Chilling injury alleviation and quality maintenance of lemon basil by preharvest salicylic acid treatment

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ABSTRACT

The purpose of this study was to alleviate chilling injury (CI) of lemon basil during storage by using preharvest salicylic acid (SA) treatment. The basil was sprayed with SA at the concentration of 0, 1, 5 or 10 mM before harvesting 24 h and was then stored at 7 °C for 4 days. Weight loss, CI score, malondialdehyde (MDA) content, electrolyte leakage (EL), superficial colour, pigments and bioactive compounds were determined. SA treatments alleviated CI and reduced the increase in EL and MDA content which the best result alleviating CI was 5 mM SA. The greenness and total chlorophyll content were maintained by SA application. After 24 h of treatment, antioxidants, total phenols and flavonoids were enhanced. During storage, SA treatment reduced the loss of bioactive compounds in the lemon basil. In conclusion, preharvest SA treatment at 5 mM alleviated CI and maintained quality of lemon basil during cold storage.

Keywords: Lemon basil; Salicylic acid; Preharvest treatment; Chilling injury; Bioactive compounds

INTRODUCTION

Basils (Ocimum sp.), one of most popular culinary herb, are widely used as a flavoring agent in Asian cuisine, especially Thai food. In Thailand, there are three commercial basils; lemon basil (Ocimum x citriodourum), sweet basil (Ocimumbasilicum) and holy basil (Ocimum sanctum) which are widely consumed in country and exported to European and other Western countries in form of cut herb. Recently, these basils have commercially grown in European countries during summer period and the consumption demand has also increased as its favorable aroma and medicinal properties. As having medicinal properties, basils are a rich sources of biologically active compounds involving phenolic compounds, flavonoids and antioxidants (Hakkim et al., 2007). Commercially, fresh fruit and vegetables are held at cold temperature during transportation, but low temperature in a mixed-load shipment cause problems for chilling-sensitive produces, especially tropical leafy herbs (Lange and Cameron, 1997). Among the three commercial basils, lemon basil is relatively high sensitive to storage at low temperature which resulted in leaf blackening which limiting its quality and shelf-life (Wongsheere et al., 2009). The chilling injury (CI) of basil is associated with the dysfunction of oil gland in leaf which the browning spots were observed and then expanded through CI development (Pongprasert and Srilaong, 2007). At 4°C, the visible blackening spots in mature lemon basil leaves is found in 12 h and then rapidly increase hereafter whilst at 12°C the blackening spots appeared in 5 days (Wongsheere et al., 2009). In other basils, the onset of blackening spots of sweet basil held at 4-5°C is 1 day (Lange and Cameron, 1997; Pongprasert and Srilaong, 2007) and that of holy basil held the 5°C noticeably appears in 3 days of storage (Niamthong et al., 2007).

It is widely recognized that CI is accompanied by dysfunction of cell membrane due to lipid peroxidation and oxidative stress. In CI temperatures, elevated levels of reactive oxygen species (ROS), particularly H2O2, have been detected (Elwan and El-Hamahmy, 2009). Generally, there is a balance between the generation of ROS and ROS scavenging system in plant. The ROS can be minimized and removed by antioxidant defense mechanisms (Gülçin et al., 2007). Recently, salicylic acid, a plant growth regulator and a safe chemical used to maintain postharvest quality, has been proven to be a major component inducing bioactive compounds such as antioxidants and...
antioxidant enzymes (Supapvanich and Promyou, 2013). Elwan and El-Hamahmy (2009) suggested that a range of antioxidant enzyme activities such as catalase (CAT), guaiacol peroxidase (G-POD) and ascorbate peroxidase (A-POD) play important roles in removing $H_2O_2$ which these enzyme activities are induced by SA. Hatamzadeh et al. (2012) suggested that SA is able to eliminate ROS and lead to decreasing lipid peroxidation. Not only does bioactive compounds induced by SA stimulating tolerance stress system in plant but these compounds also provide health benefit to human beings. However, no information has been reported on the preharvest SA application on postharvest quality and bioactive compounds of lemon basil. Thus the purpose of this study was to investigate the effect of preharvest SA application on quality maintenance and bioactive compound levels of lemon basil during cold storage.

**MATERIALS AND METHODS**

**Plant materials preparation and treatments**

Lemon basil (*Ocimum × citriodourum*) was cultivated at an agricultural demonstration plot, Faculty of Agricultural Engineering, King Mongkut’s Institute of Technology Ladkrabang, in May 2014. After 2 months of cultivation, they were sprayed with SA solution at the concentration of 0, 1, 5 or 10 mM. The lemon basils were harvested after SA spraying 24 h in morning (6:30-7:30 am) by using sharp clipper. The vegetables were then delivered to the laboratory at Department of Agricultural Education within 15 min after harvested. The vegetables were screened being free of any physical damages or disease. The vegetable leaves were cut in 15 cm length. The vegetables were cleaned by rinsing with tap water and then immersed in 50 $\mu$L $L^{-1}$ sodium hypochloride solution for 2 min. The excess surface water was removed by using a shaking basket. Twenty grams of the vegetables were contained in a laminated polyethylene plastic bag, size $22.9 \times 10.2$ cm and then the bag was sealed. The vegetable was stored at 7 ± 1 °C and 85 ± 2% RH for 4 days.

**Weight loss and superficial colour measurement**

The loss of fresh weight was monitored before and during storage. The lemon basils were weighed on initial day and then the bag was sealed. The vegetables were then delivered to the laboratory at Department of Agricultural Education within 15 min after harvested. The vegetables were screened being free of any physical damages or disease. The vegetable leaves were cut in 15 cm length. The vegetables were cleaned by rinsing with tap water and then immersed in 50 $\mu$L $L^{-1}$ sodium hypochloride solution for 2 min. The excess surface water was removed by using a shaking basket. Twenty grams of the vegetables were contained in a laminated polyethylene plastic bag, size $22.9 \times 10.2$ cm and then the bag was sealed. The vegetable was stored at 7 ± 1 °C and 85 ± 2% RH for 4 days.

**Chilling injury score measurement**

Chilling injury (CI) of lemon basil during storage was estimated using the score of CI symptoms (browning fleck) on the leaves. The CI severity was rated on a relative scale of 1 to 5 where: 1 = no CI symptom, 3 = moderate and 5 = severe (Supapvanich et al., 2012).

**Electrolyte leakage measurement**

The electrolyte leakage of lemon basil leaves was carried out using method describe by Promyou et al. (2012) with slight modification. Ten leaves of the lemon basil leaves were half cut using a sharp blade. The weight of cut lemon basil leaves was recorded. The cut leaves were rinsed with de-ionized water twice and dried using a Whatman No. 1 filter paper. The cut leaves were incubated in 30 mL of de-ionized water at ambient temperature ($28±1$ °C) and shaken for 60 min. The conductivity of the solution was immediately measured at 0 min and again at 60 min using a conductivity tester (DIST3, Hunna Instruments Inc., Romania). The sample was then frozen for 24 h and thawed. After that, the sample was boiled for 10 min. The conductivity (total conductivity) was again measured after the solution temperature drop to ambient temperature. The percentage of conductivity increase per 1 h was calculated which compared to the total conductivity.

**Malondialdehyde (MDA) content assay**

A sample of 5 g of lemon basil leaves was homogenized in 20 mL of 5 % trichloroacetic acid (TCA) and then centrifuged at 10,000×g for 20 min. The supernatant was used to assay MDA content using method described by Heath and Packer (1968) with slight modification. The reaction began when 1mL of supernatant was mixed with 2mL of 15 % TCA containing 0.5 % thiobarbituric acid (TBA). The mixture was heated at 60 °C for 30 min and immediately cooled using an ice bath for 30 min. The absorbance of the mixture was then measured at 532 and 600 nm, respectively. The MDA content was calculated using an extinction coefficient of $1.55$ mM cm$^{-1}$ as follows: MDA content = [(OD$_{532}$ - OD$_{600}$) × 2 mL × (total volume of extract (mL) × 1 mL)]/(1.55 × $10^{-3}$ × 5 g). The data were represented as nanomoles of MDA per gram fresh weight (nmolg$^{-1}$ FW).

**Chlorophylls and carotenoid contents measurements**

Chlorophylls and carotenoid contents were assayed according to the method of Kirk (1968) with slight modification. Three gram of lemon basil was homogenized with 10 mL of absolute ethanol. The sample was filtered using Whatman No. 1 filter paper and rinsed with 10 mL of absolute ethanol. The filtrate was collected and then adjusted to 50 mL using absolute ethanol. The absorbances at 654, 663 and 470 nm were measured using a Hekios UV–visible spectrophotometer (Thermo Spectronic, Cambridge, UK). Chlorophyll $a$ content was calculated as $(11.75 \times OD_{649}) - (2.35 \times OD_{663})$, chlorophyll $b$ content was calculated as $(18.61 \times OD_{649}) - (3.96 \times OD_{663})$.  

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The sum of chlorophyll \( a \) and chlorophyll \( b \) was expressed as total chlorophylls content. The data was represented as \( \mu g \) total chlorophylls content per 100 g fresh weight. Total carotene was calculated as \((1000 \times OD_{470}) - (2.27 \times \text{Chlorophyll } a) - (81.4 \times \text{chlorophyll } b)/227 \). The data was represented as \( \mu g \) carotene per 100 g fresh weight.

**FRAP value, \( H_2O_2 \) scavenging activity assays**

Antioxidant capacity was determined using ferric reducing antioxidant potential (FRAP) methods, which described by Benzie and Strain (1996). Three grams of the lemon basil leaves was homogenized with 5 mL of 80% (v/v) methanol and then 25 mL of distilled water was added. The suspension was stirred at 4 °C for 30 min and then filtered using Whatman No.1 filter paper. The filtrate was collected to determine FRAP value, \( H_2O_2 \) scavenging activity, total phenols (TP) and total flavonoids (TF) content. For FRAP assay, 0.1 mL of the extract was added into 2.9 mL FRAP reagent, consisting of acetate buffer pH 3, 10 mM 2,4,6-tripyridyl-1,3,5-triazine (TPTZ) and 20 mM ferric chloride hexahydrate in the ratio of 10:1:1. The reaction was held at ambient temperature for 30 min before measuring absorbance at 560 nm. FRAP value was expressed as \( \mu g \) Trolox equivalents per g fresh weight. \( H_2O_2 \) scavenging activity was determined using titration method described by Zhang (2000). One mL of the extract was mixed with 0.1 mL of 0.1 mM \( H_2O_2 \), followed by 2 drops of 3% ammonium molybdate, 10 mL of 2M \( H_2SO_4 \) and 7.0 mL of 1.8M KI. The mixture solution was titrated with 5 mM \( NaS_2O_3 \) until colourless. The percentage of \( H_2O_2 \) scavenging was calculated as \( [(V_o-V_f)/V_o] \times 100 \); \( V_0 \) = volume of 5 mM \( NaS_2O_3 \) used to titrate the control in present of \( H_2O_2 \) (without extract) and \( V_f \) = the volume of 5 mM \( NaS_2O_3 \) used to titrate in the present of extract.

**Total phenolic and total flavonoids content assays**

TP content was determined using the method described by Slinkard and Singleton (1977). The reaction of TP content was started when 0.1 mL of the extract was added into the solution of 1.9 mL 50% (v/v) Folin–Giolacelte reagent solution and 2 mL saturated \( Na_2CO_3 \) solution. The mixture was left at ambient temperature for 30 min. The absorbance at 750 nm was recorded. TP content was expressed in term of mg gallic acid per g fresh weight. TF content was determined using method described by Jia et al. (1999). The reaction was started when 0.5 mL of the extract was mixed with 1.5 mL of distilled water, 75 \( \mu L \) of 0.5% \( NaNO_2 \). The mixture was left for 6 min and then 150 \( \mu L \) of 10% \( AlCl_3 \cdot 6H_2O \) was added and allowed to stand for 5 min. After that, 0.5 mL of 1 M NaOH was added. The absorbance at 510 nm was recorded. The data were expressed in term of mg catechin equivalents per g fresh weight.

**Catalase, guaiacol peroxidase and ascorbate peroxidase activities assays**

Three grams of lemon basil was homogenized with 20 mL of 0.1 M phosphate buffer (pH 7) containing 0.2 g of polyvinylpolypyrrolidone (PVPP). The homogenate was filtered using Whatman No.1 filter paper. The filtrate was held in an ice bath. The assay of catalase (CAT) (EC 1.11.1.16) was determined using titrimetric method of Kar and Mishra (1976) and guaiacol peroxidases (G-P) (EC 1.11.1.7) and ascorbate peroxidase (A-P) (EC 1.11.1.11) activities were determined using method of Andrade Cuvi et al. (2011). Five mL of the mixture for CAT assay consisted of 3 mL of phosphate buffer (pH 7.0), 1 mL of 0.01 mM \( H_2O_2 \) and 1mL of enzyme extract. After incubation for 1 min at ambient temperature (27±1°C), the reaction was stopped by adding 10 mL of 2% \( H_2SO_4 \). The residual \( H_2O_2 \) was titrated against 0.01 M KMnO \(_4\) until a faint purple colour persisted for at least 20 sec. A control was the same mixture but the enzyme activity was stopped at 0 min. Unit of CAT is defined as that amount of enzyme which break down 1 \( \mu mol \) of \( H_2O_2 \)/min. The reaction mixture of G-P contained 0.6 mL of 0.5% (v/v) guaiacol, 0.5 mL of enzyme extract and 1.6 mL of phosphate buffer (pH 7.0). The reaction started when 0.3 mL of 40 mM \( H_2O_2 \) was added. The G-P activity was determined by monitoring the increase in the absorbance at 470 nm. The unit of the enzyme activity was expressed as the \( \Delta OD \) per min (unit) per g fresh weight. The mixture of A-P consisted of 0.5 mL of 5 mM ascorbic acid, 0.5 mL of enzyme extract and 1.5 mL of phosphate buffer (pH 7.0) and the reaction began when 0.5 mL of \( H_2O_2 \) was added. The absorbance at 290 nm was expressed as the amount of enzyme that oxidized 0.01 mmole ascorbate per minute (unit) per g fresh weight. The activity of CAT, G-P and A-P were expressed as unitg\(^{-1}\) FW.

**Statistical analysis**

The data are shown as the mean of four replications and S.D. bar. Data were analyzed using analysis of variance (ANOVA) which was performed with the SPSS software. The means of data were compared by the least significant difference (LSD) test at a significance level of 0.05.

**RESULTS AND DISCUSSION**

**Weight loss and chilling injury**

As the result shown in Fig. 1A, preharvest SA treatments had no effect on the loss of fresh weight during cold storage. The weight loss of lemon basil were approximately 3.18% and 4.22% on day 2 and 4, respectively (Fig. 1A). The main factor affecting the loss of fresh weight of the lemon basil during storage could be relative humidity as describe by Supapvanich et al. (2012). The CI symptom of
lemon basil was generally characterized by leaf blackening and brown spot on leaves which due to the dysfunction of oil glands (Wongsheree et al., 2009; Pongorsert and Srilaong, 2007). As expected, the CI score of lemon basil increased during stored at 7°C. The onset of CI symptom appeared on day 2 and SA treatments were able to alleviate CI of the basil leaves (Fig. 1B). We found that 5 mM SA application was the most effective in alleviating CI in the basil when compared to other concentrations. At the end of storage (day 4), the CI score of 5 mM SA-treated lemon basil was significantly lower than that of others ($P \leq 0.05$) which it was lower than 2.5 (acceptable score) whilst the CI score of the control was about 4 (Fig. 1B). This may due to the induction of defense mechanism system by SA (Supapvanich and Promyou, 2013) which the cell membranestructure of oil glands in the basil leaves was also maintained. However, the use of SA at high concentration (10 mM) caused tip-burn symptom on the shoot and young leaves (data not shown). In the similar vein, Babalar et al. (2007) and Elwan and El-Hamahmy (2009) had explained that the use of SA at a high concentration may harm the produce. This work shows that preharvest SA application at 5 mM was a proper concentration controlling CI of lemon basil during cold storage. Therefore, the basil treated with 5 mM SA was selected to investigate physicochemical factors relating to chilling injury.

**Electrolyte leakage and MDA content**

Increase in EL and MDA content are commonly accepted as indicators of CI development during refrigerated storage of fresh commodities which due to the cell membrane dysfunction (McCollum and McDonald, 1991; Luo et al., 2011). As shown in Fig 2, the EL of the basil leaves increased throughout storage and that of the control was significantly higher than that of the SA-treated basil ($P \leq 0.05$). This showed that preharvest SA application at 5 mM was effective in preventing the membrane dysfunction due to CI alleviation. Similarly, Lukaszewska and Kobyliński (2009) reported that SA application increased longevity of *Hipperstrum x chmielii* leaves by protecting membrane integrity and reducing the increase in electroconductivity of cell sap. The MDA content of both SA-treated and untreated lemon basil also increased following storage period (Fig. 2B). The increase in MDA content was related to lipid peroxidation of cell membrane which was concomitant with the increase in EL and CI during storage (McCollum and McDonald, 1991). The SA application reduced the increase in MDA content of lemon basil during storage. These show that preharvest SA application was able to alleviate CI incidence in lemon basil by maintaining cell membrane integrity and also reducing lipid peroxidation during cold exposure. In the similar vein, SA treatments delayed lipid

![Fig 1](image1.png)

**Fig 1.** Weight loss (%) (A) and chilling injury (CI) score of SA preharvest-treated lemon basil at various SA concentration during storage at 7±1 °C for 4 days. Data represent the mean of four replications ±S.D.

![Fig 2](image2.png)

**Fig 2.** Electrolyte leakage (%) (A) and malonaldehyde (MDA) content of lemon basil preharvest-treated with 0 and 5 mM SA during storage at 7±1 °C for 4 days. Data represent the mean of four replications ±S.D.
peroxidation and membrane dysfunction in pomegranate fruit (Awad et al., 2013) and plum fruit (Luo et al., 2011) during cold storage.

**Superficial colour and pigments**

Superficial colour is generally accepted as a main factor affecting the quality of leafy vegetables, especially greenness. As the results shown in Fig 3, the SA treatment maintained the greenness and chroma value of lemon basil during cold storage whereas those of the control significantly decreased ($P < 0.05$). No significant difference in brightness and hue value of both SA-treated and untreated lemon basil leaves were found over storage. Similarly, Wei et al. (2011) reported that SA treatment could maintained superficial colour of asparagus spears, due to delaying the deterioration in green colour. The changes in chlorophylls and total carotenoids content are generally related to the superficial colour. The loss of total chlorophyll content is concommitant with the loss of greenness and the occurance of yellowness of leafy vegetables (Supapvanich et al., 2012). In this study, the SA application maintained total chlorophyll content during storage whilst that of the untreated lemon basil was significantly decreased during storage ($P < 0.05$) (Fig 4). Similarly, chlorophyll content and greenness of asparagus was maintained by SA use (Wei et al., 2011). The reduced chlorphyll content in the control was concomitant with the increase in total cartenoids content as shown in Fig 4B. The total carotenoids content of SA-treated basil was maintained. These suggest that preharvest SA treatment was able to maintain superficial colour of lemon basil by retarding chlorophyll loss. These might involve with the increase in photosynthesis rate and the number of chloroplasts by SA which described by Radwan et al. (2008). Moreover, Li et al. (1992) and Kazemi et al. (2011) described that the inhibition of chlorophyll loss by SA application might due to the suppression of ACC synthase and ACC oxidase activities.

![Fig 3](image-url) **Fig 3.** Superficial colour, brightness (A), greenness (B), chroma (C) and hue value (D) of lemon basil preharvest-treated with 0 and 5 mM SA during storage at 7±1°C for 4 days. Data represent the mean of four replications ±S.D.

![Fig 4](image-url) **Fig 4.** Pigments content, total chlorophylls (A) and total carotenoids (B), of lemon basil preharvest-treated with 0 and 5 mM SA during storage at 7±1°C for 4 days. Data represent the mean of four replications ± S.D.
Total phenols, total flavonoids, FRAP value and $H_2O_2$ scavenging activity

Antioxidant activities involving FRAP value and $H_2O_2$ scavenging activity, TP and TF content which known as the biologically active compounds benefiting human health were commonly investigated in fresh commodities. After preharvest SA spray for 24 h, TP, TF and FRAP value of SA-treated lemon basil were significantly higher than those of the control ($P<0.05$); whereas, $H_2O_2$ scavenging activity of both treatments was similar (Fig. 5). TP, TF and FRAP value of both SA-treated and untreated lemon basil decreased over storage. No significant difference in TP and TF content were found at the end of storage but FRAP value of SA-treated lemon basil was higher than that of the control. We also found that $H_2O_2$ scavenging activity of SA-treated lemon basil was maintained whilst that of the control significantly decreased. Recent studies on fruit and vegetables have shown that their antioxidant efficiency may be contributed to a range of bioactive compounds such as phenolics and flavonoids content (Rapisarda et al., 1999; Wei et al., 2011) and antioxidant enzymes (Supapvanich et al., 2012). SA actively induced or maintained FRAP and free radical scavenging activity as reported for asparagus (Wei et al., 2011). In the similar vien, FRAP value, TP and TF content in SA-treated rambutan fruit (Supapvanich 2015) and peach fruit (Razavi et al., 2014) were higher than those of the untreated fruits. These confirm that SA induces biologically active compound in fresh produce which due to the stimulation of defense mechanism as described by Supapvanich and Promyou (2013). The higher bioactive compound of SA-treated lemon basil after preharvest spray 24 h might induce the tolerance to chilling temperature during storage.

CAT, G-POD, A-POD activities

CAT, G-POD and A-POD are known as the antioxidant enzymes playing important roles in removing $H_2O_2$ which recognised as a ROS (Elwan and El-Hamahmy, 2009). As shown in Table 1, CAT activity of the lemon basil was induced after SA spray for 24 h; whereas, both G-POD and A-POD activities of SA-treated lemon basil did not different from the control. During storage, all antioxidant enzyme activities of the control decreased considerably whilst those of SA-treated lemon basil remained constant. The stimulation of G-POD and A-POD activities by SA plays a key roles in preventing the increase of $H_2O_2$ in plant tissues (Kang et al., 2003; Shi et al., 2006). The high level of these antioxidant enzyme activities in the SA-treated lemon basil on day 4 were positively related to the level of $H_2O_2$ scavenging activity as shown in Fig. 5D. These were also concomitant with the CI suppression and the maintained membrane integrity as shown in Fig 1 and 2.

CONCLUSION

In conclusion, preharvest SA treatment was effectively able to delay CI symptom of the lemon basil during refrigerated storage at 7°C. Preharvest SA treatment at 5 mM showed the best results suppressing CI and maintaining superficial colour and total chlorophylls content. Regarding to CI alleviation, preharvest SA treatment retarded the increase in EL and MDA content. The higher bioactive compounds of SA-treated lemon basil played an important role maintaining postharvest quality and alleviating CI symptom during refrigerated storage.

Fig 5. Total phenolics (TP) content (A), total flavonoids (TF) content (B), FRAP value (C) and $H_2O_2$ scavenging activity (D) of lemon basil preharvest-treated with 0 and 5 mM SA during storage at 7±1°C for 4 days. Data represent the mean of four replications ±S.D.
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Author contributions

S. Supapvanich was the project director who planned the overall experiments, interpreted and analyzed data and wrote this article. R. Phnopakdee supervised basil cultivation and reviewed the article. P. Wongsuwan collected raw data and measured certain physicochemical factors.

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