Oxidative Stress and Superficial Scald of Apple Fruit

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Superficial scald of apples and pears, which manifests as brown or black patches on the fruit skin, results from necrosis of the hypodermal cortical tissue (Bain and Mercer, 1963) and is thought to be induced by oxidation products of the sesquiterpene (E,E)-α-farnesene (Anet, 1972a; Huelin and Coggiola, 1970a, 1970b; Huelin and Murray, 1966). Synthesis of sesquiterpenes in plant tissues occurs mainly via the cytosolic mevalonic acid (MVA) pathway, which is initiated by the enzyme 3-hydroxy-3-methylglutaryl-CoA reductase (HMGR) (Eisenreich et al., 2001). Studies utilizing radiolabeled precursors (Rupasinghe et al., 2001) or a statin inhibitor of HMGR (Ju and Curry, 2000a) indicated that in apple peel α-farnesene is synthesized almost exclusively via the MVA pathway, rather than the chloroplastic deoxyxylulose phosphate pathway. Fruit of scald-susceptible apple cultivars such as ‘Granny Smith’, ‘Law Rome’, and ‘Red Delicious’ typically exhibit a burst of α-farnesene synthesis shortly after they are placed in low-temperature storage, which results in a marked accumulation of the sesquiterpene in the skin and epicuticular coating during the first 8 to 12 weeks (Fig. 1a) (Anet, 1972a; Huelin and Coggiola, 1968; Whitaker et al., 1997, 1998). The concentration of α-farnesene subsequently plateaus and then declines as the skin and epicuticular coating of apple fruit). As a group these compounds were called conjugated trienes (CTs), and in a concurrent study by Huelin and Coggiola (1970a) it was shown that oxidation of α-farnesene to CTs was inhibited by the antioxidant DPA both in vitro (in hexane solution) and in vivo (in the epicuticular coating of apple fruit).

On the basis of these findings, it was long assumed that the CTs that accumulated on the surface of apples as a consequence of α-farnesene oxidation during low-temperature storage were identical to those produced during in vitro autoxidation of α-farnesene. However, Rowan et al. (1995) determined that >96% to 99% of the CTs in hexane-dip extracts of cold-stored ‘Granny Smith’ fruit were conjugated trienols, the 7E,9E

OXIDATION PRODUCTS OF α-FARNESENE

It was first proposed that oxidation products of α-farnesene are the causal agents of superficial scald in reports by Huelin and Murray (1966) and Huelin and Coggiola (1968). Anet (1969) isolated α-farnesene from apples and characterized the products of its in vitro autoxidation. The principal oxidation products were identified as 7E,9E and 7E,9Z isomers of a farnesyl 6-hydroperoxide (Fig. 2a and b, respectively) and mixed diastereoisomers of a farnesyl endoperoxide-hydroperoxide (Fig. 2c), all of which had UV spectra with a series of absorbance maxima at 250, 269, and 281 nm, reflecting the presence of three conjugated double bonds in the structure. As a group these compounds were called conjugated trienes (CTs), and in a concurrent study by Huelin and Coggiola (1970a) it was shown that oxidation of α-farnesene to CTs was inhibited by the antioxidant DPA both in vitro (in hexane solution) and in vivo (in the epicuticular coating of apple fruit).

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Fig. 1. Accumulation of α-farnesene (a) and its conjugated trieniol oxidation products (b) on the surface of scald-susceptible ‘Granny Smith’, ‘Law Rome’, and ‘Red Delicious’ apples stored in air at 0.5 °C for up to 28 weeks. Peel tissue samples were extracted with hexane and aliquots of the hexane extracts were analyzed by high-performance liquid chromatography with UV detection at 232 and 269 nm.
intriguing question of how the conjugated trienols arise in the skin and scald severity in 'Red Delicious' apples. Moreover, Rowan et al. (2000) found no correlation between MHO concentrations and ship to the scald index in 'Granny Smith' fruit. In contrast, Rupasinghe that MHO production and accumulation showed a strong positive relation-
a burst of MHO upon removal from storage, coincident with a marked
that MHO does in fact induce scald (Mir et al., 1999; Wang and Dilley,
He concluded that the levels of MHO produced in vivo were probably too
and proposed that it is formed via decomposition of a 6-alkoxy radical
subsequently determined that the volatile ketone 6-methyl-5-hepten-2-
al., 1919). In the 1950s and 1960s, several groups examined the correlation
enzymatic, c may be a poor substrate compared with a and b.

Fig. 2. Primary hydroperoxide (a and b) and endoperoxide-hydroperoxide (c)con-
jugated triene autoxidation products of ω-farnesene and their corresponding
conjugated trienol reduction products d, e, and f, respectively. During cold
storage, the Δ7,9Δ and Δ7,9Δ conjugated trienol isomers d and e, respectively,
accumulate on the surface of scald-susceptible apples, whereas the
endoperoxide-alcohol f does not. It is currently not known whether reduction of
a and b to d and e in vivo is enzymatic or nonenzymatic.

and Δ7,9Δ isomers of 2,6,10-trimethyl-2,7,9,11-dodecatetraene-6-ol (Fig. 2d and e, respectively), present in a ratio of ω9:1. Analysis of ex-
tracts from cold-stored 'Delicious' apples in my laboratory corroborated this
discovery, and again the ratio of the Δ7,9Δ isomer d to the Δ7,9Δ
isomer e was ω9:1 (Whitaker et al., 1997). These conjugated trienol
isomers were obtained in about the same ratio when Anet (1969) used
sodium borohydride to reduce, and thereby stabilize, the corresponding
6-hydroperoxides a and b that were present in a thin-layer chromatog-
raphy fraction of the ω-farnesene autoxidation products. This raises the
intriguing question of how the conjugated trienols arise in the skin and epidermal
wax of apple fruit. Rowan et al. (1995) observed that as the
6-hydroperoxide a slowly decomposed in pentane at -20 °C, ω10% was
converted to the trienol d. Perhaps this conversion is more efficient at 0 °C
in the waxy matrix on the surface of apple fruit. Another explanation
is that the 6-hydroperoxide isomers a and b are enzymatically reduced to
the corresponding alcohols d and e by a peroxidase that acts on alkyl
hydroperoxide substrates. A lipid hydroperoxide-dependent glutathione
peroxidase encoded by a gene that is upregulated in response to active
oxygen species (O2A) in sesening leaves (Navabour et al., 2003) might be
an example of such an enzyme in plant tissues. Another related ques-
tion is why do the conjugated trienols often accumulate to high levels in
apple fruit, whereas isomers of the endoperoxide-alcohol (Fig. 2f), derived
by reduction of the endoperoxide-hydroperoxide autoxidation product c,
do not accumulate. This may be due to both the extreme instability of the
endoperoxide-hydroperoxide c and its limited production. The two
occur via further oxidation of the endoperoxide a (Anet, 1969; Rowan et
al., 1995, 2001). In addition, if reduction of the peroxides to alcohols is
enzymatic, c may be a poor substrate compared with a and b.

Early research on scald indicated a key role of toxic volatiles (Brooks et
al., 1919). In the 1950s and 1960s, several groups examined the correlation
of naturally occurring apple volatiles with development of the disorder
but failed to identify a likely causal agent (Meigh, 1970). Anet (1972b)
subsequently determined that the volatile ketone 6-methyl-5-hepten-2-
one (MHO) is a major secondary product of ω-farnesene autoxidation
and proposed that it is formed via decomposition of a 6-alkoxy radical
intermediate derived from the primary autoxidation products a, b, and c.
He concluded that the levels of MHO produced in vivo were probably too
low to have a toxic effect and suggested that cell damage was most likely
caused by free radicals generated during the reaction that yields MHO.
However, more recently there has been a resurgence of the hypothesis
that MHO does in fact induce scald (Mir et al., 1999; Wang and Dilley,
2000a, 2000b). Mir et al. (1999) observed that 'Cortland' apples released
a burst of MHO upon removal from storage, coincident with a marked
intensification of scald symptoms, and Wang and Dilley (2000b) reported
that MHO production and accumulation showed a strong positive relation-
ship to the scald index in 'Granny Smith' fruit. In contrast, Rupasinghe et
al. (2000b) found no correlation between MHO concentrations and
scald severity in 'Red Delicious' apples. Moreover, Rowan et al. (2001)
did not observe an increase in scald development in apples treated with
MHO. They did, however, suggest that production of MHO by the fruit
is likely to be a sensitive indicator of oxidative processes leading to scald.
Not surprisingly, MHO is released during in vitro autoxidation of both
the CT hydroperoxides a and b (Fielder et al., 1998) and the conjugated
trienols d and e (Whitaker and Saffner, 2000). Kinetics of the autoxidation
of d and e that yielded MHO were indicative of a free radical-mediated
reaction, which was much more rapid at 20 than at 0 °C.

ROLE OF SYNTHETIC AND NATURAL ANTIOXIDANTS

The efficacy of DPA treatment as a means of scald control was dis-
covered by Smock (1955), and Huelin (1964) determined that several
related amon compounds, including dibenzylamine, dicyclohexylamine,
and N-benzylamine, were also effective. As mentioned previously,
Huelin and Coggiola (1970a) showed that DPA inhibited autoxidation of
ω-farnesene to CTs both in vitro and in vivo. Anet and Coggiola (1974)
subsequently evaluated the ability of a variety of lipophilic
antioxidants to control ω-farnesene autoxidation. They found that all
classes of antioxidants tested were capable of inhibiting autoxidation of
the sesquiterpene in vitro, whereas only DPA and related amine-type
compounds effectively blocked the accumulation of CTs in vivo. These
results prompted investigation of possible modes of action of DPA in
scald prevention other than inhibition of ω-farnesene autoxidation. Sev-
eral studies showed a reduction or delay in ω-farnesene production in
DPA-treated fruit (Du and Bramlage, 1993, 1994a; Huelin and Coggiola,
1968; Whitaker, 2000), but this was not consistently observed (Huelin
and Coggiola, 1970b; Lurie et al., 1989). There were also reports of ef-
fects of DPA on fruit physiology, including reduced rates of respiration,
ethylene production, and senescence (Du and Bramlage, 1994a; Lurie et
al., 1989; Whitaker, 2000). These effects are not necessarily persistent,
since respiration and ethylene evolution were the same in control and
DPA-treated 'Cortland' apples removed from storage after six months
(Mir and Beaudry, 1999). Recent evidence indicates that DPA reduces
respiration in plant cells by inhibiting mitochondrial electron transport
(Purvis and Gengege, 2003). It would be interesting to know if related
amine-type antioxidants that also prevent scald, such as N-isopropyl-N'
-phenyl-p-phenylenediamine (Anet and Coggiola, 1974), have a similar
effect on plant mitochondria.

Long after it was proposed that autoxidation products of ω-far-
nesene are involved in scald development, the hypothesis was modified to
include a role of endogenous antioxidants in delaying the accumulation
of CTs, particularly in late-harvest and scald-resistant fruit (Anet, 1972a;
isolated 11 lipophilic antioxidants in hexane-dip extracts of apple fruit
from 16 cultivars. In general, he observed that a substantial decline in
these antioxidants was correlated with the onset of ω-farnesene autoxi-
dation and the eventual development of scald. The lipophilic antioxidant
compounds had a pair of UV absorbance maxima ranging from 204 to 207 nm and 256 to 263 nm, which resembled UV spectra
of p-hydroxybenzoic acids and esters. Huelin and Coggiola (1970a) had
previously found a correlation between 258-nm absorbance in hexane-dip
extracts and a low incidence of scald in 'Crofton' and late-pick 'Granny
Smith' apples, and more recently Du and Bramlage (1993, 1994a) reported
similar findings for 'Cortland' and 'Delicious' apple fruit. Detection
of a broad, prominent UV maximum at 258 nm in hexane extracts of
peel tissue from scald-resistant 'Gala' apples prompted me to isolate the
compounds responsible for this UV absorbance (Whitaker, 1998).
They were identified as a family of mainly saturated, long-chain fatty-
acid esters of E- and Z-p-coumaryl alcohol, shown in Fig. 3 (Whitaker
et al., 2001), and on the basis of their UV spectra, it appears likely that
Anet’s three unidentified antioxidant compounds belonged to this group.
Although these phenolic fatty-acid esters were quite abundant in scald-
resistant 'Gala' and were found to have antioxidant activity, they were
undetectable in the epicuticular wax of equally scald-resistant 'Empire'
fruit (Whitaker, 1998). Thus, these lipophilic antioxidant compounds may
serve to delay the onset of ω-farnesene autoxidation and to generally
prevent the accumulation of free radicals and reactive oxygen species
(Anet, 1974; Barden and Bramlage, 1994; Gallarani et al., 1990; Meir
and Bramlage, 1988).
GENES AND ENZYMES INVOLVED IN α-FARNESENE SYNTHESIS

Synthesis of the acyclic sesquiterpene (E,E)-α-farnesene in apple peel tissue occurs mainly via the cytosolic MVA pathway (Ju and Curry, 2000a; Rupasinghe et al., 2001), as depicted in the simplified schematic in Fig. 4. Before the first committed step of the pathway, the conversion of HMGR-CoA to MVA by HMGR (Fig. 4, step 1), three acetetyl-CoA molecules are condensed to yield HMG-CoA. Because the pool of acetetyl-CoA increases substantially with the advent of the respiratory climacteric in apple fruit, availability of this substrate may partly determine the rate of α-farnesene production during the initial 4 to 8 weeks of storage (Rupasinghe et al., 2001). Ju and Curry (2000a) found that dip treatment of whole apples with the potent HMGR inhibitor lovastatin all but eliminated α-farnesene production. They also reported evidence that ethylene-induced transcription and translation of HMGR gene(s) is required for α-farnesene synthesis in peel tissue of apple fruit (Ju and Curry, 2000b). On the basis of these reports, and the commonly ascribed role of HMGR as the rate-limiting enzyme in isoprenoid synthesis via the MVA pathway (Goldstein and Brown, 1990; Hartmann et al., 2000), HMGR is considered to be a likely control point in α-farnesene production in apple fruit.

In plants, HMGR is typically encoded by small families of genes, designated as HMG1, HMG2, HMG3, etc., with a high degree of sequence homology among the cDNA coding regions (McCaskill and Croteau, 1997). Expression of different HMGR genes and isoforms is subject to diverse modes of regulation; expression of a given gene may be constitutive, tissue-specific, or inducible by hormones, infection, or wounding (McCaskill and Croteau, 1997; Pirironen et al., 2000). Since ethylene clearly plays a key role in the stimulation of α-farnesene production in harvested apples (Fan et al., 1999; Gong and Tian, 1998; Rupasinghe et al., 2000a; Shaham et al., 2003; Watkins et al., 2000), it is probable that expression of an HMGR gene specifically involved in α-farnesene synthesis would be strongly induced by ethylene.

Current research is aimed at identification of an ethylene-inducible HMGR gene in apple peel tissue that could be selectively suppressed to reduce α-farnesene synthesis. Complete or partial cDNAs encoding three HMGR isoforms (HMG1, HMG2, and HMG3) have been cloned from commercial apple cultivars. A full-length cDNA of HMG1 has been obtained from ‘Delicious’, ‘Granny Smith’, and ‘Law Rome’ apples (GenBank accession numbers AF315713, AY039230, and AY043490, respectively). Because this isogene is abundantly expressed at harvest and through the first 8 to 16 weeks of storage in both untreated and 1-MCP-treated fruit (Rupasinghe et al., 2001; Pechous and Whitaker, 2002), it is unlikely to play a specific role in α-farnesene synthesis. The very low level of HMG2 (AY043491) transcript in ‘Law Rome’ fruit during the interval of maximum α-farnesene synthesis suggests that it is also not involved (Pechous and Whitaker, 2002). In contrast, the pattern of expression of HMG3 in ‘Delicious’ peel tissue (AF316112) was wholly consistent with a role in the postharvest burst of α-farnesene production (Rupasinghe et al., 2001). In untreated controls HMG2 transcript increased sharply over the first 8 weeks of storage then declined, whereas in 1-MCP-treated fruit it was nearly absent. We have recently cloned a full-length cDNA of HMG2 from ‘Law Rome’ and are presently trying to corroborate the findings of Rupasinghe et al. (2001).

HMGR initiates the MVA pathway and thus potentially contributes to the synthesis of a wide variety of isoprenoids. It is logical, therefore, to also clone genes encoding enzymes that catalyze late steps in the α-farnesene biosynthetic pathway, with the expectation that their suppression would more specifically inhibit the production of the sesquiterpene. Farnesyl diphosphate (FDP) synthase performs the penultimate reaction of the 7α-farnesene biosynthetic pathway. When these facts are considered, it is apparent that the FDP synthase gene probably does not play a primary role in regulation of the α-farnesene biosynthetic pathway.

In contrast, α-farnesene synthase (α-FS) catalyzes the final step in production of α-farnesene (Fig. 4, step 3) and consequently this enzyme is the optimal target for specific inhibition of α-farnesene synthesis. Rupasinghe et al. (1998) demonstrated the presence of α-FS in peel tissue of ‘Delicious’ apples and identified the substrate as FDP. In a subsequent study, they partially purified and biochemically characterized the enzyme, but were unable to purify the protein to homogeneity due to loss of activity.

**Fig. 4.** Simplified schematic of the pathway of (E,E)-α-farnesene synthesis and oxidation in apple fruit. Genes have been cloned encoding enzymes that perform steps 1, 2, and 3, which are hydroxymethylglutaryl-CoA reductase, farnesyl diphosphate synthase, and (E,E)-α-farnesene synthase, respectively. Steps A1 and A2, respectively, represent autoxidation of (E,E)-α-farnesene to the 7E,9E conjugated triene 6-hydroperoxide and autoxidation of the 7E,9E conjugated trienol to yield MHO. The step labelled ? indicates the hypothetical enzymatic reduction of the major autoxidation product of α-farnesene, the 7E,9E conjugated triene 6-hydroperoxide, to the corresponding conjugated trienol. Abbreviations: HMGR-CoA, 3-hydroxy-3-methylglutaryl-coenzyme A; MVA, mevalonic acid; IPDP, isopentenyl diphosphate; DMADP, dimethylallyl diphosphate; GDP, geranyl diphosphate; and FDP, farnesyl diphosphate.
(Rupasinghe et al., 2000b). Moreover, in vitro assays of activity in apple tissue extracts were confounded by competing pyrophosphatase(s) that converted much of the FDP substrate to farnesol. In light of these difficulties, my laboratory chose to undertake cloning and characterization of the α-FS gene from peel tissue of scald-susceptible ‘Law Rome’ apples. We were successful in obtaining a full-length cDNA, AF51 (AY182241), and quite recently achieved functional expression of the enzyme, which synthesized (E,E)-α-farnesene almost exclusively (Pechous and Whiteker, 2004). Rupasinghe et al. (2000b) did not detect an increase in α-FS activity in ‘Delicious’ apples during the first few weeks of storage and saw no effect of 1-MCP treatment. These results are seemingly at odds with our Northern blot and RT-PCR analyses using ‘Law Rome’ peel tissue RNA, which showed over a twofold increase in AF51 expression in control fruit and nearly total disappearance of AF51 transcript in 1-MCP-treated fruit (Pechous and Whiteker, 2004). As mentioned above, assays of α-FS activity in apple peel tissue extracts are evidently subject to technical biases, so it is hoped that antibodies specific for AF51 will at least determine if there are changes in α-FS enzyme levels over the initial weeks of storage.

Using primers based on the ‘Law Rome’ AF51 sequence we were able to rapidly clone the corresponding cDNA from scald-resistant ‘Idared’ apples. As shown in Fig. 5, a similar percent increase in AF51 expression occurred in ‘Idared’ and ‘Law Rome’ during the first 4 weeks of storage, but AF51 transcript was consistently about fourfold less abundant in scald-resistant ‘Idared’ than in scald-susceptible ‘Law Rome’. This raises the interesting question of whether the promoters differ for the otherwise nearly identical AF51 genes from these two cultivars, or perhaps the rate of turnover of AF51 mRNA differs. Isolation of genomic clones of these genes including the 5′-flanking region will be required to analyze their promoters. For both AF51 and HMG2, which appear to be upregulated by ethylene, this approach could provide valuable information about ethylene response elements in the promoter region that are involved in the ethylene-induced increase in expression (Deikman et al., 1997; Solano et al., 1998).

**ALTERNATIVE HYPOTHESIS OF SCALD INDUCTION**

There are dissenting views concerning the root cause of apple scald and the genetic basis of scald susceptibility or resistance. A number of researchers have proposed that scald arises as a consequence of more general oxidative stress and that the runaway autoxidation of α-farnesene is merely a secondary manifestation of unchecked free radical reactions (Rao et al., 1998; Rupasinghe et al., 2000b). Regardless of the role of α-farnesene oxidation products, scald development almost certainly involves the adverse effects of oxidative stress that occurs with prolonged storage at chilling temperatures (Du and Bramlage, 1994b, 1995; Rao et al., 1998; Watts and Purvis, 1995; Watkins et al., 1995). Purvis and Gegogeine (2003) recently proposed that scald arises as a result of superoxide radicals generated by perturbation of the mitochondrial electron transport chain. Several studies have evaluated the accumulation of peroxides and lipid peroxidation products in relation to the activities of enzymes that detoxify AOS in peel tissue of scald-susceptible and -resistant apples. Du and Bramlage (1994b, 1995) found no marked changes in peroxidation or activities of antioxidative enzymes related to scald development in ‘Cortland’, ‘Delicious’, and ‘Empire’. In marked contrast, Rao et al. (1998) found that in fruit of susceptible and resistant ‘White Angel’ x ‘Rome Beauty’ hybrid selections the occurrence and severity of scald were strongly correlated with increasing levels of H₂O₂, increasing lipid peroxidation, and decreasing peroxidase and catalase activities. Quite recently, Shaham et al. (2003) compared the activities of five antioxidative enzymes, as well as lipophilic and water-soluble antioxidative factors, in untreated, heat-treated, and 1-MCP-treated ‘Granny Smith’ apples. In 1-MCP-treated fruit, the complete prevention of scald was better correlated with antioxidative capacity than with antioxidative enzyme activities, whereas in heat-treated fruit, a substantial delay in scald development was more closely associated with elevated antioxidative enzyme activities.

There are several studies and lines of evidence that challenge the α-farnesene oxidation−scald induction hypothesis. Rupasinghe et al. (1998, 2000b) determined that α-farnesene synthesis and content were consistently threefold greater in scald-free than in scald-developing apples. In addition, they found no correlation of α-farnesene production, α-FS activity, or conjugated triene accumulation with scald susceptibility in 11 commercial apple cultivars (although scald severity was apparently not evaluated for the fruit used in this study). Rao et al. (1998) also observed that accumulation of α-farnesene and CTs was poorly correlated with scald susceptibility among their ‘White Angel’ x ‘Rome Beauty’ hybrid selections. In a subsequent investigation with these hybrid lines (Whitaker et al., 2000), it was found that although high levels of α-farnesene and conjugated trienes occurred in the lines that scalded severely, there were scald-resistant lines that accumulated equally high levels, and lines that accumulated very low levels yet developed mild to moderate scald. On the basis of these results, it was concluded that oxidation products of α-farnesene are not essential for scald development in fruit with severely compromised antioxidative defenses, but free radicals and/or toxic volatiles generated by oxidation of α-farnesene can exacerbate scald symptoms.

**SUMMARY**

Superficial scald is a costly storage disorder that develops in fruit of susceptible apple cultivars such as ‘Granny Smith’ and ‘Law Rome’ after several months at low temperature. Despite intensive investigation, the biochemical mechanism and genetic basis of apple scald remain unknown. Evidence indicates that scald is induced by oxidative stress. The prevailing hypothesis holds that oxidation products of the sesquiterpene α-farnesene are directly involved. A dramatic rise in α-farnesene synthesis occurs shortly after apples are placed in storage, and oxidation of the accumulated α-farnesene proceeds rapidly after 6 to 8 weeks, particularly in air-stored fruit. The primary in vitro autoxidation products of α-farnesene are conjugated trienes (CTs) composed of several hydroperoxide and endoperoxide-hydroperoxide isomers. However, >95% of the CTs that accumulate in the skin of cold-stored apples are two conjugated trienes, 9E and 9Z isomers of 2,6,10-trimethyl-7,9,11-tetraen-3-ol, a reduced form of the major hydroperoxide autoxidation products. Application of synthetic CT peroxides or alcohols to apples induces symptoms indistinguishable from superficial scald. Autoxidation of CTs yields the volatile 6-methyl-5-hepten-2-one, which has been implicated as a possible causal agent for scald, but may simply be indicative of damaging free radical-mediated reactions. For the most part, synthesis of α-farnesene and accumulation of CTs during apple storage are correlated with the subsequent incidence and severity of scald development. However, exceptions have been noted, where some scald occurs despite low levels of CTs or there is no scald despite moderately high levels of CTs. Further evidence in support of the α-farnesene oxidation−scald induction hypothesis was the recent finding that prestorage treatment of scald-susceptible apples with 1-MCP, a blocker of ethylene action, greatly reduced α-farnesene synthesis and abolished scald development.
Yet it can still be argued that ethylene-induced processes other than α-farnesene production lead to scald, and it has been proposed that scald results from more general oxidative stress, perhaps triggered by disruption of mitochondrial electron transport at low temperature and the consequent production of superoxide. Molecular genetic disruption of genes controlling α-farnesene biosynthesis is a strategy that should prove or disprove the direct role of α-farnesene oxidation in the induction of superficial scald. Logical targets for gene knockouts are genes encoding: 1) a sesquiterpene pathway-specific isozyme of 3-hydroxy-3-methylglutaryl-coenzyme A reductase (HMGR), and 2) α-farnesene synthase, the last enzyme in the pathway that converts FDP to (E,E)-α-farnesene. Work is currently in progress to clone and characterize these genes from scald-susceptible and -resistant apple cultivars.

Literature Cited