Changes in Gluconasturtiin Concentration in Chinese Cabbage with Increasing Cabbage Looper Density

Fernando A. De Villena
Department of Horticultural Science, University of Minnesota, 1970 Folwell Avenue, St. Paul, MN 55108

Vincent A. Fritz
Department of Horticultural Science, University of Minnesota, 1970 Folwell Avenue, St. Paul, MN 55108; and the Southern Research and Outreach Center, University of Minnesota, 35838 120th Street, Waseca, MN 56093

Jerry D. Cohen
Department of Horticultural Science, University of Minnesota, 1970 Folwell Avenue, St. Paul, MN 55108

William D. Hutchison
Department of Entomology, University of Minnesota, 1980 Folwell Avenue, St. Paul, MN 55108

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Abstract. Changes in the levels of gluconasturtiin (2-phenylethyl glucosinolate), an aromatic glucosinolate, was used to evaluate the response of ‘Green Rocket’ Chinese cabbage (Brassica campestris ssp. pekinensis L.) to the feeding of three and five cabbage looper (Trichoplusia ni Hübner) larvae per plant. Plants were harvested 0, 10, and 17 days after infestation. The change in gluconasturtiin concentration resulting from decreased light capture from damaged leaf area was also studied. All samples were assayed for gluconasturtiin concentration using high-performance liquid chromatography. The gluconasturtiin concentration of plants subjected to five larvae per plant showed a 59% increase 10 days after infestation compared with noninfested plants. Difference in gluconasturtiin concentration between three and five larvae per plant was nonsignificant. Seventeen days after initial infestation and 7 days after larvae were removed (final harvest), gluconasturtiin concentration did not decrease compared with the previous harvest. Reduced light or leaf area removal did not significantly affect gluconasturtiin concentration.

In recent years, the consumption of cruciferous vegetables with increased recognition of their nutraceutical properties has become very important because of the associated benefit of reducing the incidence of cancer and other chronic diseases (Talalay and Fahey, 2001; Wargovich, 2000). A group of secondary compounds, called glucosinolates, is widely present in crucifers. On cellular damage or disruption, hydrolysis of glucosinolates, through the enzyme myrosinase, produces isothiocyanates, which may, in part, account for the chemopreventive properties associated with these vegetables.

The hydrolysis of the aromatic glucosinolate gluconasturtiin releases phenethyl isothiocyanate (PEITC). Isothiocyanates are known to induce phase II detoxification enzymes (Wattenberg, 1990). Previous studies have shown that PEITC provides significant chemoprevention, especially against human prostate cancer (Powolny et al., 2003; Wargovich, 2000; Xiao et al., 2003). Gluconasturtiin is present in several crucifers such as watercress (Nasturtium officinale L.) and Chinese cabbage (Brassica campestris ssp. pekinensis L.). Because of the chemopreventive properties of PEITC, it is desirable to characterize gluconasturtiin (precursor of PEITC) concentration and explore the potential of plants to produce increased amounts of this compound.

The variation in total and individual glucosinolate concentration in a plant is the result of several factors, both abiotic and biotic (Rosa et al., 1997; Smith et al., 2003). Research has also shown that glucosinolate levels are not constant throughout the phenology of a plant, particularly in early growth stages. Young photosynthetically active tissues are believed to be the major sites of synthesis or storage of indole glucosinolates (Chong and Bible, 1974; Ju et al., 1980).

The cabbage looper (Trichoplusia ni Hübner) is a generalist insect native to the United States. It has been extensively used in controlled feeding trials and it was chosen for the present study because of its voracity, lower susceptibility to pesticides, and especially because cabbage plants have been shown to tolerate moderate defoliation before a significant reduction in plant weight occurs (Andaloro and Shelton, 1981). Changes in leaf nutrients incited by herbivory depend on the timing and intensity of feeding (Rostás et al., 2002). However, little has been done to correlate the concentration of glucosinolates with the level of insect pressure.

The possibility of manipulating the metabolism of cruciferous plants to modify its glucosinolate content has generated considerable interest. The potential impact this will have on both consumers and producers will depend on the plant’s ability to offer improved nutritional quality, less susceptibility to herbivores/pathogens, increased therapeutic properties, and desirable agronomic, storage, and sensory characteristics (Rosa et al., 1997).

Our objectives were to: 1) determine if there was an effect of cabbage looper larvae feeding on gluconasturtiin concentration in Chinese cabbage by correlating it with the insect pressure imposed on the plant; 2) evaluate the effect after the larvae were removed; and 3) determine if reduced light conditions had an effect on gluconasturtiin concentration.

Materials and Methods

Two greenhouse experiments were conducted to evaluate the effect of cabbage looper larval feeding on gluconasturtiin concentration in ‘Green Rocket’ Chinese cabbage plants. This cultivar was used to model the response of cruciferous vegetables to insect herbivory because of its relatively high gluconasturtiin concentration compared with other cultivars (V.A. Fritz, personal communication). A preliminary study was conducted in 2002 to determine the effect of larval feeding of two densities of cabbage looper on gluconasturtiin concentration. A second experiment was designed to evaluate the level of gluconasturtiin over time and was conducted twice in 2003. Greenhouse conditions were maintained constant throughout all experiments (20 ± 2 °C with natural photoperiod at 45° latitude).

Effect of cabbage looper density. On 15 Oct. 2002, two seeds of ‘Green Rocket’ Chinese cabbage (American Takii, Salinas, CA) were planted 2.5 cm deep in each cell of a 53 × 27-cm germination tray (Dillen Products, Middletfield, OH) filled with Metro-Mix 200 potting mix (60% horticultural vermiculite, 40% Canadian sphagnum peatmoss, horticultural perlite, and washed sand; Scotts Co., Marysville, OH) and watered as needed. On germination, plants were thinned to one per cell. After thinning, 100 mg L−1 of a
20N–8.7P–16.6K soluble fertilizer (JR Peters, Allentown, PA) was used to fertilize plants twice a week for the first 2 weeks. From the third week after thinning until the end of the experiment, plants received a weekly application of 200 mg L$^{-1}$ of the same 20N–8.7P–16.6K fertilizer. Three weeks after sowing, plants were transplanted to greenhouse benches, filled with a mixture of 1 sterilized sandy loam soil:3 SB 300 (55% bark, 45% horticultural vermiculite, Canadian sphagnum peatmoss, horticultural perlite, dolomite limestone, and gypsum; Sungro Horticulture Canada Ltd., Vancouver, BC) by volume.

The statistical model used was a split-plot design with four replicates. Half the plants in each replicate were covered [photosynthetic photon flux (PPF): 240 μmols m$^{-2}$ s$^{-1}$] with antiphid netting (Kaplan-Simon Co., Brain-tree, MA) to prevent unwanted herbivory. This netting would also prevent unwanted predation on the test insects. The remaining half of the plants in each replicate were not covered (PPF: 400 μmols m$^{-2}$ s$^{-1}$). The difference in whether the plants were covered with the netting comprised the main plots. No unwanted herbivory was observed in any of the control plants. Larval densities comprised the subplots and were arranged as follows: 1) nontreated plants (control), 2) plants infested with three larvae, and 3) plants infested with five larvae. Larval densities were chosen based on previous observations and the need to achieve significant levels of feeding injury while preventing total defoliation. These three larval density treatments were randomly assigned within each block. Each treatment was comprised of four plants per replicate located at each corner of a 38 × 23-cm rectangle, which ran perpendicular to the length of the block. Treatments were separated from each other by four nontreated plants arranged in a similar manner.

On arrival in the laboratory, cabbage looper eggs (USDA-ARS Western Cotton Research Laboratory, Phoenix, AZ) were placed on a plastic tray in a growth chamber (Charron and Sams, 2004) observed in rapid-orthogonal growth chambers, Chagrin Falls, OH). Once the eggs hatched, first-instar larvae were placed on the adaxial surface of the leaves of 4-week-old plants using a small, moist paint brush (length, 23 cm). One larva was placed per leaf so that each was feeding initially on a separate leaf. Larvae were left on the plant for a period of 10 d, after which two plants per treatment in each replicate were harvested and processed. Before processing, leaf area measurements were taken in plants from both treatments in each replicate located at each corner of a 76 × 56-cm rectangle with three plants equally distributed on the longer edge of the rectangle. Treatments were separated from each other by six nontreated plants arranged in a similar manner. Three sequential harvests were conducted and glucosinasturin concentration at each of them was determined to observe temporal distribution. The first harvest was performed on 12 Oct. 2003 before infestation and the second harvest was conducted 10 d after infestation. On this date, all larvae were manually removed from all plants. Finally, the third harvest was performed 17 d after infestation. At each harvest, two plants were harvested from each treatment per replicate, and the roots were discarded. Before processing, leaf area measurements were recorded as described. The experiment was repeated.

Protocol for analysis of glucosinolates. Samples from all experiments were processed and analyzed using the described procedures. First, the weight of Chinese cabbage plants was recorded before washing with deionized water. A 1:15 (w:v) ratio between Chinese cabbage fresh weight and deionized water was used. Water was brought to a boil in a 400-mL heat-resistant beaker covered with aluminum foil. Boiling was conducted to heat-deactivate the myrosinase enzyme. On boiling, the Chinese cabbage was placed into the beaker, making sure it was submerged under water. Once the water containing the Chinese cabbage returned to a boil, it was allowed to continue for 3 more minutes. The beaker was then removed from heat and maintained at room temperature for 10 min. The boiled cabbage was liquefied with water in a blender for 2 min. The total volume of the homogenate was recorded and 100 mL of the homogeneous suspension was transferred to a 120-mL plastic cup and frozen at −30 °C until analyzed for glucosinolate content.

One day before analysis, frozen samples were thawed at room temperature and centrifuged at 13,000 g $\times 10$ for 1 h (J2-21M/E; Beckman Instruments, Fullerton, CA). The supernatant was transferred to a 20-mL scintillation vial (03-337-2; Fisher Scientific Co., Pittsburgh, PA) and 5.6 g of ammonium sulfate was added per 10 mL. Finally, the vials were stirred until the salt was completely dissolved and placed in a cold room (5 °C) overnight to precipitate proteins that interfere with the high-performance liquid chromatography analysis. Before analysis, samples were retrieved from the cold room and filtered using 0.2-μm syringe filters containing a surfactant-free cellulose acetate membrane (190-2520; Fisher Scientific Co.).

To analyze for glucosinasturin concentration, an intact glucosinolate protocol was used (Lewke et al., 1996, modified by J.D. Cohen and L.Y. Wong, unpublished data). Samples were analyzed using an Ultra Aqueous C$_{18}$ analytical column (4.6 mm × 15 cm, 5 μm; Restek USA, Bellefonte, PA) at 30 °C with a flow rate of 1.0 mL min$^{-1}$. Solvent A was 0.1 M ammonium acetate and solvent B was 30% methanol with 0.1 M ammonium acetate. The elution protocol was 0 to 6 min, 100% A; 6 to 21 min, a linear gradient to 70% B; and 21 to 24 min, a linear gradient to 100% B. The solvent was then held at 100% B for 11 min. After the analytical sequence, a linear gradient to starting conditions (35 to 37 min, linear gradient to 100% A) followed by a 13-min postrun hold at 100% solvent A reconditioned the column for the next injection. Samples were analyzed in duplicate.

The external standards used to identify chromatogram peaks were prepared using a serial dilution starting from a stock of 30 μL of 1.0 μg mL$^{-1}$ of phenethyl glucosinolate potassium (glucosasturin) mixed with 120 μL of distilled deionized water (ddwater) and then diluted in half three times.

Statistical analysis. Data from all experiments were analyzed using SAS (version 8.2; SAS Institute, Cary, NC). Testing for significance of main effects and interactions on all variables was conducted using analysis of variance. Data were analyzed using the PROC GLM procedure and DUNCANS for mean comparison separation.

Results and Discussion

Effect of cabbage looper density. Glucosasturin concentration significantly increased as a result of insect herbivory. However, the differences in glucosasturin concentration between the two larval treatments were not significant, regardless whether the plants were covered with netting as shown in Figure 1 (noncovered PPF: 400 μmols m$^{-2}$ s$^{-1}$ versus covered PPF: 240 μmols m$^{-2}$ s$^{-1}$) (P ≤ 0.05). Rostás et al. (2002) found no effect of caging Chinese cabbage leaves with and without Phaedon cockerelli beetles on glucosinolate concentration. However, this is in contrast to what Charron and Sams (2004) observed in rapid-cycling Brassica oleracea in which total glucosinolates decreased when PPF was...
increased from 200 to 400 \( \mu \text{mols m}^{-2}\text{s}^{-1} \). Because the difference in gluconasturtiin concentration between covered and noncovered plants was nonsignificant (Fig. 1), there was no significant influence of reduced light on gluconasturtiin production \((P \leq 0.05)\). However, it may be that differences in light levels were not large enough to influence gluconasturtiin concentration by affecting the metabolism of either the plant or insect. An analysis of variance between the amount of leaf area removed by herbivory and the related gluconasturtiin concentration in remaining leaf tissue was conducted. Although a higher population of insects per plant resulted in lower leaf area, differences in gluconasturtiin were nonsignificant (data not shown). A regression analysis found a weak correlation between gluconasturtiin and leaf area removed \((y = 254.2 - 0.14x; r^2 = 0.11; P \leq 0.05)\), so leaf area loss alone does not appear to influence gluconasturtiin concentration in Chinese cabbage. Cipollini et al. (2003) also found no correlation between leaf area removed and total glucosinolate concentration in rape \( (Brassica napus) \) plants. This suggests that the effect of larval feeding (chewing) may be more important than any amount of leaf area removed. It is more likely that Chinese cabbage plants increased their gluconasturtiin concentration on experiencing damage incited by herbivory of the cabbage looper and not necessarily as a result of a reduction in leaf area (W.D. Hutchison, personal communication). Nielsen et al. (2001) suggested that \( Arabidopsis \) lines were induced to increase their sinapin and total glucosinolate concentration regardless of the herbivore density of the specialist beetles \( Phyllotreta nemorum \) and \( P. cruciferae \). Larval feeding could have caused a translocation of gluconasturtiin from wounded to nonwounded areas as suggested previously (Agrell et al., 2003; Bodnaryk, 1992). It is also possible that there is a more complex plant:insect interaction that may have accounted for the increase of gluconasturtiin. Kessler and Baldwin (2002) found that lytic enzymes in the oral secretions of \( Pieris brassicaceae \) larvae acted as elicitors of terpenoid volatiles from cabbage leaves, although there are no published data on a similar relationship between the saliva of cabbage looper and glucosinolates in crucifers.

Overall, we concluded that indeed, there was a significant increase of gluconasturtiin attributable to larval feeding compared with the noninfested controls, but no significant difference was noted between three and five larvae per plant.

**Change in gluconasturtiin concentration after herbivory.** The experiment was conducted twice in 2003. Ten days after cabbage looper feeding, gluconasturtiin concentration in Chinese cabbage significantly increased with five larvae per plant compared with control plants in both covered and noncovered plants as shown in Figure 2. There was no significant decrease in gluconasturtiin in the noncovered plants in both larval treatments at the last harvest \((17 \text{ d after infestation})\) compared with the second harvest \((10 \text{ d after infestation})\). Once the stressor (larvae) was removed, gluconasturtiin levels were expected to return to normal, as suggested by Renwick (2002). It was also expected that gluconasturtiin levels would decline with plant age (Mewis et al., 2002). Agrell et al. (2003) has suggested that plants respond to injury within hours or days with relatively short lasting effects. However, this was not observed in our studies. Once gluconasturtiin increased after the initial herbivory treatment, concentrations remained elevated 7 \text{ d after larval removal} as shown in Figure 2. This may indicate that the increased gluconasturtiin concentration in plants subjected to herbivory results from accumulation of gluconasturtiin into a smaller pool of leaf tissue. This might suggest that there is a concentration effect and that the plant is compensating for tissue loss by increasing gluconasturtiin production per unit of leaf tissue.

To better evaluate the effect of larval feeding, we conducted a Duncan’s multiple

![Fig. 2. Effect of immediate and postherbivory (17 d after infestation) cabbage looper larvae feeding on gluconasturtiin concentration in ‘Green Rocket’ Chinese cabbage in 2003 \((P \leq 0.05)\).](image1)

![Fig. 3. Effect of immediate and postherbivory (17 d after infestation) cabbage looper larvae feeding on gluconasturtiin concentration in ‘Green Rocket’ Chinese cabbage, averaged over covering treatment, in 2003 \((P \leq 0.05)\).](image2)
range mean separation test averaged over the covering (netting) treatments as shown in Figure 3. We confirmed that the treatment using five larvae per plant induced the plants to produce significantly more gluconasturtiin than the noninfested control plants \((P \leq 0.05)\) at the second and third harvests, but we found no significance between the two larval densities used.

These results confirmed that there was a significant increase in gluconasturtiin concentration when Chinese cabbage plants were damaged by cabbage looper feeding. It may be possible that the increase is because this aromatic glucosinolate is part of a plant defense mechanism against herbivory. Rostás et al. (2002) suggested that glucosinolates act against generalists (insects that feed on several vegetable species) and not against insects that feed primarily on crucifers (specialists). The cabbage looper might be considered a generalist, because it feeds on different species of vegetables. It may also be able to rapidly detoxify the compound or be stimulated to feed more aggressively by the increase in gluconasturtiin concentration. This action could force the plant into a rapid cycle of greater production of the compound, which was observed with five larvae per plant (Agrawal, 2000). It has also been reported that the potential repellent effect of gluconasturtiin is more effective in cabbage looper adults where they have been deterred from oviposition by high concentrations of glucosinolates (Kushad et al., 2004; Mewis et al., 2002).

Because nondamaged organs of the plant elevate their glucosinolate concentration when other organs within the same plant are damaged (Agrell et al., 2003; Bodnaryk, 1992), we analyzed changes in gluconasturtiin concentration in whole plants. This is important especially for potential marketing of specialty vegetables with enhanced health benefits. The damaged parts of a plant could be removed to improve the plant appearance while still benefiting from increased levels of glucosinolates.

Under our experimental conditions, leaf area and reduced light (noncovered, \(PPF: 400 \mu\text{mol m}^{-2}\text{s}^{-1}\) versus covered, \(PPF: 240 \mu\text{mol m}^{-2}\text{s}^{-1}\)) conditions did not significantly affect gluconasturtiin concentration.

Public acceptance of vegetables with improved health benefits will depend on a large degree on a positive sensory experience. Elevated levels of glucosinolates in cruciferous vegetables are often associated with bitterness or pungency and could limit widespread embrace by the consumer. As fruits and vegetables are developed with the aim of increasing concentrations of naturally produced compounds with strong chemopreventive properties, management of compounds associated with a positive sensory reaction by consumers will be critical.

### Literature Cited


