Oxidative Metabolism in ‘Valencia’ Sweet Orange (Citrus sinensis Osbeck) Abscission Zone Tissue Treated with the Abscission Agent 5-Chloro-3-Methyl-4-Nitro-1H-Pyrazole

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Abstract. 5-Chloro-3-methyl-4-nitro-1H-pyrazole (CMNP) is an abscission agent, standardized for the mechanical harvesting of late season ‘Valencia’ sweet oranges in Florida. This work was conducted to investigate the role of CMNP to induce oxidative stress in the abscission zone (AZ) of ‘Valencia’ sweet orange. Fully mature ‘Valencia’ sweet orange trees in a commercial grove were sprayed with 2.0 mM of CMNP. The experiment was repeated three times during the Apr.–May 2013 harvest season. Fruit material and culture. Three independent experiments were conducted on fully mature 15-year-old trees of ‘Valencia’ sweet orange (Citrus sinensis (L.) Osbeck) and to understand oxidative association between FT and AZ to mediate CMNP-induced abscission cascade in citrus.

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1.9-fold higher in CMNP-treated fruit at 3DAT. Higher GR and low APOn activity reflects limited functioning of the APOD/GR cycle (e.g., APOD and GR) in scavenging of H2O2 at 3DAT. Guaiacol POD activity transiently increased at 1DAT and then declined. POD plays an important role in cell wall lignification and indole acetic acid (IAA) oxidation. The decline in POD activity may cause a decrease in lignification while higher activity made the AZ sensitive to ethylene and thus promote abscission in citrus fruit. This work also showed that CMNP-induced abscission is a collaborative effort of oxidative metabolism in flavedo tissue (FT) and AZ.

Abscission is an active process by which plants shed vegetative and reproductive parts during various developmental stages of their life cycle though a narrow zone of anatomically distinct cells that constitute AZ (Estornell et al., 2013). Besides developmental signals and environmental stresses, commercially available compounds also cause abscission in plant organs (Kossuth et al., 1978; Taylor and Whitelaw, 2001). The peel of citrus fruit consists of the colored outer flavedo and the inner white spongy albedo. However, calyx AZ is situated deeper within the albedo tissue and could be visually identified (Brown and Burns, 1998). CMNP is a pyrazole compound and demonstrated as a potential nontoxic abscission agent for the late season ‘Valencia’ sweet orange crop (Li et al., 2008). It has also been shown to aid in mechanical harvesting of citrus in Florida (Yuan and Burns, 2004). Several studies have been conducted to understand the mechanism of CMNP-induced loosening in mature citrus fruit (Kossuth et al., 1978; Li et al., 2008; Yuan and Burns, 2004). Mechanisms of abscission vary among particular plant parts (Lewis et al., 2006). These studies were focused on CMNP-induced metabolic changes in FT of citrus fruit instead of calyx AZ (Alferez et al., 2005; Alferez et al., 2006; Alferez et al., 2007; Kumar and Ebel, 2015). It has been shown that following CMNP application, there is a gradual decline in chlorophyll content with concomitant rise in total carotenoid in FT of citrus (Alferez et al., 2005). These authors also showed an increase in electrolyte leakage upon CMNP treatment, which suggests membrane degradation could be the possible mechanism for CMNP-induced abscission in citrus (Alferez et al., 2005). It has been proposed that CMNP-induced lipid signaling in FT promotes abscission in calyx AZ (Alferez et al., 2005). CMNP-mediated higher levels of phospholipase A2 (PLA2) and LOX activities were observed in FT (Alferez et al., 2005). However, inhibitory studies using aristolochic acid, which is a potent inhibitor of PLA2 and LOX revealed that lipid signaling partly (≥70%) accounts for CMNP-induced abscission (Alferez et al., 2005). This evidence suggests that in addition to lipid signaling, other unknown mechanisms also contribute to CMNP-mediated abscission. We showed that other signaling molecules are reactive oxygen species (ROS), such as O2·-, H2O2, and hydroxyl radicals (OH·), in CMNP-mediated abscission cascade (Kumar and Ebel, 2015; Sakamoto et al., 2008). Our earlier work on FT showed the generation of H2O2 at 1DAT, which is a mobile signaling molecule and has been shown to induce abscission (Kumar and Ebel, 2015; Sakamoto et al., 2008).

Several other pyrazole compounds [pyrazole pyrimidines, pyrazole hydrazine (anticancerous), methyl pyrazole (pesticide), and pyrazolo triazine (herbicides)] also share a common mode of action in treated cells by generating ROS (Graillot et al., 2012; Hassan et al., 2011; Vicentini et al., 2004). There are reports that showed a direct relationship between ROS and the abscission process (Djanaguiraman et al., 2004; Sakamoto et al., 2008). Sakamoto et al. (2008) observed that H2O2 is continuously produced by the cells of AZ in vitro following ethylene exposure. In addition, ethephon-induced abscission was suppressed by inhibitors of H2O2 in Capsicum baccatum plants (Sakamoto et al., 2008). However, higher activity of antioxidant enzymes has been shown to delay abscission in tomato (Lycopersicon esculentum Mill.) fruit (Djanaguiraman et al., 2004).

The objective of this study was to explore the role of CMNP-induced oxidative metabolism in AZ of ‘Valencia’ sweet orange (Citrus sinensis (L.) Osbeck) and to understand oxidative association between FT and AZ to further dissect the overall mechanism of abscission. This work provided preliminary data on CMNP-induced H2O2 changes in AZ of ‘Valencia’ sweet orange and also suggests the existence of oxidative cooperation between FT and AZ to mediate CMNP-induced abscission cascade in citrus.

Materials and Methods

Plant material and culture. Three independent experiments were conducted on fully mature 15-year-old trees of ‘Valencia’ sweet orange (C. sinensis Osbeck) scions on ‘Carri- zizo’ citrone rootstocks during the Apr.–May 2013 harvest season. The studies were conducted in a commercial grove (Lykes Brothers Inc., Basinger Grove, Lake Placid, FL) that applied standard cultural practices (Li et al., 2008). The trees were spaced 3.7 m × 7.6 m and were 4.7 m in height and skirted (height of the lowest branches from soil line) to 0.4 m.
Treatments. Trees were sprayed using a multifeed air-blast sprayer (Model T1000; OXBO International, Clear Lake, WI) with a vertical (5.5 m) boom oriented parallel to andarched over the outer part of the canopy (Ebel et al., 2010). CMNP was applied at a rate of 2.0 mL with 0.55 mg L⁻¹ of Activator 90 as an adjuvant (alkylphenol ethoxylate, alcohol ethoxylate, and tall oil fatty acid or liquid rosin; Loveland Products Inc., Greeley, CO). The CMNP applications were made based on air temperatures being >15.6 °C with no rain forecast for the first 24 h after application (Kossuth et al., 1978). Five trees were used for each treatment or as untreated controls at 0, 1, 2, and 3 DAT for estimation of H₂O₂ and (Kossuth et al., 1978). Five trees were used for forecast for the first 24 h after application in the cylindrical tissue and further trimmed to through the pedicel and pushed up to the peel collected from 40 fruits by using a 4-mm-

APOD, and GR activities.

Sampling AZ tissue. The calyx AZ was collected from 40 fruits by using a 4-mm-diameter cork borer. The cork borer was passed through the pedicel and pushed up to the peel (Burns, 2002). The AZ was visually identified in the cylindrical tissue and further trimmed to 6 mm x 4 mm (Brown and Burns, 1998). These samples were collected in the field and stored in liquid nitrogen at ~80 °C till further studies.

Measurement of the H₂O₂ generation. Each AZ sample [0.5 g fresh weight (FW)] was homogenized in 2 mL of 0.1% trichloroacetic acid (TCA) and centrifuged at 12,000 g, for 15 min at 4 °C. The supernatant (0.3 mL) was mixed with 1.7 mL of 1.0 M potassium phosphate buffer, pH 7.0, and 1.0 mL of 1.0 M potassium iodide solution, and then incubated for 5 min at room temperature before measuring absorbance at 390 nm. H₂O₂ concentrations were calculated using a standard curve prepared from known concentrations of H₂O₂ and were expressed in μmol·g⁻¹ FW (Velikova et al., 2000).

Measurement of lipid peroxidation. Lipid peroxidation was measured in terms of MDA concentrations using the thiobarbituric acid assay described by Heath and Packer (1968) with slight modifications. Each AZ sample (0.5 g FW) was homogenized in 2 mL of 0.1% TCA (w/v) and centrifuged at 12,000 g, for 30 min. The supernatant 1 mL was in-
cubated with 2 mL of 20% TCA (w/v) containing 0.5% (w/v) thiobarbituric acid at 1.9 (v/v) for 30 min at 95 °C. The reaction was stopped by cooling the test tubes in an ice bath for 10 min and the contents were then centrifuged at 10,000 g, for 15 min. The absorbance of the supernatant was measured at 532 nm. The value for nonspecific absorption at 600 nm was sub-
tracted and the amount of MDA-thiobarbituric acid complex was calculated from an extinction coefficient of 155 m M·cm⁻¹. The MDA con-
centration expressed in μmol·g⁻¹ FW.

Enzyme assays. AZ samples (0.5 g FW) were homogenized in 3 mL of 100 mM potassium phosphate buffer, pH 7.0, containing 0.5 M ethylenediaminetetraacetic acid (EDTA), 0.1 mM phenylmethylsulfonyl fluo-
ride, 2% (w/v) polyvinylpyrrolidone, and 5 mM ascorbate, using a prechilled mortar and pestle. The homogenate was centrifuged at 4 °C for 30 min at 15,000 g. For all enzymatic analyses, the supernatant extract was passed through Sephadex G-25, PD-10 column (Pharmacia, GE Health Care Bio-
Science AB, Uppsala, Sweden) equilibrated with 50 mM potassium phosphate buffer, pH 7.0. All enzymes activities were measured in 3 mL reaction volumes using different ali-
quets of each supernatant [25 μL (5 μg protein) for SOD, 25 μL (5 μg protein) for LOX, 50 μL (10 μg protein) for POD, 50 μL (10 μg protein) APOD, and 100 μL (20 μg protein) for GR].

SOD activity was determined by the method of Dhindsa et al. (1981). Each SOD assay reaction (3.0 mL) contained 0.2 M methionine, 1.72 mM nitroblue tetrazolium, 400 μM potassium phosphate buffer, and an appropriate aliquot (25 μL) of the gel-
filtered AZ extract, and 0.12 mM riboflavin in 50 mM potassium phosphate buffer, pH 7.8. The riboflavin was added last. The contents of each tube were thoroughly mixed and placed 30 cm below the two 75-W fluorescent bulbs (Philips, Andover, MA). The reaction was started by switching on the light for 10 min. Each tube was then covered with a black cloth immediately after switching off the light. Nonirradiated reaction mixtures containing the same AZ extract, which did not develop color, were used as controls. A blank reaction without any added AZ extract produced the maximum blue color. The absorbance of each reaction mixture was read at 560 nm. One unit of SOD activity was defined as the amount of enzyme that caused 50% inhibition of the initial rate of the reaction in the absence of enzyme (Dhindsa et al., 1981). The specific activity of SOD was expressed in units min⁻¹·mg⁻¹ total soluble protein (TSP; Bradford, 1976).

APOD activity was assayed in 3 mL re-
action mixtures containing each containing 50 mM potassium phosphate buffer, pH 7.0, 0.1 mM EDTA, 1.0 mM H₂O₂, 0.25 mM ascorbic acid, and 50 μL of AZ extract. APOD activity was measured by the rate of oxidation of ascorbate at 290 nm using extinction coefficient of 2.8 mM⁻¹·cm⁻¹ (Nakano and Asada, 1981). The specific activity of APOD was expressed in μmol ascorbic oxidized min⁻¹·mg⁻¹ TSP.

GR activity was measured in 3 mL re-
anction mixtures containing 50 mM potassium phosphate buffer, pH 7.5, 3 mM 5,5-dithio-
-bis-2-nitrobenzoic acid, 0.1 mM EDTA, 2 mM nicotinamide adenine dinucleotide phosphate, and 100 μL of AZ extract. Each reaction was started by adding 0.67 mM oxidized glutathii-
one. The increase in absorbance at 412 nm at 25 °C was recorded using a period of 5 min (Smith et al., 1988). The specific activity of GR was expressed in ΔA₄₁₂ min⁻¹·mg⁻¹ TSP.

LOX activity. LOX specific activity increased at 1 DAT and then transiently de-
clined at 2 DAT in AZ of treated fruit and thereafter increased (Fig. 2A). Highest LOX specific activity (1.33 ΔA₄₃₄ min⁻¹·mg⁻¹ TSP) was observed at 1 DAT.

APOD activity. The changes in APOD specific activity were nonsignificant up to 2 DAT in AZ of CMNP-treated and control fruit (Fig. 2B). Significant reduction (50%) in APOD activity was observed at 3 DAT in

Results

H₂O₂ concentrations. CMNP treatment increased the formation of H₂O₂ in AZ of fruit at 3 DAT (Fig. 1A). The rise in H₂O₂ production was 1.3-fold in comparison with controls at 3 DAT. However, H₂O₂ concentra-
tion remained constant (≈1.8 μmol·g⁻¹ FW) in AZ of control fruit throughout the sampling period.

Lipid peroxidation. The concentrations of MDA were similar in AZ for both of CMNP-
treated and control fruit throughout the abscis-
sion process (Fig. 1B). The levels of MDA were ≈24 mmol·g⁻¹ FW in AZ of both control and treated fruit.

SOD activity. SOD specific activity declined severely (65.0%) to 2.97 units min⁻¹·mg⁻¹ pro-
tein at 3 DAT from 4.47 units min⁻¹·mg⁻¹ TSP at 0 DAT in AZ of treated fruit (1C). The decline was 75.0% at 2 DAT. Total SOD activity remained constant (≈4.5 units min⁻¹·mg⁻¹ TSP) in AZ of control fruit.

LOX activity. LOX specific activity in-
crease at 1 DAT and then transiently de-
clined at 2 DAT in AZ of treated fruit and thereafter increased (Fig. 2A). Highest LOX specific activity (1.33 ΔA₄₃₄ min⁻¹·mg⁻¹ TSP) was observed at 1 DAT.

APOD activity. The changes in APOD specific activity were nonsignificant up to 2 DAT in AZ of CMNP-treated and control fruit (Fig. 2B). Substantial reduction (50%) in APOD activity was observed at 3 DAT in

Statistical analysis. Three separate exper-
iments were arranged in a randomized com-
plete block design and five trees were used for each treatment. Each assay was repeated five times on AZ from 45 fruit of each plant. Means values for fruit from each plant were determined and subjected to one-way analy-
sis of variance and separated using the protected least significant difference test using SAS (SAS Institute, Cary, NC) at P ≤ 0.05. se of the mean were also calculated.
AZ of treated fruit. APOD activity in AZ of control fruit was \(\approx 0.23 \text{ \mu mol \ asc. min}^{-1} \text{mg}^{-1} \text{TSP} \) during the abscission period.

**GR activity.** CMNP enhanced the GR specific activity at successive stages of abscission process (Fig. 2C). The highest GR activity (0.93 \(\Delta A_{412} \text{ min}^{-1} \text{mg}^{-1} \text{TSP} \)) was observed at 3 DAT. The increase in GR activity was 1.3-fold and 1.6-fold at 1 and 2 DAT, respectively. GR activity in AZ of control fruit remained constant (\(\approx 0.50 \Delta A_{412} \text{ min}^{-1} \text{mg}^{-1} \text{TSP} \)) at all the stages.

**POD activity.** POD specific activity increased transiently at 1 DAT in AZ of treated fruit and thereafter declined (Fig. 2D). The rise in POD activity was 1.3-fold at 1 DAT. The lowest POD activity was 0.66 \(\text{\mu mol H}_2\text{O}_2 \text{ min}^{-1} \text{mg}^{-1} \text{TSP} \) at 3 DAT.

**Discussion**

Several mechanisms have been proposed to elucidate the role of CMNP in citrus abscission. However, most of the investigations were restricted to the FT of fruit (Alferez et al., 2005, 2006, 2007; Kumar and Ebel, 2015). It has been reported that following CMNP treatment, FT generates an unknown lipid signal that facilitates abscission in citrus (Alferez et al., 2005).

Our earlier work showed that in addition to lipid signals, CMNP also causes formation of ROS in FT (Kumar and Ebel, 2015). Herein, we further confirmed the occurrence of oxidative events in AZ tissue of ‘Valencia’ sweet orange following CMNP application. Progressive decline in fruit detachment force (FDF) was observed following CMNP treatment, which showed the occurrence of cell wall loosening in AZ tissue (Kumar and Ebel, 2015). The decline in FDF was not associated with \(\text{H}_2\text{O}_2 \) accumulation in AZ tissue up to 2 DAT (Fig. 1A). However, it could be mediated by higher levels of \(\text{H}_2\text{O}_2 \) (14–16 \(\text{\mu mol g}^{-1} \text{FW} \)) in FT (Kumar and Ebel, 2015). We proposed that CMNP induce a cooperative action between FT and AZ in terms of \(\text{H}_2\text{O}_2 \) generation and SOD activity. Lower accumulation (1.8–2.0 \(\mu \text{mol g}^{-1} \text{FW} \)) of \(\text{H}_2\text{O}_2 \) between 0 and 2 DAT in AZ was associated with decline (74% at 2 DAT and 65% at 3 DAT) in total SOD specific activity at these stages of abscission (Fig. 1C). However, this decline seems to be complemented by higher SOD specific activity (\(\approx 10 \text{ units min}^{-1} \text{mg}^{-1} \text{TSP} \) at 1–3 DAT) in FT (Kumar and Ebel, 2015). Higher SOD activity produced a copious amount of \(\text{H}_2\text{O}_2 \) (1 DAT: 16 \(\mu \text{mol g}^{-1} \text{FW} \); 2 DAT: 14 \(\mu \text{mol g}^{-1} \text{FW} \); 3 DAT: 13 \(\mu \text{mol g}^{-1} \text{FW} \)) in FT (Kumar and Ebel, 2015). \(\text{H}_2\text{O}_2 \) is a mobile signal and can easily diffuse through membranes and maintained up to 16–18 \(\mu \text{mol g}^{-1} \text{FW} \) of \(\text{H}_2\text{O}_2 \) (FT + AZ) in AZ (Kumar and Ebel, 2015; Makino et al., 2004). Alternatively, lower SOD activity in AZ could increase the production of \(\text{O}_2^- \). Free radical-mediated cell wall loosening has been observed during abscission, seed germination, cell elongation, and fruit ripening (Cohen et al., 2015; Fry et al., 2001). In addition, there is a possibility of generation of highly reactive...
OH in AZ through the action of PODs (Cohen et al., 2014). PODs are heme-containing enzymes and can generate OH (Cohen et al., 2014). Higher POD activity was detected in AZ tissue at 1 DAT (Fig. 2D) that could generate OH via the Fenton reaction. FT derived H₂O₂ (16 μmol·g⁻¹ FW) might be the substrate for this reaction at 1 DAT (Kumar and Ebel, 2015). The Fenton reaction uses H₂O₂ to oxidize the metal ions cuprous (Cu⁺) and ferrous (Fe²⁺), and thus generate OH in biological systems (Cohen et al., 2014). Hydroxyl radical can directly break or weaken the polysaccharide chains in AZ of longan (Dimocarpus longan Lour.) fruit during carbohydrate starvation (Yang et al., 2015).

CMNP-induced oxidative stress involves the following sequence of characteristic events during abscission: 1) higher levels of H₂O₂ in FT can mediate OH formation in AZ, 2) higher activity of POD at 1 DAT in AZ tissue can also generate OH, 3) suppression of SOD in AZ can generate O⁻, and 4) H₂O₂ generation at 3 DAT in AZ can further enhance formation of OH. All these findings suggest the role of ROS in CMNP-induced abscission in citrus.

The levels of MDA concentration remained constant (22–24 nmol·g⁻¹ FW; Fig. 1C) in AZ in comparison with FT (20–40 nmol·g⁻¹ FW; Kumar and Ebel, 2015). However, specific LOX activity increased at 2 DAT in AZ that might generate lipid signals or precursor for JA biosynthesis (Fig. 2A; He et al., 2002). It has been well established that early steps in JA biosynthesis are catalyzed by the products of LOX1, LOX2, LOX3, and LOX4 genes in Arabidopsis (He et al., 2002). The expression of these genes was several-fold higher during leaf senescence in Arabidopsis thaliana (He et al., 2002). Hartmond et al. (2000) reported that exogenous application of methyl jasmonate (MJ; 10 mM) causes reduction in FDF (<40 N) in ‘Hamlin’ and ‘Valencia’ sweet oranges (C. sinensis Osbeck). MJ application also accelerates production of internal ethylene in these genotypes (Hartmond et al., 2000). However, CMNP application induced 10 times higher production of ethylene in comparison with MJ (U. Hartmond, unpublished data). We suggest that LOX-mediated biosynthesis of JA and subsequent production of ethylene may be a part of CMNP-induced abscission program in AZ tissue.

The pattern of CMNP-induced changes in specific activity of APOD and GR are tissue specific and differ in FT (Kumar and Ebel, 2015) and AZ (Fig. 2B and C). APOD and GR activities declined throughout the abscission in FT but in AZ, APOD activity remained constant up to 2 DAT and then declined by 50%. GR activity profile in AZ responded opposite to FT and showed a rise in specific activity at 1 DAT (26%), which further increased up to 3 DAT (82%). APOD-GR cycle regulates cellular redox in plant cells and protects the cell and selected cell organelles from the deleterious effects of H₂O₂ (Perl-Treves and Perl, 2002). Low APOD and high GR activity in AZ showed an impaired functioning of APOD-GR cycle, which is responsible for...
shown to elevate the levels of ethylene in as IAA oxidases and can cause local decline in in FT and AZ to lower the total concentration of IAA oxidase (Kumar and Ebel, 2015). This observation showed a collaborative action of POD in FT and AZ to lower levels of APOD-GR cycle in FT and AZ further maintained oxidative reduct (Kumar and Ebel, 2015), and 3) complementary higher POD activity in FT and AZ leads to lower levels of IAA (Kumar and Ebel, 2015).

**Conclusions**

In summary, CMNP induce a plethora of oxidative events in AZ tissue as described in Fig. 3. We suggest that further experimentation is required to dissect the role of individual ROS in CMNP-induced abscission process in citrus.

**Literature Cited**

Alferez, F., L. Pozo, and J.K. Burns. 2006. Physiological changes associated with senescence and abscission in mature citrus fruit induced by 5-chloro-3-methyl-4-nitro-1H-pyrazole and ethylene application. Physiol. Plant. 127:66–73.

Alferez, F., S. Singh, A.L. Umbach, B. Hockema, and J.K. Burns. 2005. Citrus abscission and Arabidopsis plant decline in response to 5-chloro-3-methyl-4-nitro-1H-pyrazole are mediated by lipid signaling. Plant Cell Environ. 28:1436–1449.

Alferez, F., G.Y. Zhong, and J.K. Burns. 2007. A citrus abscission model addresses anoxia- and senescence-related gene expression in Arabidopsis. J. Exp. Bot. 58:2451–2462.

Almagro, L., L.V.G. Ros, S. Belchi-Navarro, R. Bru, A.R. Barceló, and M.A. Pedreno. 2009. Class III peroxidases in plant defense reactions. J. Exp. Bot. 60:377–390.

Asada, T. 1999. The water-water cycle in chloroplast: Scavenging active oxygen species and dissipation of excess photons. Annu. Rev. Plant Biol. 50:601–639.

Bradford, M.M. 1976. A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Anal. Biochem. 72:248–254.

Brown, G.E. and J.K. Burns. 1998. Enhanced activity of abscission enzymes predisposes oranges to invasion by Diplodia natalensis during ethylene degreening. Postharvest Biol. Technol. 14:217–227.

Burns, J.K. 2002. Using molecular biology tools to identify abscission materials for citrus. HortSci. 37:459–464.

Calmkam, I. and M. Marschner. 1992. Magnesium deficiency and high light intensity enhance activities of superoxide dismutase, ascorbate peroxidase, and glutathione reductase in bean leaves. Plant Physiol. 98:1222–1227.

Cohen, M.F., S. Gurung, G. Birada, H.-Y.N. Helman, and H. Yamazaki. 2015. Bimodal effect of hydrogen peroxide and oxidative events in nitrite-induced rapid root abscission by the water fern Azolla pinnata. Front. Plant Sci. 6(518):1–8.

Cohen, M.F., S. Gurung, J.M. Fukuto, and H. Yamazaki. 2014. Mediated free radical attack in the apoplast: A hypothesis for roles of O, N, and S species in regulatory and polysaccharide cleavage events during rapid abscission by Azolla. Plant Sci. 217:120–126.

Dhindsa, R.A., P. Plumb-Dhindsa, and T.A. Thorpe. 1981. Leaf senescence: Correlated with increased permeability and lipid peroxidation and decreased levels of superoxide dismutase and catalase. J. Exp. Bot. 32:93–101.

Dixon, D.P., I. Cummins, D.J. Cole, and R. Edwards. 1998. Glutathione-mediated detoxification system in plants. Curr. Opin. Plant Biol. 1:258–266.

Djungarguren, M., D.D. Devi, A.K. Shanker, J.A. Sheeba, and Y.U. Barathram. 2004. The role of nitrophenol on delaying abscission of tomato flowers and fruits. Food Agr. Environ. 2:183–186.

Doderer, A., I. Kokkelink, S. Van Der Veen, B.E. Valk, A.W. Schram, and A.C. Douma. 1992. Purification and characterization of two lipoygenase isoenzymes from germinating barley. Biochim. Biophys. Acta 1120:97–104.

Ebel, R.C., J.K. Burns, K.T. Morgan, and F.M. Roka. 2010. Abscission agent application and canopy shaker frequency effects on mechanical harvest efficiency of sweet orange. Hort-Science 45:1079–1083.

Estornell, L.H., J. Agusti, P. Merelo, M. Talon, and F.R. Tadeo. 2013. Elucidating mechanisms underlying organ abscission. Plant Sci. 199–200:48–60.

Fry, S.C., J.C. Dumville, and J.G. Miller. 2001. Fingerprinting of polysaccharides attacked by hydroxyl radicals in vitro and in the cell walls of ripening pear fruit. Biochem. J. 357:729–737.

Grailiot, V., F. Tomaartij, J.P. Cravedi, and M. Audebert. 2012. Evidences of the in vitro genotoxicity of methyl-pyrazole pesticides in human cells. Mutat. Res. 748:8–16.

Hartmmond, C., R. Yuan, J.K. Burns, A. Grant, and W.J. Kender. 2000. Citrus fruit abscission induced by methyl-jasmonate. J. Amer. Soc. Hort. Sci. 125:547–552.

Hassan, G.S., H.H. Kadry, S.M. Abou-Seri, M.M. Ali, and A.E. El-Din Mahmoud. 2011. Synthesis and in vitro cytotoxic activity of novel pyrazolo[3,4-d]pyrimidines and related pyrazole hydrazones towards breast adenocarcinoma MCF-7 cell line. Bioorg. Med. Chem. 19:6808–6817.

He, Y., H. Fukushige, D.F. Hildebrand, and S. Gan. 2002. Evidence supporting a role of jasmonic acid in Arabidopsis leaf senescence. Plant Physiol. 128:876–884.

Heath, R.L. and L. Packer. 1968. Photoperoxidation in isolated chloroplasts. I. Kinetics and stoichiometry of fatty acid peroxidation. Arch. Biochem. Biophys. 125:189–198.

Henry, E.W. 1975. Peroxidase in tobacco abscission zone tissue. III. Ultrastructural localization in thylakoids and membrane bound bodies of chloroplast. J. Ultrastruct. Res. 52:289–299.

Kossuth, S.V., R.H. Biggs, and V.M. Winkler. 1978. Uptake and distribution of the herbicide 5-chloro-3-methyl-4-nitro-1H-pyrazole in ‘Valencia’ and ‘Hamlin’ oranges. J. Amer. Soc. Hort. Sci. 103:20–22.

Kumar, N. and R.C. Ebel. 2015. Oxidative metabolism in ‘Valencia’ sweet orange (Citrus sinensis (L.) Osbeck) flavedo tissue treated with the abscission agent 5-chloro-3-methyl-4-nitro-1H-pyrazole (CMNP). J. Hort. Sci. Biotechnol. 90(4):413–418.

Lewis, M.W., M.E. Leslie, and S.J. Liijegren. 2006. Plant separation: 50 ways to leave your mother. Curr. Opin. Plant Biol. 9:59–65.

Li, K.-T., J.K. Burns, and J.P. Sypertsen. 2008. Recovery from physiological stress by foliar application of fruit-loosening abscission compounds to citrus. J. Amer. Soc. Hort. Sci. 133:535–541.
Licciardello, C., N. D’agostino, A. Traini, G.R. Recupero, R. Frusciante, and M.L. Chiusano. 2014. Characterization of the glutathione S-transferase gene family through ESTs and expression analyses within common and pigmented cultivars of Citrus sinensis (L.) Osbeck. BMC Plant Biol. 14:39.

Makino, N., K. Sasaki, K. Hashida, and Y.A. Sakkura. 2004. A metabolic model describing the H2O2 elimination by mammalian cells including H2O2 permeation through cytoplasmic and peroxisomal membranes: Comparison with experimental data. Biochim. Biophys. Acta 1673:149–159.

Nakano, Y. and K. Asada. 1981. Hydrogen peroxide is scavenged by ascorbate specific peroxidases in spinach chloroplast. Plant Cell Physiol. 22:887–892.

Perl-Treves, R. and A. Perl. 2002. Oxidative stress: An introduction, p. 1–33. In: D. Inze and Van Montagu (eds.). Oxidative stress in plants. Taylor & Francis, London, UK.

Sakamoto, M., I. Munemura, R. Tomita, and K. Kobayashi. 2008. Involvement of hydrogen peroxide in leaf abscission signaling, revealed by analysis with an in vitro abscission system in capsicum plants. Plant J. 56:13–27.

Smith, I.K., T.L. Vierheller, and C.A. Thorne. 1988. Assay of glutathione reductase in crude tissue homogenates using 5,5′-dithiobis (2-nitrobenzoic acid). Anal. Biochem. 175:408–413.

Taylor, J.E. and C.A. Whitelaw. 2001. Signals in abscission. New Phytol. 151:323–339.

Velikova, V., I. Yordanov, and A. Edreva. 2000. Oxidative stress and some antioxidant systems in acid rain-treated bean plants: Protective roles of exogenous polyamines. Plant Sci. 151:59–66.

Vicentini, C.B., D. Mares, A. Tartari, M. Manfrini, and G. Forlani. 2004. Synthesis of pyrazole derivatives and their evaluations as photosynthetic electron transport inhibitors. J. Agr. Food Chem. 52:1897–1906.

Yang, Z., X. Zhong, Y. Fan, H. Wang, J. Li, and X. Huang. 2015. Burst of reactive oxygen species in pedicle-mediated fruit abscission after carbohydrate supply was cut off in longan (Dimocarpus longan). Front. Plant Sci. 6(360):1–10.

Yuan, R. and J.K. Burns. 2004. Temperature factor affecting the abscission response of mature fruit and leaves to CMN-pyrazole and ethephon in ‘Hamlin’ oranges. J. Amer. Soc. Hort. Sci. 129:287–293.