h.

2.2. Experimental design

145 After transplant, six boxes (36 plants) remained with the nutrient solution containing 0 146 μ M Al and six other boxes (36 plants) received the nutrient solution containing 1480 μ M Al. The plants grew in these conditions for 90 days, and non-destructive measurements (leaf gas 148 exchange) and destructive measurements (leaf water potential (Ψ_w) , relative leaf water content (RWC), NCED expression and ABA and its metabolites) were performed at 1, 7, 15, 30, 60 150 and 90 days after treatment (DAT). Using predawn (Ψ_{nd}) and midday (Ψ_{md}) leaf water potential and transpiration rates measured in the afternoon, we also estimated the hydraulic conductivity 152 from roots to the leaves (K_L) .

153 The excision of leaves for measuring Ψ_w , leaf discs for RWC and the collection of root tips for gene expression analysis were not performed on the same plants used for measuring leaf gas exchange, so that harmed plants did not interfere in the results of gas exchange rates. 156 Leaf discs for RWC and the collection of leaf pieces and root tips were performed within 60 s, so that these variables interfered as little as possible in each other. In addition, the present study did not involve repeated measurements on the same plants through time, as one box (with 6 plants) per treatment was used on each evaluation date. The leaf pieces and root tips were collected, and their RNA was extracted for measuring NCED gene expression. Using six extra plants (0 DAT and 90 DAT), the biomass of leaves, stems, roots and the total plant biomass were assessed in order to check the severity of the Al treatment.

2.3. Analysis

2.3.1. Leaf gas exchange

166 CO₂ assimilation (*A*; μ mol m⁻² s⁻¹) and transpiration (*E*; mmol m⁻² s⁻¹) rates, stomatal 167 conductance $(g_s; \text{mol m}^{-2} \text{ s}^{-1})$ and intercellular CO_2 (*Ci*; µmol mol⁻¹) were measured using an open gas exchange system (LI-6400xt; LI-COR, Lincoln, NE, USA). The water use efficiency 169 (*WUE* = A/E) and intrinsic water use efficiency (*iWUE* = A/gs) were also calculated. The CO₂ 170 concentration entering the leaf cuvette (LCF chamber; 2 cm^2 , LI-COR) averaged 400 µmol 171 mol⁻¹, as provided by the 6400-01 CO₂ mixer (LI-COR), as this was the air CO₂ concentration accepted for the experimental site when the study was performed. Measurements were taken between 9h and 11:30h on cloudless days, as it is the best period for measuring gas exchange parameters (Feistler and Habermann, 2012). We also measured gas exchange in the afternoon (13h-15h) in order to calculate the estimated hydraulic conductivity from roots to the leaves 176 (K_L) .

177 The PPFD in the leaf cuvette was provided by an artificial LED light source (6400-40) 178 LCF, LI-COR), which was set to provide 90% red and 10% blue light at 1500 µmol photons m^{-2} s⁻¹, as this value saturates *A* for *C. limonia* as observed in *A/Ci* curves (Silva et al., 2018). The vapor pressure deficit (VPD) inside the leaf cuvette was similar to the external environment (inside the greenhouse), which was not higher than 1.5 kPa and relative humidity was approximately 65% on the days of measurement.

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184 **2.3.2. Water relations**

185 Ψ_{pd} and Ψ_{md} (under maximum VPD) were measured (MPa) by the pressure chamber method 186 (Turner, 1981), using a 3005F01 Plant Water Status Console (Soil Moisture, Santa Barbara, 187 CA, USA) chamber.

The estimated hydraulic conductivity from roots to the leaves $(K_L; \text{ mmol H}_2\text{O m}^{-2} \text{ s}^{-1})$ 189 MPa⁻¹) was determined by the method proposed by Hubbard et al. (2001) , which is based on 190 Ohm's Law. For this, the following equation was applied:

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191 \t KL = l
$$

 E_{14h} / (Ψ_{pd} – Ψ_{md}),

192 where *E*14h is the transpiration rate (*E*) measured between 13:00h and 15:00h under maximum 193 VPD; Ψ_{pd} is assumed as the soil water potential (Ψ_{soil}), and Ψ_{md} is Ψw measured under 194 maximum VPD. Although the plants were grown directly on nutrient solution, the $\Psi_{\text{pd}} = \Psi_{\text{soil}}$ 195 principle is still accepted because Ψ_{pd} is measured before sunrise in non-transpiring plants and, 196 therefore, $\Psi_{\rm pd}$ represents the plant's capacity to rehydrate overnight (Turner, 1981). This 197 method was previously used for measuring *K*L in *Citrus sinensis* (Magalhães Filho et al., 2009) 198 and *C. limonia* (Cavalheiro et al., 2020) grown on nutrient solution.

199 For measuring RWC (%), leaf discs were collected at 13h-15h from plants of both 200 treatments and calculated as:

201 RWC = $[(FM-DM)/(TM-DM)] \times 100$,

 where FM is the fresh mass (immediately measured after collection); TM is the turgid mass, measured after rehydrating samples for 24 h in 100 mL deionized water inside amber flasks (to avoid photosynthetic activity); and DM is the dry mass, measured after oven-drying the 205 discs at 60°C for 48 h, according to Silva et al. (2018).

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207 **2.3.3. Gene expression analysis**

208 Leaf pieces ($\approx 1 \text{ cm}^2$) or root pieces (1 cm in length), totaling 100 mg (fresh mass) for 209 each leaf or root sample $(n = 4)$, were collected at 13h-15h, frozen in liquid nitrogen and stored at -80ºC for future analysis. Total RNA was extracted from leaf and root samples using the 211 RNeasy Plant Mini Kit (Qiagen, Hilden, Germany). Total RNA (2 µg) was treated with RNase- free TURBO DNase (Ambion, Carlsbad, USA) and reverse transcribed to cDNA using oligo- dT and the GoScript Reverse Transcription System kit (Promega Corp., Madison, WI, USA), according to the manufacturer's protocol (Life Technologies, Carlsbad, CA, USA). Gene expression analysis was carried out by quantitative real-time PCR (qRT-PCR) with SYBR green GoTaq q-PCR Master Mix (Promega Corp., USA), using Applied Biosystems QuantStudio 3 (Life Technologies, Carlsbad, CA, USA). The primers of *ClNCED1*, *ClNCED3* and *ClNCED5* (Table 1) used in the experiment were previously used in *Citrus* species (Agusti et al., 2007, Bassene et al., 2009), including *C. limonia* (Neves et al., 2013). As reference genes, we used glyceraldehyde-3-phosphate dehydrogenase (*GAPC2*) and elongation factor 1-alpha (*EFα*) (Table 1), which were proposed by Mafra et al. (2012) and used previously by Silva et 222 al. (2019). Amplification efficiencies were calculated for each primer using Miner software (Zhao and Fernald, 2005).

- For calculating the relative expression, Ct (cycle threshold) values of each sample, which were determined by the average of three technical replicates, were converted into relative quantities (RQ) according to Pfaffl (2001), using the following equation:
- 227 $RQ = E\Delta Ct$,

228 where E is the primer efficiency, and Δ Ct is the difference between control Ct value for the evaluated gene and Ct value of the given sample. A normalization factor (NF) for each sample was calculated by the geometric mean of the RQ values of *GAPC2* and *EFα*. Normalized- relative quantity (NRQ) of each sample was calculated as the ratio of the sample RQ and the appropriate NF. Individual fold change values were determined by dividing the sample NRQ by mean values of NRQ that were obtained from the calibrator, i.e., root samples of plants not exposed to Al. Following this, fold change in the control group always shows a mean value of 1. Four independent biological replicates (plant samples) were used to calculate mean for each time point and treatment combination.

2.3.4. Quantification of ABA and metabolites

 ABA and its metabolites, phaseic acid (PA), dihydrophaseic acid (DPA), (+)-7′- hydroxy-ABA (7′-OH ABA) and ABA-β-D-glucosyl ester (ABA-GE) were analyzed via liquid chromatography/tandem mass spectrophotometry (LC/MS-MS) on a SciexExionLC coupled 242 with a QTRAP 6500+ mass spectrophotometer, following the method proposed by Morris et al. (2019).

 Leaf and root samples (500 mg DM) were ground to a powder (in 2 mL micro- centrifuge tubes containing two 5 mm acid-rinsed glass balls) in a Star-Beater (VWR) at 30 246 Hz for 2 min. The powdered material, \sim 20 mg for leaf samples and \sim 50 mg for roots, with 1 ng of internal standards added, were extracted using a Star-Beater at 30 Hz for 2 min with 1 mL of ice-cold methanol/formic acid/water solvent (60:5:35 v/v). The internal standards used 249 were: [²H₄]-abscisic acid (-)-5,8',8',8' (d₄-ABA); [²H₃]-phaseic acid (-)-7',7',7'(d₃-PA); [²H₅]-250 abscisic acid glucose ester $(+)$ -4,5,8',8',8' (d₅-ABA-GE); [²H₃]-dihydrophaseic acid $(-)$ -7',7',7' 251 (d₃-DPA); and $[^{2}H_{4}]$ -7'-hydroxy-abscisic acid (\pm)-5,8',8',8' (d₄-7OH-ABA). After extraction, the samples were left on ice in the dark for 20 min and, subsequently, the plant material was 253 pelleted by centrifugation at $24,000 \times g$ at 4°C for 10 min. The supernatant was pipetted into 15 mL conical centrifuge tubes and the solvent was evaporated overnight in a freeze-drier (- 105˚C; Scanvac, CoolSafe 110-4 Pro). Samples were reconstituted (vortexed for 2 min at 1400 rpm, sonicated for 30s and a final vortex at 2000 rpm for 3 min) in 1 mL of 5% acetonitrile in 10 mM ammonium formate (pH 3.4, adjusted with formic acid). Finally, the samples were 258 filtered through 4 mm nylon filters $(0.2 \mu M)$ pore size, Whatman) into silanised amber HPLC vials.

 Calibration samples consisted of a series of non-deuterated compounds (from 0.1 to ng mL⁻¹), each with deuterated compounds (constant 1 ng mL⁻¹). Extracts and calibration 262 samples (20 μ L) were injected with an auto-sampler into a Luna 3 μ m C18(2) 100 x 2 mm 263 (Phenomenex) column with guard column at 30° C. The aqueous mobile phase (A) consisted of 2% acetonitrile in 10 mM ammonium formate, and the organic mobile phase (B) was 95% 265 acetonitrile and 0.1% formic acid. The ratios of mobile phase A:B for separation of compounds 266 was as follows (at a flow rate of 600 μL min⁻¹): 0-1.5 min at 96:4; 1.5-7 min at 87.4:12.6 and 7-10 min at 74:26. The column was then cleaned as follows, 10-10.5 min at 60:40; 10.5-10.6 min at 50:50 and 10.6-11.6 min at 0:100. The column was then equilibrated from 11.6-13 min at 96:4 before injection of the next sample. Analyst 1.6.3 and MultiQuant 3.0.2 (Sciex, Singapore) software was used for acquisition and quantification, respectively.

2.3.5. Biometric parameters

 At 0 and 90 DAT, after separating the plant parts into leaves (plus petioles), stems and roots, the number of leaves was counted, and the shoot length was measured with a ruler. The total leaf area per plant (LA) was measured with an area meter (LI-3100C, LI-COR, USA). Total root length, root surface area and root diameter were measured using a scanner (Epson perfection v700 photo, Suwa, Japan), which was coupled to a computer running the 278 WinRHIZOTM software (Regent Instruments, Canada). The biomass of organs was measured 279 on a 0.001g precision scale (AR2140, OHAUS, USA) after oven-drying the samples at 60° C until constant mass.

2.3.6. Aluminum quantification

 Dry samples of leaves and roots were sent to a plant nutrition laboratory at University of São Paulo (ESALQ, USP, Piracicaba, Brazil) where these were ground and digested in a solution of sulfuric:nitric:percloric acids (1:10:2, v/v/v). After digestion, Al concentrations 286 were determined by the atomic absorption spectrophotometer method (Sarruge and Haag, 1974) and expressed as mg Al per kg dry mass.

2.3.7. Data analysis

 Leaf gas exchange parameters (*A*, *gs*, *E*, *Ci*, *WUE* and *iWUE*), RWC and biomass of 291 organs were measured using six plant replicates. Leaf water potential (Ψ_{pd} and Ψ_{md}), estimation 292 of hydraulic conductivity from roots to the leaves (K_L) , gene expression of NCED and ABA metabolites were assessed using four plant replicates.

294 A student's t-test (∞ = 0.05) was used, separately for each evaluation date (1, 7, 15, 30, 295 60 and 90 DAT), to test differences between 0 and 1480 μ M Al for each variable, as well as when testing differences in plant biomass and Al concentration in plant organs between 0 and 90 DAT within each treatment.

3. Results

3.1. Biometric parameters

 As expected, Al reduced the size of plants (Supplementary material; Fig. S1). At 90 DAT, Al significantly limited the main root length (-48%) (Fig. 1A), root surface area (-62%) (Fig. 1B) and root biomass (-65%) (Fig. 1D), while the root diameter was enhanced in plants exposed to Al (+25%) (Fig. 1C).

 From 0 to 90 DAT, the leaf number (Fig. 2A), leaf area (Fig. 2B) and leaf biomass (Fig. 2C) increased by 31%, 83% and 59%, respectively, in plants exposed to Al and 140%, 504% and 393%, respectively, in control plants. At 90 DAT, Al significantly decreased the leaf number (-45%) (Fig. 2A), leaf area (-70%) (Fig. 2B) and leaf biomass (-68%) (Fig. 2C). Thus, Al inhibited root growth, leaf initiation, leaf expansion and organ biomass accumulation, but caused root thickening.

3.2. Leaf gas exchange

 Compared to plants not exposed to Al, values of *A* (Fig. 3A), *gs* (Fig. 3B) and *E* (Fig. 3C) decreased from 7 DAT, and at 90 DAT these parameters were 71%, 78% and 60% lower in plants exposed to Al. On the other hand, *Ci* values increased in plants exposed to Al from 15 DAT, being 55% higher at 90 DAT (Fig. 3D). The *WUE* was the same between the treatments throughout the study (Fig. 3E), while *iWUE* was higher in plants exposed to Al from 30 DAT, being 108% higher at 90 DAT (Fig. 3F). This data is consistent with stomatal closure leading to reductions of *A*. That is, a larger effect of Al toxicity on *gs* than on *A* resulted in an increase in *A*/*gs* (*iWUE*).

3.4. Water relations

 \mathcal{Y}_{pd} was lower in plants exposed to Al throughout the study, although this was not statistically significant (Fig. 4A). However, plants exposed to Al showed significantly lower \mathcal{H}_{md} (Fig. 4B) and RWC (Fig. 5A) when compared to control plants from 7 DAT onwards. At 326 90 DAT, Al reduced Ψ_{md} from -1.2 to -2.2 (Fig. 4B) and RWC from 89% to 67% (Fig. 5A). *K*L was also lower in plants exposed to Al from 7 DAT, being 80% lower than control plants at 90 DAT (Fig. 5B). Thus, Al compromised plant water status.

3.5. NCED gene expression

 In the leaves, Al enhanced NCED genes expression over time (Fig. 6A, 6C and 6E), while in the roots Al caused a peak for *ClNCED5* (Fig. 6F) and *ClNCED3* (Fig. 6D), most pronounced in the latter. The Al-induced up-regulation of leaf *ClNCED3* started at 15 DAT, being 78-fold higher at 90 DAT (Fig. 6C). For leaf *ClNCED1* and *ClNCED5*, significant up- regulation occurred at 60 and 90 DAT, and on these dates up-regulation was of approximately 80- (Fig. 6A) and 35-fold higher (Fig. 6E) than the control, respectively. In the roots, Al caused up-regulation of *ClNCED3* on all dates except for 7 DAT, reaching a peak of 16-fold higher than control plants at 30 DAT (Fig. 6D). Root *ClNCED5* was significantly up-regulated by Al (4-fold higher) only at 60 DAT, while no up-regulation in root *ClNCED1* was induced by Al (Fig. 6B). Thus, Al up-regulated the key genes of ABA biosynthesis (*ClNCED3*) in the roots at 1 DAT (Fig. 6D), while decreases in leaf hydration were detected at 7 DAT for *gs* (Fig. 3B), \mathcal{H}_{md} (Fig. 4B), RWC (Fig. 5A) and K_L (Fig. 5B).

3.6. Abscisic acid (*ABA*) *accumulation in leaves and roots*

 In general, Al increased ABA concentrations in leaves and roots (Fig. 7A, 7B). In plants exposed to Al, [ABA]leaf increased from 7 DAT, being 4.7-times higher than the control at 90 DAT (Fig. 7A). [DPA]leaf and [7ˊOH ABA]leaf increased in plants exposed to Al from 15 DAT, being 1.3- and 1.5-times higher than the control, respectively, over this period (Fig. 7E, 7G). [PA]leaf and [ABA-GE]leaf, however, were higher in plants exposed to Al only at 90 DAT, being 2.6- and 2.0-times higher, respectively, when compared to the control (7C, 7I). Therefore, Al caused a consistent increase in [ABA], [DPA] and [7ˊOH ABA] from the first week of the study, while [PA] and [ABA-GE] increased only after 90 days of Al exposure.

 In the roots, Al caused a peak of [ABA]root (3-times higher than the control; Fig. 7B), [PA]root (7-times higher; Fig. 7D) and [ABA-GE]root (3.3-times higher; Fig. 7J) at 7, 1 and 30 DAT, respectively. After these peaks, the concentration of these metabolites in the roots decreased, but remained higher in plants exposed to Al at 15 and 30 DAT (ABA; Fig. 7B), 30 357 and 60 DAT (PA; Fig. 7D) and until 90 DAT (ABA-GE; Fig. 7J). [DPA]_{root} of plants exposed to Al was 2.0-times higher than control plants only at 90 DAT (Fig. 7F), while [7ˊOH ABA]root showed no pattern, with variable values between treatments (Fig. 7H).

 Thus, Al increased ABA in the roots immediately after Al exposure and, in the leaves, Al induced a consistent accumulation, especially for ABA, DPA and 7ˊOH ABA. In the leaves of plants exposed to Al, ABA accumulation seems to be associated with *ClNCED3* (Supplementary material; Fig. S2). In addition, in Al-treated plants, [ABA]leaf is driving major part of *gs* responses (Supplementary material; Fig. S3), which also corroborates low values of 365 RWC, Ψ_{md} and K_{L} of plants exposed to Al.

3.7. Aluminum retention in plant organs

 As expected, Al concentration in the roots was approximately 10 times higher than the leaves of plants exposed to Al (Fig. 8). From 0 to 90 DAT, leaf and root Al concentration increased by seven- (Fig. 8A) and 15-times (Fig. 8B), respectively, in control plants when compared to those treated with Al.

4. Discussion

 Hydraulic signals, in the form of turgor changes in the leaves, and hormonal signaling have been proposed to control *g^s* (McAdam and Brodribb, 2015; Huber et al., 2019). Plant hormones can influence the Al toxicity and development of symptoms (Kopittke, 2016), as well as these compounds can mediate the Al resistance, especially in the root environment

 (Massot et al., 2002; Yang et al., 2017). The well-known role of ABA in causing stomatal closure, the up regulation of ABA biosynthesis upon changes in cell turgor and water availability (McAdam et al., 2016; Sussmilch et al., 2017; Zhang et al., 2018) makes ABA a candidate for causing the decrease in *gs* during Al toxicity. In the present study, we tested whether ABA accumulation in roots and leaves could be responsible for the Al-induced low *g^s* usually found in *Citrus limonia* exposed to Al (Banhos et al., 2016; Silva et al., 2018; Cavalheiro et al., 2020).

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4.1. Effect of Al on plant water relations

 Our results show that *A* and *E* were progressively reduced in plants exposed to Al (Fig. 3A and 3C), and these reductions could be explained by the low *gs* values (Fig. 3B). In our previous studies with this same species under the same Al concentration, low *A* values were largely explained by low *gs* rather than decreased photochemical performance (Banhos et al., 2016; Silva et al, 2018). In addition, Al-induced reduction in *gs* has been observed in other *Citrus* plants, including 'Cleopatra' tangerine (-30%; Chen et al., 2005b) and 'Sour Pummelo' (-40%; Jiang et al., 2008). Therefore, the decrease in *gs* seems to be a key response in plants exposed to Al.

 Plants adjust their xylem pressure with concomitant stomatal regulation (Creek et al., 2020; Rodriguez-Dominguez and Brodribb, 2019). In the present study, the decrease in *gs* of plants exposed to Al was not sufficient to maintain the leaf water status, as evidenced by low 398 values of Ψ_{md} (Fig. 4B) and RWC (Fig. 5A) from 7 DAT. K_L represents the plant capacity to supply water to the mesophyll (Rodríguez-Gamir et al., 2019) and since its value dropped by 80% under Al toxicity (Fig. 5B), the ability of the Al-treated plants to transport water to the leaves was dramatically impaired. Indeed, root hydraulic conductance (*Lpr*) of *Solanum lycopersicum* (tomato) also declined proportionally to the increase of Al in nutrient solution (Gavassi et al., 2020). Reductions in *KL* and low expression of aquaporins (PIP family) in *C. limonia* exposed to 1480 μM Al found by Cavalheiro et al. (2020) have been associated with fibrous xylem vessels (Banhos et al., 2016) and more lignin deposition in the vascular cylinder (Silva et al., 2019) of *C. limonia* grown under the same Al toxicity conditions. The root apex senses Al toxicity (Ryan et al., 1993; Horst et al., 2010), and, despite being anatomically "disconnected" from the xylem, the longer the exposure of root tips of *C. limonia* to Al, the more lignin deposition is found in their vascular cylinders (Silva et al., 2019), although the mechanism(s) of signaling between Al perception and xylem damage is unclear. The vascular cylinder was also the most affected part of the root of maize plants, and their proto- and 412 metaxylem did not reach full maturation under $300 \mu M$ Al (Batista et al., 2013). In addition, 413 ten-times more Al was found in root tips (1 cm long) of maize plants exposed to 50 μ M Al 414 when compared to plants not exposed to Al after 24h (Souza et al., 2016). This same proportion was found in the roots of plants exposed to Al when compared to control plants, at 90 DAT (Fig. 8B). Furthermore, once Al is firmly bound to a root cell wall, where it is the site of primary lesion (Kopittke et al., 2015), it does not seem to be released (Rangel et al., 2009), and it could cause anatomical damage to the cortex and xylem of plant roots, as observed by Batista et al. (2013), Banhos et al. (2016) and Silva et al. (2019). Taken together, these results suggest that Al impairs the plant capacity to transport water to the leaves. A key question, however, is whether the impairment of root and vascular function leads directly to declining shoot water status and productivity (reduced *gs*, *A* and biomass), or whether this is controlled by early hormonal signals.

4.2. The role of ABA and its metabolites in short-term responses

 In the present study, root ABA increased at 1 DAT (Fig. 7B) relative to control, prior 427 to any significant decreases in Ψ_{pd} , Ψ_{md} , RWC or K_L (Fig. 4 and 5); and ABA kept increasing in the root until 7 DAT (Fig. 7B). PA concentration was also higher at 1 DAT, compared to control plants (Fig. 7D); this could also contribute to physiological responses because PA showed biological activity *in vitro*, activating members of the ABA receptor family, albeit with a lower affinity than ABA (Weng et al. 2016). Although not presenting biological activity, ABA-GE was higher in the roots of plants exposed to Al on all dates, including 1 DAT, although showing a peak at 30 DAT (Fig 7J). ABA-GE is considered an ABA metabolite and can be transported symplastically from the cytosol of root cells to xylem parenchyma cells and 435 be released to xylem vessels (Priest et al., 2006) as a root-to-shoot signal (Sauter et al., 2002). Therefore, the extra ABA-GE produced from 1 DAT in the Al-treated roots could have been transported, de-glycosylated and have contributed to the ABA level and stomatal closure at 7 DAT. Analysis of gene expression in the early period 1-7 DAT indicated that *ClNCED3* responded to the Al treatment in the roots, but the increase was small (Fig. 6D). This small increase could have contributed to the rise in ABA, PA and ABA-GE. Alternatively, an activation of NCED enzyme activity at the protein level, or a change in transport processes between root and shoot might explain the early spike in ABA given that there was only a small increase in *ClNCED3* gene expression. Furthermore, the catabolic product PA increased at 1

 DAT (Fig. 7D), suggesting that slower ABA catabolism was not the reason for ABA accumulation in the roots of Al-treated plants.

 Overall, the very early rise in root ABA, PA and ABA-GE caused by Al toxicity appeared to precede the decline in *gs*, and so is potentially the cause of stomatal closure. However, this reduction in *gs*, preventing water loss, was not able to stop a decline in leaf water status, presumably caused by the impact of Al on root water transport function as suggested by Batista et al. (2013), Banhos et al. (2016), Silva et al. (2019), Cavalheiro et al. (2020) and Gavassi et al. (2020), and also supported by the 80% lower *K*L in plants exposed to Al (Fig. 5B). The coincident reduction in *gs* and shoot water status at 7 DAT means that we cannot exclude a hydraulic signal as the cause of stomatal closure.

4.3. The role of ABA and metabolites in longer-term responses

 In the leaves of plants exposed to Al, ABA progressively increased until the end of the study (Fig. 7A) with the first significant increase occurring at 7 DAT. This was accompanied 458 by similar trends of DPA from 15 DAT (Fig. 7E), but surprisingly [DPA]_{leaf} peaked 25-fold higher in absolute concentration than [ABA]leaf, suggesting a high rate of catabolism to DPA that has previously been observed in *Citrus* leaves under drought and soil flooding stresses (Jitratham et al., 2006; Arbona et al., 2017). ABA accumulation in the leaf occurred later and more progressively than in the root, and after, or co-incident with, the decline in water relations (RWC, Ψmd and *KL* all down at 7 DAT). Therefore the leaf ABA and PA was likely increased as a secondary consequence of the Al toxicity, where lack of water supply to the shoot (low *K*L) led to reduced shoot water status which then stimulated accumulation of ABA and PA and 466 reinforced stomatal closure: in plants exposed to Al, [ABA]_{leaf} was inversely correlated with *gs* values (Supplementary material; Fig. S3).

 The expression of genes encoding NCED did not fully explain the increase in ABA and DPA in the leaf. The first increase was for *ClNCED3* at 15 DAT, after the rise in ABA, and this initial increase for *ClNCED3* was small (2-fold), with the peak occurring at 90 DAT (Fig. 471 6C). However, the linear correlation (R^2) between *CINCED3* and [ABA]_{leaf} in plants exposed to Al was 0.875 (Supplementary material; Fig. S2). This suggests that progressive increase in [ABA]leaf may be explained by *ClNCED3* expression. In the leaf, *ClNCED1* and *ClNCED5* also showed up-regulation of approximately 80- and 30-fold higher than control plants, respectively, at 60 and 90 DAT (Fig. 6A and 6E), but this rise was even later than for *ClNCED3*; therefore it did not explain the initial increase in ABA from 7 DAT in the leaf, but may have contributed to the continued and accelerated increase in ABA and PA which showed a sharp rise between 60 and 90 DAT. The increase in *ClNCED3* expression from 15 DAT, and of *ClNCED1* and *ClNCED5* from 60 DAT, was probably driven by reduced cellular water status in the leaf since orthologs of this gene are known to respond in this way. In *Arabidopsis*, the orthologous *AtNCED3* is predominantly induced by drought and controls endogenous ABA content in this condition (Endo et al., 2008; Hao et al., 2009), but *AtNCED5* and *AtNCED3* participate together in water deficit response (Frey et al., 2012). In *Citrus*, *NCED1* was up- regulated by drought in leaves of *C. sinensis* (Rodrigo et al., 2006; Xian et al., 2014) and *C. reshni* (Zandalinas et al., 2016), as also observed here in the leaves of plants exposed to Al (Fig. 6A). Up-regulation of *ClNCED5* was previously observed in leaves of *C. limonia* 487 submitted to 40 days of drought (Neves et al., 2013).

 Thus, only the later, but not the early increase in ABA and DPA, could be explained by *NCED* gene expression in the leaves; other mechanisms, such as reduced catabolism, post-transcriptional control or redistribution of ABA would need to be invoked.

4.4. Impact of reduced growth rate on physiological responses

 Critics could still argue that the conspicuous decrease in root growth parameters caused by Al at 90 DAT (Fig. 1A, 1B and 1D) could have acted as a physical limitation for water uptake, which could not maintain leaf transpiration, explaining the low *gs* values. However, this low root growth was followed by reduced leaf number (Fig. 2A), leaf area (Fig. 2B) and leaf biomass (Fig. 2C), which would have greatly reduced the demand for water transport from 498 the smaller root system. Similarly, tomato plants exposed to 0, 25 and 50 μ M Al showed similar root/leaf area ratio, reinforcing that the decrease in the root size is compensated by a low shoot growth (Gavassi et al., 2020).

 We have measured biometric parameters only at 90 DAT, but it seems unlikely that reduced root growth could have occurred at 7 DAT and have caused low *gs* values due to fewer roots responsible for (less) water uptake. Further evidence in this regard deserves investigation.

5. Conclusions

 We showed that Al triggered *ClNCED3* expression and ABA biosynthesis in the roots 507 1 day after Al exposure in *C. limonia*, before impairments in leaf hydration (low Ψw , RWC and *gs*) could be observed. In addition, leaf ABA concentration increased from 7 to 90 DAT and this could be partially explained by the increased expression of *ClNCED3*, *ClNCED1* and *ClNCED5* in this organ. Stomatal closure occurred concomitantly with the increase of ABA

- concentration and this result provides further evidence of the role of ABA modulation of plant hydration under Al stress.
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Tables

Table 1. List of gene primers used for qRT-PCR analysis in *Citrus limonia*.

Figure legends

Fig. 1. Main root length (A), root surface area (B), root diameter (C) and root biomass (D) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences ($P < 0.05$) between 0 and 1480 μ M Al. For plants not exposed to Al, distinct uppercase letters indicate significant differences (P < 0.05) between 0 and 90 DAT; for plants exposed to Al, distinct lowercase letters indicate significant differences (P < 0.05) between 0 and 90 DAT. Columns are mean values (n = 6, \pm SE).

Fig. 2. Leaf number (A), area (B) and biomass (C) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences ($P < 0.05$) between 0 and 1480 μM Al. For plants not exposed to Al, distinct uppercase letters indicate significant differences ($P \le 0.05$) between 0 and 90 DAT; for plants exposed to Al, distinct lowercase letters indicate significant differences ($P \le 0.05$) between 0 and 90 DAT. Columns are mean values ($n = 6, \pm SE$).

Fig. 3. Leaf gas exchange and water use efficiency of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. (A) $CO₂$ assimilation, (B) stomatal conductance, (C) transpiration, (D) intercellular $CO₂$, (E) water use efficiency and (F) intrinsic water use efficiency. For each evaluation date, asterisks indicate significant differences (*P*< 0.05) between 0 and 1480 μM Al. Circle symbols are mean values ($n = 6, \pm SE$).

Fig. 4. Leaf water potential at predawn (Ψ_{pd}) (A) and midday (Ψ_{md}) (B) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences (*P*< 0.05) between 0 and 1480 μM Al. Circle symbols are mean values ($n = 4, \pm SE$).

Fig. 5. Relative leaf water content (A) and estimated hydraulic conductance from roots to the leaf (B) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences (*P*< 0.05) between 0 and 1480 μM Al. Circle symbols are mean values ($n = 6, ± SE$).

Fig. 6. Foldchange of normalized expression level of *ClNCED1*, *ClNCED3* and *ClNCED5* in leaves (A, C, E, respectively) and root tips (B, D, F, respectively) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences (*P*< 0.05) between 0 and 1480 μM Al.The dotted line represents the control group, showing always the mean value of 1, and foldchange is that between control and Al treatment. Circle symbols are mean values $(n = 4, \pm SE)$.

Fig. 7. Abscisic acid (ABA) and its metabolites in leaves (left columns) and roots (right columns) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences ($P \le 0.05$) between 0 and 1480 μM Al. Circle symbols are mean values (n = 4, \pm SE). (PA: phaseic acid; DPA: dihydrophaseic acid; 7′OHABA: (+)-7′-hydroxy-abscisic acid; ABA-GE: abscisic acid glucosyl ester).

Fig 8. Aluminum concentration in leaves (A) and roots (B) of *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al. For each evaluation date, asterisks indicate significant differences ($P < 0.05$) between 0 and 1480 μM Al. For plants not exposed to Al, distinct uppercase letters indicate significant differences ($P \le 0.05$) between 0 and 90 DAT; for plants exposed to Al, distinct lowercase letters indicate significant differences ($P \le 0.05$) between 0 and 90 DAT. Columns are mean values ($n = 6, \pm SE$).

Appendix A. Supplementary data

Additional supporting information may be found in the online version of this article.

Fig. S1. Morphological details of shoots, leaves and roots of *C. limonia* grown for 90 days in nutrient solution containing 0 (on the left) and 1480 μM Al (on the right).

Fig. S2. Individual readings (replicates; $n = 4$ plants) of leaf abscisic acid concentration ([ABA]leaf) and *ClNCED3* expression (Foldchange) in *C. limonia* grown for 90 days in nutrient solution containing 1480 μM Al.

Fig. S3. Individual readings (replicates; $n = 4$ plants) of leaf abscisic acid concentration ([ABA]leaf) and stomatal conductance (*gs*) in *C. limonia* grown for 90 days in nutrient solution containing 0 and 1480 μM Al.

Highlights:

Aluminum (Al) toxicity inhibits root growth and reduces the stomatal conductance (*gs*) The 9-*cis*-epoxycarotenoid dioxygenase (NCED) enzyme catalyzes the abscisic acid (ABA) Roots of *Citrus limonia* exposed to Al up-regulated *ClNCED3* before *gs* was reduced Up-regulation of *ClNCED3*, *ClNCED1* and *ClNCED5* matched ABA leaf levels Al triggers ABA biosynthesis, which is associated with the low *gs*

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