Abstract. Seedless table grapes (Vitis vinifera L.) cv. Crimson Seedless were exposed to (E)-2-hexenal vapor during cold storage to determine its potential as a fumigan for long-term control of postharvest mold. Fruit were fumigated with 0.86 or 1.71 mmol (100 or 200 µL neat compound, respectively) (E)-2-hexenal per 1.1-L container for 2 weeks during 2 °C storage. Containers were moved to 20 °C storage after 4, 8, and 12 weeks for determination of mold incidence and berry quality over 12 days. The headspace concentration of (E)-2-hexenal, measured by gas chromatography, reached a maximum of 2.5 and 4.2 µmol·L⁻¹ for 0.86 and 1.71 mmol per container, respectively, after 1 day and declined to <1 µmol·L⁻¹ for both treatments by 14 days. Upon removal from cold storage at 4, 8, and 12 weeks, the incidence of mold was significantly lower for (E)-2-hexenal–treated fruit. Control of mold by (E)-2-hexenal fumigation persisted through 12 days of 20 °C storage, even though mold generally increased in all treatments. The two levels of (E)-2-hexenal were similar in their suppression of mold. Fumigation did not affect O₂ or CO₂ concentrations within the containers, nor were fruit firmness or soluble solids content affected. Postharvest fumigation of seedless table grapes with the natural volatile compound (E)-2-hexenal shows promise for control of mold.

Table grapes may be cold stored for as long as 5 months prior to marketing, but postharvest decay organisms, such as gray mold (Botrytis cinerea Pers.), can become a significant problem limiting prolonged storage (Nelson, 1991). Repeated fumigation with SO₂ is commonly used to control postharvest molds on grapes. The technique has been refined to minimize sulfite residues left on the fruit (Luvisi et al., 1990, 1991), which is of concern because of allergic responses to sulfite residues. However, as application rates are lowered, control of postharvest mold is reduced (Smilanick and Henson, 1992; Smilanick et al., 1990). Alternate fumigants have shown potential for postharvest decay control on grapes. Hydrogen peroxide vapor effectively reduces mold on B. cinerea–inoculated table grapes without adverse effects on berry quality (Forney et al., 1991; Rij and Forney, 1995). Acetic acid vapor can also reduce mold to negligible levels on grapes (Moyls et al., 1996; Sholberg and Gaunce, 1995). Recently, we demonstrated that a group of natural volatile products from plant tissues effectively suppressed mold development on grapes during short-term storage (Archbold et al., 1997). We have focused on one compound of this group, (E)-2-hexenal, because of its ubiquity (Hatanaka, 1993) and its antifungal activity against B. cinerea (Fallik et al., 1998; Hamilton-Kemp et al., 1992). Other related products of the lipoyxygenase-lyase pathway, such as hexenal and (Z)-3-hexenal, can also suppress mold development on fruit during postharvest storage (Archbold et al., 1997; Caccioni et al., 1995; Song et al., 1996; Vaughan et al., 1995).

The objective of this work was to determine the effect of fumigation with (E)-2-hexenal on incidence of mold and berry quality of seedless table grapes during long-term storage.

Materials and Methods

‘Crimson Seedless’ grapes were harvested in California and shipped overnight to the Univ. of Kentucky on two occasions. The grapes were neither precooled nor fumigated with SO₂ prior to shipment. Upon arrival, the grapes were immediately used in the experiments.

Clusters of fruit (>150 g) that fit into 150-mL plastic clamshell containers were cut from larger clusters. The clusters were selected on the basis of uniform size, color, firmness, and freedom from evident defects or diseases. The effect of both vapor phase concentration of (E)-2-hexenal and duration of exposure to the vapor were tested in this study. Liquid (E)-2-hexenal at 0 (control), 0.86, or 1.71 mmol (100 or 200 µL, respectively) of neat compound in a 10-mL glass vial was placed within each container. Containers were loosely overwrapped, hermetically sealed in low-density polyethylene film (Type OF50, film thickness 0.0021 µm; Respire Films, West Conshohocken, Pa.) for a total volume of 1.1 L, and stored at 2 °C. The effect of duration of (E)-2-hexenal treatment was evaluated by removing the vial of (E)-2-hexenal from a subset of containers at 4 or 8 weeks. Thus, after 12 weeks of cold storage, fruit had been exposed to 0, 0.86, or 1.71 mmol (E)-2-hexenal for the initial 4, 8, or 12 weeks.

The headspace vapor concentration of (E)-2-hexenal within three overwrapped containers of each of the treatments was sampled through the film at 1, 3, 7, and 14 d of storage, as previously described (Fallik et al., 1998). At 4, 8, and 12 weeks of cold storage, the O₂ and CO₂ concentrations within the sealed bags were measured. A 5-mL aliquot of the atmosphere was collected using a 5-mL syringe, the hole was resealed with tape, and the sample was measured with an oxygen/carbon dioxide analyzer (model ZR 892 HS; Illinois Instruments, McHenry, Ill.).

To fumigate fruit for different durations, vials of (E)-2-hexenal were removed from a subset of clamshell containers at 4 and 8 weeks, before their cold-storage period had ended. Because opening the film-wrapped containers restored the O₂ and CO₂ to ambient levels, all packages were opened and clamshells were briefly removed at each interval. The clamshells were returned to the same film bag, which was resealed, and the packages were returned to cold storage.

After 4, 8, and 12 weeks of 2 °C storage, five containers of each treatment were removed from storage. After measuring the O₂ and CO₂ levels, the film and remaining vials were removed, and the containers were placed at 20 °C. Mold incidence was evaluated on three containers for each treatment every other day through 12 d. The relative number of infected fruit per container was estimated and assigned to one of five groups: no mold = 1; 10% or less of the berries with mold = 2; 11% to 30% of the berries with mold = 3; 31% to 70% of the berries with mold = 4; and >70% of the fruit with mold = 5. Three experienced evaluators independently rated each container. The mean rating of each container on each evaluation date was used for statistical analysis.

For analysis of weight loss during the experiment, all containers were weighed at the start, upon removal from cold storage, and after 4, 7, and 12 d of 20 °C storage. The percentage of weight loss at each time was calculated for each container.

Berries were sampled for quality analyses at the start of the experiments, upon removal from cold storage, and after 4 d of 20 °C
storage. Ten berries were randomly selected at the start of the experiment and from each of the two remaining nonrated containers stored at 20 °C. Berry firmness was measured using a force gauge (model DFM 10; John Chatillon and Sons, Greensboro, N.C.) mounted on a model LTC test stand (John Chatillon and Sons) and equipped with a 13-mm-wide flat head. The force required for a 2-mm compression of the berry lying horizontally on the stage with slow manual advancement of the head was recorded. Berry total soluble solids (TSS) content was then measured using a refractometer (AO Scientific Instruments, Keene, N.H.).

Two shipments of grapes from separate commercial companies were used in this work. Five containers per commercial company were set up with each shipment. Three were used for evaluation of mold incidence and weight loss and the remaining two were used for sampling berries for quality analyses. Mean compression and TSS values for each container were derived from the sampled berries.

Data were blocked by shipment date and subjected to analysis of variance (ANOVA). Ratings data were found to have acceptable homogeneity of variance and normality (SigmaStat; SPSS, Chicago), so they were not transformed. The main effects of (E)-2-hexenal level, duration of treatment, and their interaction, as well as trends within each of the main effects, were determined by ANOVA (SAS Institute, Cary, N.C.). Only main effect means for (E)-2-hexenal levels are presented, because no other effects were significant.

Results and Discussion

Fumigation of ‘Crimson Seedless’ grapes with (E)-2-hexenal significantly reduced the incidence of mold as determined upon removal of the fruit from cold storage after 4, 8, and 12 weeks (Figs. 1 and 2). The two levels of (E)-2-hexenal, 0.86 and 1.71 mmol, were equally effective. Mold developed on (E)-2-hexenal–fumigated fruit during 20 °C storage, but it remained significantly below control values through 12 d of shelf storage.

The headspace concentration of (E)-2-hexenal in ‘Crimson Seedless’ grapes after 12 weeks of 20 °C storage (Fig. 3). It was also greater in containers with no fruit than in those containing fruit, indicating that the fruit absorbed the compound. Prior work indicated both a similar trend of headspace concentration over time and absorption of compound with treatment of a single berry (Archibald et al., 1997). The film used to overwrap the containers also probably absorbed (E)-2-hexenal, contributing to the decline in headspace concentration over time (Sadler and Braddock, 1991). Headspace concentration of (E)-2-hexenal was greater with 1.71 mmol in the container than with 0.86 mmol. By the end of 2 weeks of storage, the headspace concentrations in the two treatments had declined more than 3-fold from their levels at Day 1. All bags were vented at 4-week intervals to prevent development of anaerobic conditions, and the headspace levels of (E)-2-hexenal were probably not subsequently re-established at levels near 1 µmol·L⁻¹. Subsequent attempts to measure headspace levels by GC analysis were prevented by interfering peaks, possibly due to release of plasticizers by the film.

Treatment with (E)-2-hexenal did not affect the O₂ and CO₂ levels in the sealed bags (data not shown). The O₂ levels were 15.8% ± 0.3% and CO₂ levels were 5.1% ± 0.1% after each 4-week interval of cold storage. Weight loss was also unaffected by (E)-2-hexenal treatment. Berry weight was 0.9% ± 0.1% less on removal from cold storage than at the start, and loss increased with time at 20 °C, exceeding 5% after 4 d at 20 °C. Treatment with (E)-2-hexenal also had no effect on berry firmness, which declined as percent weight loss increased. Berry TSS content was not affected by fumigation treatment or length of cold storage, averaging 19.4% ± 0.1% overall, but increased 1.4% ± 0.1% after 3 d of 20 °C storage, probably because of water loss by the berries. Overall, fumigation with (E)-2-hexenal had no adverse impact on these important berry quality traits.

Similar postharvest studies with strawberry (Fragaria ×ananassa Duch.) resulted in sig-
significant stimulation of mold development on that fruit (Fallik et al., 1998). It was estimated from headspace measurements that values of (E)-2-hexenal below 0.5 μmol·L⁻¹ led to stimulation of B. cinerea mycelial development in vitro, while values above it accompanied inhibition of mold. Headspace concentration of (E)-2-hexenal in containers of strawberry fruit, set up identically to those in the present study (E) with 0.86 mmol of compound, was 0.37 μmol·L⁻¹ after 2 d of cold storage, too low for inhibition of mold and over 5-fold below those measured with grapes. As noted previously, strawberry has a significantly greater capacity for absorption and metabolism of (E)-2-hexenal than does grape (Archbold et al., 1997). Thus, a greater headspace concentration of (E)-2-hexenal could be sustained with grapes than with strawberries, and mold was inhibited, in contrast to the results with strawberry. Although B. cinerea was the primary mold observed on control fruit, others such as Rhizopus and Penicillium were also noted, so (E)-2-hexenal probably suppressed all types present.

Introduction of an adequate amount of (E)-2-hexenal into the table grape storage environment suppressed mold development. However, complete mold suppression was not achieved, and, since levels of the compound declined from the start, repeated introduction of (E)-2-hexenal may achieve better long-term suppression. Or, (E)-2-hexenal may be combined with SO₂ to reduce fumigation frequency with the latter compound and thereby allay concerns over sulfate residues. The similarity in response to the two levels of (E)-2-hexenal indicates that the higher level provided no advantage, but a minimum appropriate level might be below that provided by 0.86 mmol of compound. The ability of (E)-2-hexenal to suppress mold organisms and its presence as a part of the volatile profile of grapes, as well as of other fruits and vegetables (Nijsen et al., 1996), makes it and similar natural volatile products attractive potential fumigants.

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