Evaluation of Ethyl Formate, Phosphine, and Their Combination to Disinfest Harvested Celery against Purple Scum Springtails

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SUMMARY. Export celery (Apium graveolens var. dulce) from Australia has been affected by a natural infestation of purple scum springtails (Hypogastrura vernalis). These insects live inside the celery head, contaminating fresh celery, but do not cause any visible damage. As a result, purple scum springtail-infested celery has led to rejection for export with an impact on market value for fresh produce. In this study, fumigation with ethyl formate (EF), phosphine (PH3), and their combination on mortality of purple scum springtails in naturally infested celery was evaluated. Laboratory experiments were conducted using concentrations of 50, 60, and 90 mg L⁻¹ of PH3 for 2, 4, and 6 hours; and 20, 30, and 40 mg L⁻¹ of EF combined with 1 mg L⁻¹ of PH3, for 2 and 4 hours at the laboratory temperature 25 °C. Complete control was achieved at 90 mg L⁻¹ of PH3 for 2 hours; however, phytotoxicity was observed in celery treated by PH3 at 2.5 mg L⁻¹ achieved 100% mortality within 6 hours, and no phytotoxicity was evident. Mortality of 100% was achieved also at 30 and 40 mg L⁻¹ EF combined with 1 mg L⁻¹ of PH3 for 2 and 4 hours exposure time; however, phytotoxicity occurred with EF alone treatments and with the combination. From these data, we conclude that PH₃ alone has potential as a fumigant for the preshipment treatment of celery infested with purple scum springtails.

Celery is an intensively cultivated and valuable horticultural export product from Australia. Celery is grown in most states with the main production areas in Victoria, Western Australia, and Queensland. There are cultivated areas of ≈235 ha per year (Horticulture Australia, 2009). Australian celery production in 2015 was more than 60,000 t, with a value of 50.1 million Australian dollars. In 2017, celery production for export increased to 3557 t (Horticulture Innovation Australia, 2017). However, this growing market has been threatened by a natural infestation of purple scum springtails, which significantly affected market access as purple scum springtails are considered a quarantine pest in many Middle East countries. Australian Quarantine does not allow export of plant products containing living organisms.

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Ethyl formate is a naturally occurring compound that has been evaluated as a fumigant for stored grain, fruit, and vegetable applications and has been commercially adopted because of its valuable characteristics of rapid kill and consumer safety (Agarwal et al., 2015; Kim et al., 2013; Ren and Mahon, 2006; Ren et al., 2008). However, EF is highly absorbed in products such as strawberries (Fragaria × ananassa), apples (Malus × domestica), and several types of cut flowers because of their high moisture content and because EF is highly soluble in the water (Agarwal et al., 2015; Lee et al., 2013; Simpson et al., 2004). Some studies (Tarr et al., 2007) indicated that EF, as well as EF in combination with 10% carbon dioxide (CO₂), could be used for control of sawtoothed grain beetle (Oryzaephilus surinamensis) and confused flour beetle (Tricholium confusum) on dry vine fruit. Different levels of EF combined with low concentration of PH₃ 0.5 mg·L⁻¹ for 2 h have resulted in high mortality of three species of aphis (cotton aphid (Aphis gossypii), green peach aphid (Myzus persicae), and turnip aphid (Lipaphis erysimi)), there was significantly different in the lethal concentration LCT₅₀ and LCT₉₀ values in comparison with EF and PH₃ alone (Lee et al., 2014). Also, EF can cause high mortality in adults and nymphs of onion thrips (Thrips tabaci) on onion (Allium cepa) after 2 h exposure (Van Epenhuijsen et al., 2007).

Phosphine is a commercially available fumigant, which has been used for decades to control insects in stored grain. PH₃ in combination with CO₂, such as ECO₂FUME*** (2% PH₃ + 98% CO₂) Cytec Industries, Dongbu Hannong Co., Seoul, Korea), has been used for disinfection of postharvest horticultural products (Jamieson et al., 2012; Williams et al., 2000). PH₃ fumigation at low temperature has been shown to have high efficacy for postharvest control of western flower thrips (Frankliniella occidentalis) on lettuce (Lactuca sativa), broccoli (Brassica oleracea var. italica), asparagus (Asparagus officinalis), and strawberry (Liu, 2008). However, fumigation with PH₃ has been known to cause a significant reduction in shelf and vase life of cut flowers with both 5.5 and 11 mg·L⁻¹ for 6 h exposure time to four types of cut flowers for controlling greenhouse thrips (Heliothrips haemorrhoidalis) and green peach aphids (Karunaratne et al., 1997).

There has been no published work on control of purple scum springtails on harvested celery. Under our current study, we researched the efficacy of EF, PH₃, and their combination for control of purple scum springtails on celery. In this preliminary study, optimal fumigant concentration, exposure period, and evaluation of phytotoxic damage to celery bunches are reported.

**Materials and methods**

**Celery and target pest**

Celery infested with purple scum springtails was supplied by the Mandogalup Celery Farm (Sumich Group, Kwinana, Western Australia, Australia) located 23 km south of Perth, Western Australia (lat. 32.20°S, long. 115.84°E). Celery bunches were weighed with a digital balance, and their weight ranged between 686 and 1020 g per bunch.

The purple scum springtail species found in Western Australia were identified taxonomically by Majer et al. (2014) and current species have been confirmed by the Department of Primary Industries and Regional Development by mounting a number of specimens and following the Greenslade et al. (2014) classification key. Celery samples were stored in a cold room at 15 °C at Murdoch University (Murdoch, Australia) for 1–2 d.

**Reagents and apparatus**

Ethyl formate 97% purity was supplied by Sigma-Aldrich Co. (Castle Hill, Australia). A cylindrical PH₃ gas supplied with nitrogen (N₂) [1.4% PH₃ with 98.6% N₂] was supplied by BOC Australia (North Ryde, Australia). Ten-liter gas sampling bags (SKC Tedlar; Air-Met Scientific, Perth, Australia) were used for the introduction of gas during fumigation. The fumigation chamber had two gas sampling ports located on the side of the fumigation chamber. Each lid was fitted with gas-tight seals, which were pressure-tested before use.

For testing pressure halving time of the fumigation chambers, a digital manometer was connected to one of the compression fittings after securing the metal drum with lids. An air pump was used to apply an increased pressure of 250 Pa to the sealed drum via another compression fitting. Once the digital manometer displayed a value of 250 Pa, the air pump was turned off and the ball valve tap closed. The air pressure was then turned on until the value displayed was 125 Pa, giving the drum’s “pressure half-life.” A half-life of more than 10 min indicated a well-sealed chamber suitable for fumigation.

**Determination of EF and PH₃ concentration**

Ethyl formate was analyzed using a portable GC (Companion 600; DPS Instruments, Rancho Cucamonga, CA) installed with a flame ionization detector (FID). A 30 m × 0.53-mm (i.d.) 0.5-µm capillary column (Zebron model ZB-WAX, B13844, part no. 7HK-G007-17; Phenomenex Co., Castle Hill, Australia) was used at an oven temperature of 95 °C. N₂ carrier gas was used at a flow rate of 6 mL·min⁻¹ at 103 Pa. The concentrations of fumigants were monitored at different time intervals (depending on exposure time for each chambers to create negative pressure in preparation for fumigation.

A 100-μL syringe (005250; SGE, Melbourne, Australia) was used for the injection of gas samples into the gas chromatograph (GC) and transfer liquid EF into fumigation chambers. A 5-μL syringe (SR-GT; SGE) was used to transfer liquid EF to prepare gas standards.

**Fumigation chamber**

A cylindrical stainless-steel chamber (60 × 36 cm i.d.) with a capacity of 61 L was used. Each chamber was equipped with compression fittings (Swagelok Co., Perth, Australia) for attachment to a diaphragm pump between the top and bottom rim of the chamber. The removable metal lids were fitted with a centrally located septum for the introduction of gas during fumigation. The fumigation chamber had two gas sampling ports located on the side of the fumigation chamber. Each lid was fitted with gas-tight seals, which were pressure-tested before use.

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Phosphine concentrations were determined at timed intervals using an GC (HP 5890, Series II; Hewlett-Packard Co., Wilmington, DE) equipped with a flame photometric detector (FPD) with a phosphorus filter following isothermal separation on a 30 m × 0.25-mm (i.d.), 0.25-μm, capillary column (Varian part no. CP8944; Sigma-Aldrich Co.) at oven temperature 50 °C and inlet temperature 105 °C. For FPD, the flow of hydrogen and air was 50 and 100 mL·min⁻¹, respectively. Hydrogen was used as the carrier gas with a constant flow of 2 mL·min⁻¹. After fumigation processes, two replicate injections were administered through each gas sampling port and injected into GC (two injections from the top and also two other injections from the bottom port of the fumigation chamber). Both EF and PH₃ concentrations were calculated based on peak areas against external EF and PH₃ gas standards.

**Fumigation of purple scum springtail infested celery with EF**

Three treatments consisting of 50, 60, and 90 mg·L⁻¹ of EF plus an untreated check were investigated for 1, 2, and 3 h the duration of exposure. Celery bunches were transferred after storage for 1–2 d from the 15 °C cold room to the 23 to 25 °C laboratory for up to 4 h before fumigation. Two replication fumigation chambers were used to treat the celery. Each chamber contained between 8545 and 9261 g for 10 bunches of celery held together with rubber bands and the bottom part of celery facing down (Fig. 1), with each chamber having ≈8% loading ratio of celery. Treatments of fumigation were carried out sequentially, starting with the first, second, and the third concentration of fumigants. The remaining celery bunches from the first experiment were stored at 15 °C cold room for next experiments. Ethyl formate experiments with two replication containers were repeated a second time for confirmation of phytotoxicity only by using fresh celery bunches that were stored for less than 24 h at 15 °C cold room. The untreated controls were prepared the same way in duplicate as EF treatments, except that no fumigant was applied. Before fumigation, the chambers were sealed tightly with locking rings attached to the lid.

Gas sampling bags were used to prepare fumigant at experimental concentrations. Calculated volumes of liquid EF were injected into 1-L gas sampling bags and immersed in a hot water bath at 90 °C for 10 min for complete vaporization of EF. Vaporized EF inflated the gas sampling bags. To facilitate injection of vaporized EF into experimental chambers, ≈3 L of air was sucked out using an air pump connected to the sealed chamber. The EF concentrations prepared in the gas sample bag were sucked into the chamber under negative pressure to balance the air pressure with ambient. Ethyl formate gas standard was prepared in a 1-L erlenmeyer flask by taking out the estimated amount of air from the flask and injecting a calculated amount of liquid EF to the small piece of filter paper attached under the tight lid fitted with a rubber septum of the erlenmeyer flask. Headspace samples were taken from both gas sampling ports (top and bottom) of the fumigation chamber immediately and then 2 and 4 h later for each concentration (50, 60, and 90 mg·L⁻¹) along with gas standards and injected in duplicate into GC-FID. The fumigation was conducted at a laboratory temperature between 23 and 25 °C.

**Fumigation of infested celery with PH₃**

Celery samples were brought from the 15 °C cold room and left for ≈4 h to bring to ambient laboratory temperature of 23 to 25 °C before fumigation. Ten celery bunches were placed in each chamber, with a loading ratio of 9.2%. About 3 L of air were removed from each chamber before PH₃ injection and balanced with ambient air pressure. The same procedure and the number of celery bunches were applied for untreated check treatments.

Four concentrations of PH₃ were used: 1, 1.5, 2, and 2.5 mg·L⁻¹ plus an untreated check treatment with three replications for each treatment. The GC reading was recorded immediately followed by 2, 3, 4, 5, and 6 h after treatment commenced. Gas standards were prepared in an erlenmeyer flask (1 L) by removing the calculated amount of air and injecting a calculated amount of PH₃ into the erlenmeyer flask. All headspace samples were collected from the top and bottom ports in each of the fumigation chamber and injected in duplicate into the GC along with PH₃ standards.

**Fumigation of infested celery with the mixture of EF and PH₃**

To test the effect of combined EF and PH₃ on purple scum springtails, 10 bunches of celery were placed into two replication chambers similar to the previous (EF treatment) section. All experiments of the mixture of EF and PH₃ were repeated twice by using fresh celery bunches that were stored for less than 24 h at 15 °C cold room to confirm phytotoxicity only. The liquid EF was vaporized by using the gas sampling bag in hot water bath ≈90 °C for 10 min until the bags were inflated by EF vaporization. Phosphine of required concentration was stored in separate gas sampling bags. Both EF and PH₃ were injected into the chamber with EF as described previously. Three treatments of EF + PH₃ concentrations 20, 30, and 40 mg·L⁻¹ EF, with 1 mg·L⁻¹ PH₃, were used along with an untreated check treatment. All samples were taken from each (top and bottom) port of chamber immediately, 2 and 4 h later along with standards injected into the GC in duplicate.

**Mortality assessment of purple scum springtails**

The bioassay samples were retrieved at the end of the fumigation treatment by opening and aerating the fumigation chambers inside the fume hood for ≈30 min. The fumigated celery samples were divided into two lots from each drum. Half of the sample lot from each drum was used for counting purple scum

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Fig. 1. Fumigation chamber [61 L (16.1 gal)] used for celery fumigations.
springtail mortality. The second half of the sample lot from each drum was stored at 15 °C and 65% to 70% relative humidity for product quality and phytotoxicity studies.

After aeration, the mortality of purple scum springtails was evaluated by transferring five celery bunches from each replication to a white plastic tray for mortality assessment. This was carried out by dismantling all leaves of celery from external to internal leaves and carefully transferring all dead and live insects into a 9-cm glass petri dish to ensure that all purple scum springtails were recovered. A magnifying glass was used to determine dead and live purple scum springtails. Similarly, mortality assessment was conducted for untreated check treatments.

**Evaluation of fumigant phytotoxicity**

Phytotoxicity for EF, PH3, and their combination treatments was evaluated by cutting the treated and untreated celery bunches in half and looking for any damage or color change relative to untreated control samples. The observation of fumigant damages was taken every day from day 2 until day 7.

**Sensory and taste evaluation**

Sensory and taste evaluation were carried out by selected 20 volunteers (10 males and 10 females, between the ages of 30 and 60 years) to check and taste the treated and untreated celery. Before conducting sensory and taste tests, PH3 fumigated celery was aired for 2 h in under fume hood. The fumigated and untreated control celery samples were washed, leaves removed, and cut into pieces (15 cm long) for sensory and taste test. Treated and untreated celery were placed in two blank coded plates and the panel was asked to randomly pick the celery and rate their visual attributes related to color and taste related to flavor (characteristic flavor and odor), texture (crispness and juiciness), and smells from a scale of 1 to 9 (1 = dislike extremely to 9 = like extremely) (Barrett et al., 2010; Raffo et al., 2006; Yommi et al., 2013).

**Statistical analysis.** Mortality for all treatments was calculated with one-way analysis of variance (ANOVA) using SPSS Advanced Statistics software (version 24.0; IBM Corp., Armonk, NY) at the least significant difference level (5%) and Tukey’s 95% confidence intervals. The variation of all treatments and standards were assessed with duplicate injections and gas chromatography data. The average and sd of EF, PH3, and their combination absorption by celery were calculated on the basis of peak areas against the external gas standards of EF and PH3 using Microsoft Excel 2010 (Microsoft Co., Perth, Australia). The sensory evaluation results were treated by ANOVA at a significant level of 5%.

**Results**

**Sorption of fumigants by celery.** The concentration of EF and PH3 in the fumigation chambers was measured during 4 h of fumigation with EF and EF + PH3 and 6 h for PH3. The concentrations of fumigants are shown in Figs. 2–4. All concentrations of EF decreased rapidly within 0.5 h (Fig. 2) as EF was absorbed by the commodity. The concentration of EF in the 90 mg L⁻¹ treatment recorded a decline of 70% from the initial dose and more than 80% for the initial EF concentrations of 50 and 60 mg L⁻¹. After 2 h of fumigation, 85% to 90% of EF was absorbed by the commodity for all applied concentrations of EF. For the PH3 fumigation after 2 and 6 h, ≥40% to 50% and 60% to 70% of PH3 was absorbed by the commodity for applied concentrations of 1, 1.5, and 2 mg L⁻¹, respectively. However, for 2.5 mg L⁻¹ PH3, ≥22% and 32% of the initial applied concentration was lost over 2 and 6 h, respectively (Fig. 3).

For the EF + PH3 combination fumigation treatments, less than half of EF and ≥50% PH3 was lost within 2 h (Fig. 4). Generally, after 4 h of fumigation, ≥90% of EF and ≥40% of PH3 were lost from the headspace of fumigation chambers (Figs. 1–4).

**Bioassay of PH3 alone on purple scum springtails.** The total number of purple scum springtails tested varied in all treatments because of natural infestation. The numbers of pests ranged between 132 and 238 purple scum springtails per treatment. With celery fumigated in chambers with EF at the concentration of 50 mg L⁻¹, the mortality of purple scum springtails was 82.77%, 91.58%, and 93.28% for 1, 2, and 4 h exposure, respectively (Table 1). The second treatment of EF at a concentration of 60 mg L⁻¹ for 1, 2, and 4 h exposure showed observed mortality of 98.21%, 97.46%, and 96.04%, respectively. In the third treatment of EF, with a concentration of 90 mg L⁻¹, the observed mortality was 98.73% and 100% for 1 and 2 h exposure, respectively. These bioassay results show that there is a significant difference in the level of mortality between 1, 2, and 4 h exposure with EF alone compared with the untreated check. Also, there is a significant difference in the level of mortality among the concentrations 50, 60, and 90 mg L⁻¹ for all fumigation times. There was no significant difference in mortality between the concentrations of 60 and 90 mg L⁻¹ for any duration of fumigation. The high mortality of 100% was achieved only at 90 mg L⁻¹ after 2 h exposure.

The results of this study indicate that increasing EF concentration plays a more significant role than the duration of fumigation on mortality of purple scum springtails. There was a significant effect of concentration and time on purple scum springtail mortality (F = 5.773, P < 0.007). The relationship between concentration and time was assessed by variance analysis and accounted for $R^2 = 0.999$ and adjusted $R^2 = 0.998$.

**Bioassay of PH3 alone on purple scum springtails.** In the PH3-only treatment, varying numbers of purple scum springtails (80–217) were tested per treatment on purple scum springtail-infested celery. The relatively low and variable number of purple scum springtails per treatment is due to the celery being naturally infested.

Initially, two treatment of PH3 concentrations (1 and 1.5 mg L⁻¹) were tested (Table 2). The mortality achieved was 24.33%, 24.26%, and 26.40% at 1 mg L⁻¹ for 2, 4, and 6 h exposure time, respectively, and 37.20%, 20.90%, and 35.80% at 1.5 mg L⁻¹ for 2, 4, and 6 h exposure time, respectively. Other treatments of PH3 concentrations used two concentrations that were 2 and 2.5 mg L⁻¹ for 2, 4, and 6 h exposure. The results show mortality of 64.70%, 70.73%, and 73.23% at 2 mg L⁻¹ of PH3 and 53.86%, 73.63%, and 100% achieved at 2.5 mg L⁻¹ for 2, 4, and 6 h exposure, respectively. Mortality varied significantly by concentration and time of exposure.
exposure (F = 4.663, P < 0.001). The relationship between concentration and time variance accounted for $R^2 = 0.940$ and adjusted $R^2 = 0.912$.

**Bioassay of PH$_3$ plus EF in combination on purple scum springtails.** To reduce phytotoxicity on EF-treated celery, the dose of EF was decreased to 20, 30, and 40 mg L$^{-1}$ and combined with 1 mg L$^{-1}$ PH$_3$. The combination of PH$_3$ and EF was tested on purple scum springtail-infested celery with three concentrations of EF tested: 20, 30, and 40 mg L$^{-1}$ each combined with 1 mg L$^{-1}$ of PH$_3$. The number of target pests ranged between 101 and 211 insects per treatment.

The mortality of purple scum springtails was 100% for 2 and 4 h exposure at 40 mg L$^{-1}$ of EF mixed with 1 mg L$^{-1}$ PH$_3$ compared with the untreated check treatment. Mortality of 98.8% and 100% mortality was achieved at 30 mg L$^{-1}$ of EF mixed with 1 mg L$^{-1}$ PH$_3$ for 2 and 4 h exposure, respectively (Table 3). Mortality at 20 mg L$^{-1}$ of EF with 1 mg L$^{-1}$ of PH$_3$ was 89.5% and 95.5% for 2 and 4 h exposure, respectively. The interaction between concentration and time on purple scum springtail mortality was significant (F = 16.56, P < 0.001). The relationship between concentration and time was assessed by variance analysis and accounted for $R^2 = 1.000$ and adjusted $R^2 = 1.000$.

**Effect of fumigants on celery quality and phytotoxicity.** It was observed that EF fumigation can result in significant phytotoxicity of celery. Symptoms are changes in the color of leaves especially inside layers of young leaves, which turn brown. Our results indicate that all concentrations of EF used in this study affected celery and caused phytotoxicity. Damage was apparent in two ways: first, only young leaves were damaged at low EF concentrations, and second, both old and young leaves of celery were damaged at high concentrations of EF. Importantly, celery subjected to PH$_3$ treatments alone at various doses showed no evident phytotoxicity.

Damage to celery appeared within 1 d of treatment with 20 mg L$^{-1}$ EF plus 1 mg L$^{-1}$ PH$_3$. The browning of leaves appeared almost immediately at 30 and 40 mg L$^{-1}$ doses of EF. The quality of celery was reduced by wilting of outer foliage and yellowing of inside layered leaves by EF and the mixture of EF+PH$_3$ after 2–5 d of treatment depending on the concentration of fumigant.

The scores of sensory test for both treated and untreated celery ranged between seven and nine for all attributes (Table 4), corresponding to “like moderately” to “like extremely.” There were no statistically significantly different ($P >$
Discussion

Purple scum springtails are a common insect in Australia, occasionally found on fresh celery. This has led to export rejection to Middle East countries where it is a quarantine pest requiring fumigation. Our investigations showed that EF, PH₃, and their combination had a significant effect on purple scum springtail mortality. Experimental concentrations of fumigants EF, PH₃, and their combination were found to decrease significantly over time, which may be attributed to the absorption by celery bunches because of the high surface area and very high moisture content of the commodity (Figs. 1–4). The sorption of fumigants EF and PH₃ is known to depend on several conditions such as the type of commodity, target pest, temperature, moisture content, particle size and composition, exposure time, and fumigant concentration.

Table 1. The mortality of purple scum springtails after 1, 2, and 4 h exposure to ethyl formate (EF) at different concentrations in fumigation chambers containing celery.

| EF concn. (mg L⁻¹) | Time of exposure (h) | N | Mortality [mean ± SE (%)] | 95% CI (range) |
|-------------------|---------------------|---|---------------------------|---------------|
| 0                 | Untreated check     | 1 | 147                       | 0 ± 0 a        | 0             |
|                   |                     | 2 | 168                       | 0 ± 0 a        | 0             |
|                   |                     | 4 | 135                       | 0 ± 0 a        | 0             |
| 50                |                     | 1 | 132                       | 82.77 ± 1.17 b | 79.95–85.59   |
|                   |                     | 2 | 166                       | 91.58 ± 0.90 c | 89.42–93.74   |
|                   |                     | 4 | 164                       | 93.28 ± 0.10 cd | 93.03–93.53 |
| 60                |                     | 1 | 238                       | 98.21 ± 1.45 de | 94.71–101.71 |
|                   |                     | 2 | 230                       | 97.46 ± 2.07 cde | 92.47–102.44 |
|                   |                     | 4 | 156                       | 96.04 ± 1.87 cde | 91.55–100.53 |
| 90                |                     | 1 | 140                       | 98.73 ± 1.03 de | 96.25–101.22 |
|                   |                     | 2 | 193                       | 100.0 ± 0 c   | 100–100       |

LSD₀.05

\[1\ \text{mg} \cdot \text{L}^{-1} = 335 \text{ ppm.}\]
\[^{a}\text{Total number of purple scum springtails.}\]
\[^{b}95\% \text{ confidence interval (CI).}\]
\[^{c}\text{Mean mortality followed by the same letters in each group were not significantly different based on Tukey’s differences multiple range test, } P > 0.05.\]
\[^{d}\text{Least significant difference at } P = 0.05.\]
dose (Berck, 1968; Dhaliwal, 1975). Furthermore, Reddy et al. (2007) reported that PH3 concentration might vary according to food commodity types and found that the PH3 concentration in the headspace of the commodities varied from 0 to >2000 ppm. The higher moisture content of fresh products may be one of the contributing factors for sorption of PH3, as outlined in their studies where PH3 was applied to 74 products with high moisture content. This result is consistent with previous experiments that demonstrated that EF and PH3 can be significantly absorbed by the commodity or broken down to by-products because of the high moisture content of the commodity (Agarwal et al., 2015; Reddy et al., 2007). Furthermore, the toxicity to the pest can be increased with longer exposure time (Lee et al., 2015; Liu, 2011; Ren et al., 2011; Van Epenhuisen et al., 2007).

In the current study, mortality of purple scum springtails varied among different EF concentrations, with 100% mortality achieved at 90 mg L⁻¹ within 2 h. The results for EF indicate a significant increase in purple scum springtail mortality with increasing exposure to EF. This result is consistent with findings of other experiments which showed that vacuum fumigation of packaged lettuce with 0.5%, 1.0%, and 1.5% of EF for 2 h had little impact on the quality of lettuce while achieving high mortality of green peach aphid. Ethyl formate is approved for treatment of an export paprika (Capsicum annuum) market, as well as tomatoes (Solanum lycopersicum) in Korea (Kim et al., 2013). In addition, EF achieved high mortality against cotton aphids and two-spotted mites (Tetranychus urticae) (Kim et al., 2013; Lee et al., 2013; Stewart and Aharoni, 1983).

Table 1 shows the different percentage mortality of purple scum springtails with EF fumigation alone. A high dose of EF of 90 mg L⁻¹ achieved 100% mortality, compared with relatively low mortality at a concentration of 50 mg L⁻¹ of only 91.58% over the same exposure period. These results of high mortality on purple scum springtails may be attributable to fumigant penetration.
Table 4. Effect of fumigation on quality and sensory scores (1 to 9 hedonic scale) of celery.

| Quality attribute         | Treatment   | N | Score [mean ± SE (1 to 9)] | 95% CI (range) | P value |
|---------------------------|-------------|---|---------------------------|----------------|---------|
| Crispness                 | PH₃ treated | 20| 8.05 ± 0.05               | 7.95–8.15      | 0.574   |
|                           | Control     | 20| 8.00 ± 0.07               | 7.85–8.15      |         |
| Juiciness                 | PH₃ treated | 20| 7.50 ± 0.19               | 7.08–7.92      | 0.134   |
|                           | Control     | 20| 7.75 ± 0.14               | 7.45–8.05      |         |
| Flavor and taste          | PH₃ treated | 20| 7.85 ± 0.19               | 7.44–8.26      | 0.702   |
|                           | Control     | 20| 7.95 ± 0.17               | 7.59–8.31      |         |
| Color and appearance      | PH₃ treated | 20| 8.30 ± 0.16               | 7.96–8.64      | 0.661   |
|                           | Control     | 20| 8.20 ± 0.15               | 7.87–8.58      |         |
| Aroma and smell           | PH₃ treated | 20| 7.45 ± 0.56               | 6.91–7.99      | 0.387   |
|                           | Control     | 20| 7.75 ± 0.22               | 7.27–8.23      |         |

*Number of panelists who determined the sensory scores.

1 – dislike extremely and 9 – like extremely.

95% confidence interval (CI) for mean.

PH₃ = phosphine.

into each celery bunch. This result is consistent with results of the past studies that showed 100% mortality against Asian meal moth (Plodia interpunctella) and confused flour beetle by using EF alone on dried vine fruit and also high EF concentration caused high mortality on the eucalyptus weevil (Gonipterus platen sis) in export apples (Agarwal et al., 2015; Tarr et al., 2007).

Fumigation of celery bunches with PH₃ alone was studied, and 100% mortality was achieved at 2.5 mg L⁻¹ after 6 h fumigation. Lower doses of 1, 1.5, and 2 mg L⁻¹ also caused mortality on purple scum springtails, but levels of mortality were less (Table 2). These results are consistent with Liu (2009) who reported that PH₃ fumigant caused 100% mortality at 1.4 mg L⁻¹ after 24 h fumigation for western flower thrips on different vegetables such as lettuce, broccoli, and asparagus, which were successfully fumigated at low temperature (2 °C) without injury to these vegetables. For controlling lettuce aphids using PH₃ fumigation, 2.4 mg L⁻¹ of PH₃ achieved 100% mortality over 72 h fumigation at low temperature (2 °C) (Liu, 2009). Moon (2012) reported that concentrations of PH₃ should be more than 2 g m⁻³ for >24 h exposure for control of cotton aphid at 8 °C on cut flowers and nursery stock.

Current studies show high mortality over short fumigation periods, with the toxicity of EF plus PH₃ being much higher compared with PH₃ or EF alone. Low concentration of EF (30 and 40 mg L⁻¹) mixed with PH₃ (1 mg L⁻¹) at 2 and 4 h exposure gave 100% mortality compared with the high dose of EF or PH at the same exposure time when used alone (100% mortality achieved at 90 mg L⁻¹ EF alone for 2 h exposure and also with PH₃ at 2.5 mg L⁻¹ alone at 6 h).

Hence, we believe that there is a synergistic effect between EF and PH₃ for control of purple scum springtails and the combination produces a greater effect than either EF or PH₃ alone. At concentrations of 30 and 40 mg L⁻¹ of EF combined with 1 mg L⁻¹ PH₃, 100% mortality was achieved compared with 20 mg L⁻¹ of EF combined with the same amount of PH₃, which gave less mortality over the same exposure time. This result shown in Table 3 is consistent with Lee et al. (2015) who demonstrated a synergistic effect between EF and PH₃ for control of cotton aphids when applied in combination. They showed that the fumigant combination can cause high toxicity against all stages of citrus mealybug (Planococcus citri) and less damage to the commodities at low temperature (8 °C) for 4 h (Yang et al., 2016). There have been a number of research studies looking to reduce the fumigation period and increase PH₃ effectiveness with mixtures of different gases. For example, Liu (2008, 2011) found that exposure time of PH₃ can be reduced at low temperature and still control several postharvest pests, such as western flower thrips and grape mealybug (Pseudococcus maritimus), by using oxygenated PH₃.

All fumigants were effective against purple scum springtails in given short treatment times for exporting fresh produce with respect to disinfecting produce of live insects. However, EF fumigant alone and in combination with PH₃ affected the quality of celery and caused phytotoxicity symptoms. The observation of celery phytotoxicity from EF and the combination of EF and PH₃ demonstrated that high sorption of EF by the commodity is phytotoxic, even at low concentrations of EF or short fumigation times. Phytotoxicity of the fumigant to the commodities is very much dependent on the type of commodity treated, and the interaction of fumigant or its by-product with the chemical constituents of the commodities. These results are supported by Weller and Graver (1998) and Zhang and Van Epenhuijsen (2004) who reported that EF causes high phytotoxicity on cut flowers. In some previous studies, less damage to commodities was shown by using EF/PH₃ in combination achieving high mortality of target pests (Lee et al., 2015; Yang et al., 2016). Lee et al. (2014) indicated high mortality of insect pests by a combination of EF and PH₃ in naturally infested strawberries and cut flowers with less apparent phytotoxicity. Agarwal et al. (2015) have shown no apparent toxicity of EF to apple fumigation.

In the case of PH₃ fumigation alone, there was no difference in celery quality or phytotoxicity observed compared with untreated celery. This is consistent with the result obtained by Liu (2008). Previous studies have shown that PH₃ fumigation of vegetables is effective and safe for postharvest and commercial applications for insect control. Therefore, PH₃ remains one of the best fumigants for vegetable fumigation because it causes less phytotoxicity compared with other gases (Horn et al., 2005; Lee et al., 2012; Liu, 2008). Likewise, PH₃ treatments
alone at various doses achieved high mortality on purple scum springtails and maintained commodity at high quality. The sensory evaluation of fumigated celery bunches were similar to that of untreated control. There was no negative effect on quality parameters. Several studies indicated that PH₃ fumigation causes no significant effect on sensory evaluation in broccoli, tomato, and green pepper (C. annuum) (Ertürk et al., 2018; Liu, 2008).

Based on our results, treatments on purple scum springtails in celery, EF alone and in combination with PH₃, can control the target pest of purple scum springtails; however, the combination is phytotoxic to the commodity. By contrast, PH₃ alone achieved 100% kill of the target pest with no phytotoxic effect on the commodity.

Based on these findings, we conclude that PH₃ has excellent potential for pre-shipping treatment of export celery for control of purple scum springtails with little or no adverse effect on celery quality.

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