Expression of Ethylene Biosynthesis and Signaling Genes during Differential Abscission Responses of Sweet Orange Leaves and Mature Fruit

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Abstract. When applying abscission agents to tree fruit to facilitate harvest, it is desirable to loosen fruit and not leaves or other organs, but mechanisms controlling leaf and fruit drop are not fully understood. The effect of 450 μL L−1 ethephon (ethylene-releasing agent) alone or in combination with 1-methylcyclopropene [1-MCP (ethylene perception inhibitor)] on leaf and mature fruit abscission of ‘Valencia’ sweet orange (Citrus sinensis) was studied. Leaf abscission increased and fruit detachment force (FDF) decreased significantly especially 4 days after ethephon treatment. Leaf drop rose to over 80% 7 days after application, whereas FDF was only 30% less than untreated control fruit. When 1-MCP was combined with ethephon and applied to ‘Valencia’ sweet orange canopies, leaf abscission was greatly reduced, but reduction in FDF proceeded unabated. We hypothesized that differential response of ‘Valencia’ sweet orange fruit and leaves to 1-MCP was correlated with expression of ethylene biosynthetic and signaling genes and their downstream action. Partial or full-length nucleotide sequences were obtained for ‘Valencia’ sweet orange homologs of CsACS1, CsETR2, CsETR3, constitutive triple response-1 (CsCTR1), ethylene insensitive-2 (CsEIN2), and ethylene insensitive 3-like-1 (CsEI1L) and 2 (CsEIIL2). Ethephon application increased expression of biosynthesis genes CsACS1 and CsACO and receptors CsERS1 and CsETR2 in the abscission zones of leaves and mature fruit. Ethephon-induced increase in gene expression was completely suppressed by 1-MCP application in all but CsACS1 and CsACO in fruit abscission zones. Although gene expression was suppressed initially, CsACS1 and CsACO expression in fruit abscission zones treated with 1-MCP in the presence or absence of ethephon increased over the 7-day measurement period, suggesting that CsACS1 and CsACO expression were negatively regulated by basal ethylene production in this tissue. However, 1-MCP treatment alone did not loosen fruit, indicating that CsACS1 and CsACO played minor roles in fruit abscission. To determine if the difference in ethylene sensitivity was the basis of differential response to ethephon within the same organ, potted ‘Valencia’ sweet orange plants were treated with ethylene, and rates of blade and petiole drop and detachment forces at the laminar and petiolar abscission zones were studied. Although leaf blades abscised earlier than petioles, the force of detachment was similar, indicating no differences in ethylene sensitivity. Overall, the most significant difference between fruit and leaf abscission zones was seen in the expression of CsACS1 and CsACO genes, but the expression pattern was poorly correlated with abscission.

Abscission is a coordinated process of separation of organs such as leaves, flowers, and fruit from the parent plant. Abscission occurs in abscission zones due to the dissolution of cell walls by the activity of hydrolase enzymes cellulase and polygalacturonase, resulting in organ detachment. Since the initial work of Horton and Osborne (1967) associating cellulase and polygalacturonase, resulting in organ detachment. Since the initial work of Horton and Osborne (1967) associating cellulase activity with abscission of bean (Phaseolus vulgaris) explants, cell wall hydrolase activity has been associated with abscission in other plants, including citrus (Citrus spp.) (Greenberg et al., 1975; Ratner et al., 1969). Ethylene is a plant hormone that plays a major role in regulating the abscission process (Bleecker and Patterson, 1997; Burg, 1968; Jackson and Osborne, 1970). Application of exogenous ethylene hastens abscission by inducing expression of cellulase and polygalacturonase genes in peach (Prunus persica) fruit (Bonghi et al., 1992) and tomato (Solanum lycopersicum) leaves, fruit, and flowers (del Campillo and Bennett, 1996; Kalaitzis et al., 1995). In citrus, gene expression and enzymatic activity of cellulase and polygalacturonase increased during ethylene-induced abscission of mature citrus fruit (Burns et al., 1998; Greenberg et al., 1975; Huberman and Goren, 1979). The regulatory role of ethylene in abscission was demonstrated in transgenic plants. Overexpressing the ethylene biosynthesis gene 1-amino-cyclopropane-1-carboxylate synthase (ACS) in tomato plants resulted in premature flower abscission, while a delay in abscission was noted in Nr plants (mutant of ethylene receptor LeETR3) and in plants expressing antisense transcripts of the ethylene receptor LeETR1 (Lanahan et al., 1994; Whitelaw et al., 2002).

The response or sensitivity to ethylene can be controlled at steps involving ethylene biosynthesis and ethylene perception. Ethylene biosynthesis begins with the conversion of S-adenosylmethionine (SAM) to ACC by ACS. ACC is oxidized to ethylene by ACO. The conversion of SAM to ACC by ACS is considered the rate-limiting step in ethylene biosynthesis, but regulation at the ACO step has also been reported (Alexander and Grierson, 2002). The structure of ACS is similar to the subgroup I family of pyridoxal 5'-phosphate (PLP)-dependent aminotransferases and PLP is an essential cofactor for ACS activity (Mehlta et al., 1993). ACS is encoded by a multigene family and different isoforms are differentially regulated (Barry et al., 2000; Oetiker et al., 1997; Peck and Kende,
Ethylene perception begins with ethylene binding at the receptors located in the endoplasmic reticulum (Nehring and Ecker, 2004). Many ethylene perception receptors were elucidated using the triple response of arabidopsis (Arabidopsis thaliana) seedlings when exposed to ethylene. Bleecker et al. (1988) isolated the first ethylene receptor ETR in arabidopsis. Screening for ethylene-insensitive plants using the triple response led to the isolation of four additional ethylene receptors in arabidopsis: ETR2 (Sakai et al., 1998), ERS1 (Hua et al., 1995), ERS2 (Hua et al., 1998), and EIN4 (Hua et al., 1998; Roman et al., 1995). The N-termini of the receptor proteins contain transmembrane domains, with ETR1 and ERS1 having three transmembrane domains and ETR2, ERS2, and EIN4 having four transmembrane domains (Nehring and Ecker, 2004). The C-termini contain histidine kinase and receiver domains and share similarity with the bacterial two-component signal transduction system (Chang et al., 1993). ERS1 and ETR1 receptors have conserved protein motifs in the histidine kinase domain and are classified as subfamily I receptors, whereas subfamily II receptors (ETR2, ERS2, and EIN4) have one or more protein motifs missing. ERS1 and ERS2 do not have the receiver domains. Downstream of the receptors in the signaling pathway is CTR1 (Kieber et al., 1993). CTR1 interacts with the receptors and together they act as negative regulators in ethylene signaling. Downstream of CTR1 is EIN2, which is a positive regulator in ethylene signaling pathway (Alonso et al., 1999). Downstream of EIN2 is EIN3, which is located in the nucleus. EIN3 belongs to a multigene family designated as EIL proteins. EIN3 and EILs are transcription factors that ultimately activate ethylene-responsive genes. Several studies have examined how ethylene treatment impacted expression of these perception and signaling genes in an attempt to identify which may have important roles in downstream biological responses. In general, subfamily I receptors are thought to play a more dominant role in ethylene signaling (Cancal and Larsen, 2002).

Ethephon is an ethylene-releasing compound used to accelerate abscission. Ethephon is an effective abscission agent in citrus, plum (Prunus domestica), cherry (Prunus cerasus), and olive (Olea europaea) (Bukovac et al., 1969; Burns, 2002; Martin et al., 1981). However, excessive leaf abscission occurs at concentrations required for effective fruit loosening in citrus (Burns, 2002) and olive (Burns et al., 2008). Pozo et al. (2004) used 1-MCP, an ethylene perception inhibitor that irreversibly binds to ethylene receptors, to reduce ethphon-induced leaf drop with minimal effect on ethylene-induced fruit loosening. Application of guanfacine, an agonist of G-protein-coupled \( \alpha_{2A} \)-adrenoreceptors, also reduced ethephon-induced leaf drop but not fruit loosening (Yuan et al., 2005). These data suggest that different abscission control mechanisms may exist in leaf and mature fruit of citrus. Furthermore, ethylene-independent abscission has been observed in several plants. Studies with ethylene-insensitive mutants revealed that ethylene-independent abscission occurs in arabidopsis (Patterson and Bleecker, 2004). Moreover, the lack of ethylene involvement was observed during flower abscission in tulip (Tulipa hybridra) and orchid (Cymbidium sp.) (Sexton et al., 2000; van Doorn, 2002).

For effective use of mechanical harvesting in citrus, it is necessary to use abscission agents that preferentially increase fruit loosening but have very little effect on leaf abscission. Thus, understanding mechanisms of abscission in leaf and mature fruit can provide insight into the abscission process and can assist with selecting an effective abscission compound with uniform efficacy. The objective of this study was to test if differential expression of ethylene biosynthesis and signaling genes were correlated with differential abscission of leaves and mature fruit. In this study, genes involved in ethylene signaling and biosynthetic pathways were cloned from citrus and their expressions were analyzed during abscission in leaf and fruit tissues. We demonstrate that most ethylene biosynthetic and perception gene expression explored in this study were responsive to ethephon application in one or more tissues examined, that failure of 1-MCP to suppress fruit loosening was not correlated with ACS or ACO expression in fruit abscission zones, and that differential timing of subtending organ drop at two spatially distinct leaf abscission zones was not correlated with ethylene sensitivity.

Materials and Methods

Plant materials and treatment. ‘Valencia’ sweet orange trees [17 years old on Swingle citrumelo (Citrus paradisi × Poncirus trifoliate) rootstock] were selected from an experimental grove located at the Citrus Research and Education Center, Lake Alfred, FL. Trees were sprayed until runoff with 450 \( \mu \)L L\(^{-1} \) ethephon or 5 nm of 1-MCP or a combination of both using a motorized back-pack sprayer on 28 Apr. 2006 as described (Pozo et al., 2004). Control trees were sprayed with water and two trees were used for each treatment. The experiment was repeated on 30 Apr. 2008. Data from the two seasons had similar trends and hence the average from both seasons is shown for leaf drop, fruit detachment force, and gene expression. Leaf abscission and FDF were measured immediately after treatment and after 6 h and 1, 2, 4, and 7 d of application. Ten branches (each with about 100 leaves; \( \approx 1 \) year old) per treatment were tagged to follow leaf abscission. Leaf count was taken on these branches and the percentage of leaf drop was calculated. FDF was measured on five mature fruit (percentage soluble solids/percentage acid ratio of juice was \( \approx 15.0 \)) per replication with four replicates using a digital force gauge (Force One; Wagner Instruments, Greenwich, CT).

Reports indicate that abscission may be initiated in organ abscission zones and/or their subtending organs (Alferez et al., 2005; Beyer, 1975), thus leaf blade, laminar abscission zones (LAZ), fruit peel, and fruit abscission zones (FAZ) were used for this study. Samples of leaf blade, LAZ, fruit peel, and FAZ tissues were collected in three replicates during the first season (four replicates during the second season) immediately after treatment and 6 h and 1, 2, 4, and 7 d after application. Leaf blade and LAZ were collected from 10 leaves/replicate at each sampling time. Midsection of the leaf blade was used and LAZ were excised using a razor blade as described (Yuan et al., 2005). Fruit peel and FAZ were collected from four fruit/replicate. Fruit flavedo (the colored portion of the peel) was removed from the equatorial area of each fruit using a potato peeler. FAZ were removed from fruit using a 1-cm-diameter cork borer. The pedicel, calyx, and albedo surrounding the abscission zone were trimmed away using a razor blade, leaving about the 4-mm-thick FAZ intact. Tissues were frozen in liquid nitrogen immediately after sampling and stored at \(-80 \) °C until needed.
RNA extraction. RNA was extracted from 0.5 g of flavedo, 0.5 g of leaf blade, four FAZ (≈0.3 g), and 10 LAZ (≈0.2 g). The frozen tissue was ground into fine powder and suspended in 1 mL of cold TRI Reagent (Molecular Research Center, Cincinnati, OH). Chloroform (200 μL) was added and the mixture vortexed thoroughly. Samples were incubated on ice for 15 min and centrifuged at 20,000 g for 15 min at 4 °C. RNA was extracted from the aqueous fraction using a RNeasy mini kit (Qiagen, Valencia, CA). First-strand cDNA was synthesized from 1 μg of RNA using SuperScript III Reverse Transcriptase (Invitrogen, Carlsbad, CA) in a 20-μL reaction.

Gene cloning. Full-length nucleotide sequences of four putative ethylene receptors were cloned from leaf blade tissues of ‘Valencia’ sweet orange. The full-length sequence of CsERS1 (1905 bp) was obtained from the NCBI database (accession no. AF092088). A partial sequence of CsETR1 was available in NCBI (accession no. AJ276294) and a full-length sequence (2223 bp; accession no. GQ339592) was amplified using the 5’ RACE and 3’ RACE systems (Invitrogen) and cloned. A partial sequence of CsETR2 was available in a citrus mature fruit and leaf EST database (J.K. Burns, unpublished data) and a partial sequence of CsETR3 was obtained from the publically available citrus HarvEST database (University of California, 2009). Full-length sequences of CsETR2 (2295 bp; accession no. GQ339593) and CsETR3 (2292 bp; accession no. GQ339594) were amplified by 5’ RACE and 3’ RACE protocols and cloned. Partial sequences of CsCTR1, CsEIN2, and CsEIL2 were obtained from HarvEST database. A partial sequence of CsEIL1 was obtained from mature fruit and leaf EST database and a partial sequence of CsEIL1 (1845 bp; accession no. GU981740) was obtained from HarvEST database. A partial sequence of CsACO (2292 bp; accession no. GU981740) was cloned. A partial sequence of CsEIL2, CsERS1, CsETR2, and CsEIL2 was cloned. Partial sequences of CsERS1, constitutive triple response 1 (CsCTR1), ethylene response 1 (CsEIN1), ethylene insensitive 2 (CsEIL2), and glyceraldehyde 3-phosphate dehydrogenase (CsGAPDH) have occurred in each replicate. CsGAPDH and the genes of interest had similar efficiency of amplification. Relative gene expression was calculated using the comparative C_T (threshold cycle) method (Livak and Schmittgen, 2001). For each sample, the C_T value for the gene of interest was subtracted from the C_T value of CsGAPDH (ΔC_T). ΔC_T was calculated by subtracting ΔC_T of one T0 control replicate from the ΔC_T of all individual replications for all treatments at each time point. Relative gene expression was calculated using the equation 2^-ΔΔC_T. Dissociation curve generated after the amplification indicated that a single amplicon was produced in each reaction.

Ethylene sensitivity in leaf abscission zones. Citrus leaves have two abscission zones. One abscission zone is located between the leaf blade and the petiole (LAZ) and the second is located between the petiole and the stem (petiolar abscission zone). When abscission agents are applied, leaf blades first abscise due to abscission processes at the LAZ and later followed by abscission of the petioles due to abscission processes at the petiolar abscission zone. To study differences in ethylene sensitivity of LAZ and petiolar abscission zones during ethylene-induced abscission, two 7-year-old potted ‘Valencia’ sweet orange trees on Swingle citrumelo rootstock were placed in a closed chamber and treated with 5 μL·L⁻¹ ethylene for 88 h. The trees were grown in 12/12-h light/dark cycles (light: 0600–1800 hr; dark: 1800–0600 hr) and were watered daily. The temperature and humidity of the chamber were monitored every 15 min using an automatic data logger (HOBO; Onset Computer Corp., Bourne, MA). Leaf blade and petiole drop were counted twice a day at 1000 and 1800 hr on 20 branches (at least 30 leaves/branch).

In another experiment, two 7-year-old potted ‘Valencia’ sweet orange trees on Swingle citrumelo rootstock were placed in a chamber with 5 μL·L⁻¹ ethylene for 24 h and then transferred to ethylene-free air. The plants were held under the same light, temperature, and humidity conditions as above.

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Table 1. Primer sequences, length of amplicon, and primer concentrations used to assess ethylene biosynthesis and perception gene expression for real-time PCR in Citrus sinensis. The genes are 1-amino-cyclopropane-1-carboxylate synthase 1 (CsACS1) and 2 (CsACS2), 1-amino-cyclopropane-1-carboxylate oxidase (CsACO), ethylene response sensor 1 (CsERS1), ethylene response 1 (CsETR1), 2 (CsETR2), and 3 (CsETR3), constitutive triple response 1 (CsCTR1), ethylene insensitive 2 (CsEIL2), ethylene insensitive 3-like 1 (CsEIL1), and 2 (CsEIL2), and glyceraldehyde 3-phosphate dehydrogenase (CsGAPDH).

| Gene       | Forward primer (bp) | Reverse primer (bp) | Final concn of each primer (μM) | Length of amplicon (bp) |
|------------|---------------------|---------------------|---------------------------------|------------------------|
| CsACS1     | 5’TTCGAAATCCACTAGGCACAACCT-3’ | 5’-CAACGCTCTGTAACCTTGGAGA-3’ | 0.5                             | 140                    |
| CsACS2     | 5’-GATGGCGTTATGGCAGTGTA-3’ | 5’-GACAAATTCCATCGTCCGA-3’ | 0.4                             | 134                    |
| CsACO      | 5’-AAGATGGCCCTATTGATTG-3’ | 5’-TCACCGAGGTGACACAAAC-3’ | 0.4                             | 61                     |
| CsERS1     | 5’-TTGTGGGACTGACTGACTTCAAGC-3’ | 5’-ATGACAAAAAAGCAACAAGC-3’ | 0.4                             | 104                    |
| CsETR1     | 5’-TCGTCAAGCAGAATCTCTGTTG-3’ | 5’-GCGTTTAATCTGACTGGACA-3’ | 0.8                             | 120                    |
| CsETR2     | 5’-AACCTTCCCATTCACTAG-3’ | 5’-TCACCGTGACTGAATAAACCTTG-3’ | 0.5                             | 109                    |
| CsETR3     | 5’-GGCCATAATGCAAGGAATATTAAAGAG-3’ | 5’-CGTGAAGCCATCTGATAGTGG-3’ | 0.3                             | 112                    |
| CsCTR1     | 5’-GTGGATGCGACCGAAGTT-3’ | 5’-GAAATTTCGCAAGGGTTTGGCAG-3’ | 0.4                             | 114                    |
| CsEIL2     | 5’-GAAAGGATGATGATGAACTAGGAT-3’ | 5’-GAGCCGGACCATCAGACAT-3’ | 0.2                             | 117                    |
| CsEIL1     | 5’-ACAGAGCAAGGATGGAATGGTGTTG-3’ | 5’-TCTTGGTTGCGGACATCTTC-3’ | 0.4                             | 90                     |
| CsGAPDH    | 5’-GGGCTGAAAGTATGACCAAACT-3’ | 5’-CTAAAGGTTTGTTGTTCTGCTA-3’ | 0.5                             | 75                     |
Leaf blade and petiole drop were counted on six tagged branches (at least 30 leaves/branch) every 12 h for 84 h. Laminar and petiolar detachment force were measured on 30 leaves every 12 h using a digital force gauge (DS2–11; Imada, Northbrook, IL).

**Statistical analysis.** Analysis of variance (ANOVA) was performed using SAS (version 9.1; SAS Institute, Cary, NC). Standard error of the mean was calculated and represented as vertical bars in the figures.

**Results**

**Structure of the cloned citrus ethylene receptors.** Full-length sequences of CsETR1, CsETR2, and CsETR3 (accession nos. GQ339592, GQ339593, and GQ339594, respectively) were obtained from RNA isolated from ‘Valencia’ sweet orange. The predicted protein sequence from the longest cDNA open reading frame for CsERS1, CsETR1, CsETR2, and CsETR3 contained 634, 740, 764, and 763 amino acids with calculated molecular weights of 71, 83, 85, and 86 kDa, respectively. Although CsERS1, CsETR1, and CsETR3 had a predicted pI of 6.5, CsETR2 had a higher pI of 8.0. Based on von Heijne transmembrane prediction (von Heijne, 1992), CsERS1 and CsETR1 contained three transmembrane domains in their N-termini, whereas CsETR2 and CsETR3 had four such domains (Fig. 1). CsERS1 and CsETR1 had histidine kinase domains containing five conserved motifs (H, N, G1, F, and G2). CsETR2 lacked all five conserved motifs in the histidine kinase domain. Only the H motif was present in the histidine kinase domain of CsETR3. CsETR1, CsETR2, and CsETR3 contained the receiver domain with the conserved aspartate residue. The receiver domain was absent in the C-terminus of CsERS1. A structural comparison between the predicted amino acid sequence of the cloned citrus ethylene receptor genes and arabidopsis and tomato heterologs is presented (Fig. 1).

**Differential effect of 1-MCP on leaf and fruit abscission.** Ethephon-induced leaf abscission was 12% after 2 d of application and increased rapidly to 73% after 4 d of application (Fig. 2A). When 1-MCP was applied in combination with ethephon, leaf abscission was delayed and reduced to 6% after 4 d of application and increased to only 22% after 7 d. Visual examination for an additional 2 weeks indicated no further increase in leaf abscission in this treatment combination. Fruit loosening was evident 4 d after ethephon application (Fig. 2B) and FDF was reduced to 6.4 kg of force compared with 10.5 kg of force in untreated control fruit. When 1-MCP was combined with ethephon, fruit loosening was reduced slightly on day 2, and FDF fell to 7.8 kg of force on day 4 after application. However, FDF had fallen to similar levels in both treatments after 7 d of application. 1-MCP treatment alone did not cause leaf abscission or fruit loosening.

**Expression of ethylene biosynthesis genes.** Expression of CsACS1 in peel was ethylene responsive (Fig. 3A). Peak expression occurred 6 h after application and then fell to control levels soon thereafter. A slight but measurable increase in gene expression occurred in FAZ 6 h after ethephon application, but rose 22-fold 24 h after application and then decreased (Fig. 3B). CsACS1 gene expression in peel was suppressed if 1-MCP was combined with ethephon and applied to the canopy; however, expression in FAZ was only delayed. CsACS1 expression in leaf blade and LAZ tissue was ethephon responsive; with peak expression measured 4 d after application (Fig. 3, C and D, respectively). Gene expression was suppressed in leaf tissues when 1-MCP was applied in combination with ethephon. Timing of increased CsACS1 gene expression in leaf tissues coincided with increased leaf drop, while in fruit tissues, increased gene expression preceded fruit loosening. Increased expression in FAZ in response to 1-MCP alone suggests that basal ethylene production may be inhibitory to CsACS1 expression. CsACS2 expression was...
Effect on to control levels after day 4 (Fig. 3F). Ethephon had very little a maximum of 5.7-fold 1 d after application and then falling (not shown).

The production may inhibit CsACO expression in leaf blade (Fig. 4C), ethephon induced maximum and 1 and 2 d after treatment and dropped to control levels respectively.

CsERS1 was ethylene responsive in all four tissues examined (data not shown). Expressed in FAZ reached a maximum (6.8-fold) 6 h after application of ethephon and remained high through day 4, after which expression decreased (Fig. 3E). Treatments with 1-MCP alone or in combination with ethephon delayed the rise in CsACO expression. Like CsACS1 expression, basal ethylene production may inhibit CsACO expression in FAZ. In LAZ, ethephon treatment increased CsACO expression, reaching a maximum of 5.7-fold 1 d after application and then falling to control levels after day 4 (Fig. 3F). Ethephon had very little effect on CsACO expression in leaf blade and fruit peel (data not shown).

Expression of ethylene receptor and signaling genes. CsERS1 was ethylene responsive in all four tissues examined (Fig. 4). In fruit peel (Fig. 4A) and FAZ (Fig. 4B), expression increased to a maximum 6 h after application and gradually declined to control levels by 4 and 2 d after application, respectively. CsERS1 expression in response to ethephon increased gradually in leaf blade (Fig. 4C) and LAZ (Fig. 4D) to a maximum of 2.1- and 2.6-fold after 2 and 1 d of application, respectively. CsETRI expression was induced by ethephon only in fruit peel where the maximum increase in expression (2.2-fold) occurred after 1 d of application and decreased to control levels on day 4 (data not shown). Similar to CsERS1, expression of CsETR2 was ethylene responsive in all four tissues (Fig. 5). Ethephon-induced expression in fruit peel, FAZ, and LAZ (Fig. 5 A, B, and D) was higher than control after 6 h and 1 and 2 d after treatment and dropped to control levels by day 4. In leaf blade (Fig. 5C), ethephon induced maximum increase in CsETR2 expression after 1 d of application and levels in leaf blade remained higher than the control until the end of the experiment. Expression of CsETR3 was unaffected by ethephon (data not shown). Ethephon-induced increases of receptor gene expression were counteracted by 1-MCP. Expression of CsERS1 and CsETR2 was generally lower than control when 1-MCP was applied alone or in combination with ethephon, while expression of CsETR1 and CsETR3 genes was similar to the control. The expression of CsCTR1, CsEIN2, CsEIL1, and CsEIL2 was variable but not significantly altered by ethephon and/or 1-MCP treatment in any of the tissues (data not shown).

Ethylene sensitivity in leaf abscission zones. When potted trees were treated with continuous ethylene exposure, abscission of leaf blades was observed 24 h after exposure, whereas abscission of petioles began after 40 h (Fig. 6A). Abscission of leaf blades increased rapidly to 95% after 40 h of continuous ethylene exposure, and after 64 h, all leaf blades
Fig. 6. Percentage of abscission of leaf blade (A) and petiole (B) in ‘Valencia’ sweet orange when treated with 5 μL·L⁻¹ ethylene for 88 h followed by transfer to ethylene-free air (B). Detachment force (C) of leaf blade and petiole was measured on organs from potted trees treated as in (B). Time of watering is indicated by t, and white and black bars indicate light and dark conditions, respectively. Vertical bars through symbols represent se. Where bars are not visible, symbols are larger than se.

Absced. Abscission of bladeless petioles steadily increased to 93% by 72 h and reached 100% by 88 h. When potted trees were treated with ethylene for 24 h and then transferred to ethylene-free air, leaf blade abscission began 12 h after ethylene removal, whereas petioles started to abscise 36 h after ethylene removal (Fig. 6B). After 60 h of transfer from ethylene, 57% and 13% abscission of leaf blades and petioles was observed, respectively. Leaf blade and petiole detachment forces, however, indicated that reduction in detachment force due to ethylene treatment was similar at both abscission zones (Fig. 6C). The detachment force in both abscission zones reduced from about 1 kg of force to 0.6 kg of force at 12 h and reached a minimum of 0.1 kg of force by 24 h.

Discussion

Ethylene receptors are classified into two groups based on conserved motifs present in the histidine kinase domain. Subfamily I receptors contain all conserved motifs in the histidine kinase domain, while subfamily II receptors lack one or more of the conserved motifs. Based on these criteria, CsERS1 and CsETR1 are classified as subfamily I receptors, while CsETR2 and CsETR3 are classified as subfamily II receptors. In arabidopsis, subfamily I receptors play more important roles than subfamily II receptors. Wang et al. (2003) demonstrated that loss of function in the ers1;etr1 double mutant was not compensated by overexpression of any of the three subfamily II receptors. Furthermore, physical interaction between subfamily I receptors and CTR1 was stronger than that of ETR2 (subfamily II) receptor and CTR1, suggesting an important role of subfamily I receptors in ethylene signaling (Cancel and Larsen, 2002). Because an inverse relationship exists between receptor abundance and ethylene sensitivity (Klee, 2004), differences in the abundance of CsERS1 and/or CsETR1 between leaf and fruit tissues could explain differential ethylene sensitivity in these tissues.

CsERS1 is structurally similar to AtERS1 and LeETR3, having three transmembrane domains, all conserved motifs in the histidine kinase domain, and lacking the receiver domain (Fig. 1; Hua et al., 1998; Klee and Tieman, 2002). CsETR1 has three transmembrane domains, a conserved histidine kinase domain, and a receiver domain, which is similar to AtETR1, LeETR1, and LeETR2. CsETR2 and CsETR3 have four transmembrane domains and the receiver domain. CsETR2 lacks all the conserved motifs in the histidine kinase domain, while CsETR3 contains only the H motif. Similar to CsETR2, all the motifs in the histidine kinase domain are absent in AtERS2 and LeETR5, but unlike CsETR2, the receiver domain is absent in AtERS2. CsETR3 is similar to AtEIN4 with four transmembrane domains, only the H motif in histidine kinase domain, and a receiver domain. Although receptors are redundant, they could have unique functions because they exhibit differential expression patterns in various tissues (Hua et al., 1998; Lashbrook et al., 1998; Tieman and Klee, 1999). Nf (LeETR3) expression increased 10- to 20-fold during tomato fruit ripening, and reproductive tissues, flowers, and young developing fruit had higher expression of LeETR4, LeETR5, and LeETR6 when compared with vegetative tissues. CsERS1 and CsETR2 are structurally similar to LeETR3 and LeETR5, respectively, and may play a greater role in ethylene signaling in citrus fruit than in leaves. Furthermore, differences in the levels of different receptors in vegetative and reproductive tissues could result in differential ethylene sensitivity in citrus fruit and leaves.

Citrus leaves and fruit abscised when treated with ethephon, but foliage appeared to respond more readily by dropping over 80% of the total leaves after 7 d. Apart from this differential intensity of response to ethephon, differential response to 1-MCP was also observed. Ethephon-induced leaf drop was suppressed by application of 1-MCP in combination with ethephon, while it had little effect on fruit loosening. Similar results were observed with application of 1-MCP (Pozo et al., 2004) or guanfacine, an agonist of G-protein-coupled α₂A*-adrenoreceptors (Yuan et al., 2005). Because 1-MCP is a gas, penetration of 1-MCP into bulky fruit tissues may have been restricted, but less so in leaf tissues. Absorption of 1-MCP was reported to vary between different plant tissues (Nanthachai et al., 2007). These differences were due to various cellular components such as oils, polysaccharides, and lignin that alter 1-MCP absorption. 1-MCP was absorbed faster and in greater amounts in high-lipid avocado (Persea americana) fruit than apple (Malus domestica) of lower lipid content (Dauny et al., 2003). Choi and Huber (2009) reported that 1-MCP absorption ranged from 34% to 94% in fruit and vegetable tissues, depending on the type of polysaccharide, with xyloglucan exhibiting the lowest absorption (34%) followed by cellulose (38%), starch (49%), and esterified pectin (94%). Among the aliphatic components of lipid-derived polymers, citrus leaves contain a higher percentage of fatty alcohols and fatty acids than citrus fruit peel, while peel has a higher percentage of ω-hydroxy acids, dicarboxylic acids, and polar acids than leaves (Espelie et al., 1980). Pectin content was slightly higher in citrus fruit flavedo (270 mg·g⁻¹ dry weight) than in leaf blade (220 mg·g⁻¹ dry weight; H.-L. Liao and J.K. Burns, unpublished...
data). In whole fruit, pectin is expected to be much higher than
leaves due to the highly pectinaceous albedo. Nonetheless, our
gene expression data indicated that 1-MCP penetrated flavedo
and fruit abscission zone tissues sufficiently because ethephon-
induced receptor gene expression was suppressed. Thus,
differences in oil, poly saccharide, lignin, and pectin content
cannot solely account for differential response observed in fruit
and leaves.

Alternatively, regeneration of new receptor sites could play
a role in the differential abscission response. When ethylene
and 1-MCP are coapplied to tissues, 1-MCP preferentially
binds to the receptors, but there is no 1-MCP binding preference
between receptor types (Hall et al., 2000). Because 1-MCP was
applied as a gas to citrus canopies in an open environment, it
would be available only for a short duration, while release of
ethylene into plant tissues from a liquid ethephon solution could
take place after several hours of application (Domir and Foy,
1978; Perry and Gianfagna, 1987). Though more than 50% of
ethephon was dissociated to ethylene within 24 h of application,
ethylene release continued to occur for 96 h in leaves of tobacco
[Nicotiana tabacum (Domir and Foy, 1978)] and peach (Perry
and Gianfagna, 1987). Several studies indicated that new
receptors were synthesized after 1-MCP was bound to available
receptors (Binder et al., 2004; Blankenship and Dole, 2003). In
avocado, two applications of 1-MCP at 10-d intervals were
required to prevent fruit softening (Pesis et al., 2002), and
1-MCP was effective in increasing vase life of hibiscus
(Hibiscus rosa-sinensis) flowers only by continuous exposure
for 15 h (Reid et al., 2002). Application of 1-MCP delayed
ripening of avocado and tomato by 2 weeks and 5 to 10 d,
respectively (Feng et al., 2000; Wills and Ku, 2002). If new
receptors were synthesized and incorporated into the endoplasmic
reticulum, the rate of new receptor regeneration must have
varied between tissues. The amount or rate of receptor re-
generation in mature citrus fruit could be much lower than in
leaves, creating a situation of lower new and functioning receptor
numbers in fruit. Because low receptor number is associated with
greater ethylene sensitivity (Hua and Meyerowitz, 1998; Klee,
2001). Though the mechanism is not clear in CTR1 and EIN2,
post-transcriptional regulation may play a role in citrus as postulated in tomato and
arabidopsis (Gao et al., 2003; Guo and Ecker, 2003; Kieber
et al., 1993; Klee, 2004; Leclercq et al., 2002; Tieman et al.,
2001). Though the mechanism is not clear in CTR1 and EIN2,
post-transcriptional regulation of EIN3 and EIL protein was
described by Guo and Ecker (2003). Degradation of EIN3
protein occurs through an ubiquitin/proteasome pathway me-
diated by two F box proteins, EBF1 (EIN3-binding F box
protein 1) and EBF2. This pathway is inactive in the presence of
ethylene, resulting in increased accumulation of EIN3 protein.
Analyzing the protein expression of CTR1, EIN2, and EILs
could help to explain differential abscission responses in citrus.

Taken together, leaves and fruit respond differentially
to ethephon and/or 1-MCP. Although differences were seen in
timing and intensity of gene expression, the most significant
differences were measured in ethylene biosynthesis gene
expression in FAZ and LAZ. During the course of this study, it was
shown that the timing of ethephon-induced abscission associated
with the two abscission zones located in leaves was different.
Soon after ethephon application in citrus, the leaf blade abscised
first, followed days later by the subtending petiole. However,
differences in timing of abscission of the blade and petiole could
not be attributed to differences in ethylene sensitivity. Although
abscission occurred earlier in the LAZ than the petiolar
abscission zone when treated with ethylene, the reduction in
detachment force at both locations was similar. Because force is
directly proportional to mass (force = mass \times acceleration) and
the average weights of leaf blade and petiole are 579 and 36 mg, respectively, much more downward force was exerted by the leaf blade than by the petiole. The total force exerted by the leaf blade may not be transferred to the petiolar abscission zone due to the presence of the weakened LAZ. This might have caused earlier abscission of the leaf blade at the LAZ than at the petiolar abscission zone. Further work on quantifying the number of receptors in leaf and fruit tissues and the regeneration capacity of receptors would more fully elucidate the basis of this differential abscission response.

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