Lettucenin A and Its Role against *Xanthomonas Campestris*

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Abstract
Lettucenin A is the major phytoalexin produced in lettuce after being elicited by biotic or abiotic elicitors. The production of lettucenin A in leaf can be induced by 5% of CuSO4 and 1% of AgNO3. A clear inhibition zone where the fungi *Aspergillus niger* failed to develop on TLC plates dipped in hexane: ethyl acetate (1:1, v/v) at Rf 0.45 was observed. Lettucenin A was detected at a retention time of approximately 5.3 min after being injected into the HPLC run with isocratic solvent system containing water: acetonitrile ratio 60:40, (v/v). In vitro antibacterial study with *Xanthomonas campestris* results showed this pathogen has different sensitivity to all tested concentrations of lettucenin A. The bacteria was more sensitive to higher concentration of lettucenin A (333, 533 and 667 µg ml⁻¹), compare to lower concentrations such as 67 µg ml⁻¹. Thus, the relationship between the bacteria growth rate and lettucenin A concentration was negatively correlated. However, the bacteria growth rate continues to increase after two hours of incubation. Hence, it is suggested that *X. campestris* may have the ability to detoxify lettucenin A. The success or failure of *X. campestris* to invade lettuce may very well depend on the balance between accumulation and degradation of lettucenin A at the invading sites of lettuce. In summary, lettucenin A may play an important role in the resistance of lettuce against microbial colonization.

Keywords: *Lactuca sativa*, Lettucenin A, *Xanthomonas campestris*

1. Introduction
Lettuce (*Lactuca sativa*) is a member in the Family of Compositae or Asteraceae that is grown for its leaves, which are either eaten raw or cooked (Grubben and Sukprakarn, 1994). It is a popular type of vegetable consumed in Malaysia. The production of lettuce in Sabah, Malaysia increased from year 2002 to year 2003 (Department of Agriculture 2003a, 2003b) thus increasing exports from 1997 to 2000 (Jipanin et. al., 2001).

However, the growth of lettuce is always hampered by bacterial spots caused by *Xanthomonas campestris* pv. *Vitians* (Barak et. al., 2002). Plants being attacked by this pathogen developed symptoms like spots on leaves and thus decreasing its commercial value in the market. In Malaysia, farmers usually apply large quantities of chemical to control this disease, thus increasing the cost of production. Pesticides also caused contamination to soil, water and air which is hazardous to human health (Agrios, 2005; Barak et. al., 2002). New awareness to reduce the usage of chemical pesticides by developing alternative strategies or technologies in order to improve plant disease resistance and control of pathogens are being promoted.

On the other hand, there is a positive linear relationship between the amount and speed of accumulation of phytoalexin (secondary antimicrobial compound) produced and the degree of disease resistance. This relationship is also known as
quantitative relationship. When there is a higher rate of phytoalexin accumulation, the smaller lesion size and lesser number of the bacterial cells is found in the host plant (Chong et. al., 2006a; 2006b; Chong et. al., 2007b; Mansfield, 2000; 2007b). In this study, the role of lettucenin A was tested with different concentrations on *Xanthomonas campestris* to verify the antimicrobial effect of lettucenin A to *X. campestris*.

2. Materials and Methods

2.1 Plant materials and pathogens culture

Seeds of lettuce were sown in trays containing sterilized moistened soil at 25°C. Later, three weeks-old seedlings were transferred from trays to pots. Elicitations were done when the plants reached eight weeks old. *Aspergillus niger* was kindly provided by Queen Elizabeth Hospital, Sabah and maintained on Potato Dextrose Agar (PDA) at 25°C in Plant Technology Laboratory of Universiti Malaysia Sabah. *Xanthomonas campestris* was obtained from Department of Agriculture, Tuaran, Sabah and maintained on Nutrient Agar (NA) at 25°C.

2.2 Elicitation and Extraction of lettucenin A

Production of lettucenin A was induced with abiotic elicitors. Leaves of lettuce were elicited by spraying the leaves with 5% of CuSO₄ or 1% of AgNO₃. The leaves were incubated at 25°C for three days and then subjected to extraction by methanol 60% overnight. Extracts were filtered through Whatman No. 1 filter paper, pooled and evaporated with a rotary evaporation at 40-45°C. Homogenates were re-extracted three times with chloroform, pooled and reduced to one ml.

2.3 Detection of lettucenin A and TLC Bioassays

Thin Layer Chromatography (TLC) plates (Merck Kieselgel 60 F₂₅₄ silica gel) were used throughout this study to detect the presence of antifungal properties of the leaf extracts and to separate lettucenin A from the crude extracts. The solvent system used was hexane: ethyl acetate (1:1, v/v). Lettucenin A gives off a bright yellow florescence when examined under UV radiation with wavelength peak of 365nm. Chromatograms were sprayed with the conidia suspensions of *Aspergillus niger* in potato dextrose broth for bioassays. Replicated chromatogram plates were prepared and sprayed with 2,4-dinitrophenylhydrazine reagent (2,4-DNPH), which gave a pink coloration after reacting with lettucenin A. Retardation value (Rf) for all bioactive and reactive bands were calculated and recorded.

2.4 Isolation of Lettucenin A

The confirmation of lettucenin A was based on the retention time (Rt) and UV absorption spectrum using High Performance Liquid Chromatography (HPLC) as described by Bennett et al., 1994. 10 µl and 50 µl of lettucenin A were injected into HPLC (Perkin Elmer Series 200), with SUPERCOIL™ LC-18 Analytical Column, 4.6 mm x 250 mm, 5µm. The presence of lettucenin A was analyzed using isocratic solvent system 60: 40 (v/v) water: acetonitrile, running for 15 min at 25 °C with a flow rate of 1.0 ml min⁻¹. The retention time (Rt) for lettucenin A was recorded and compared with the Rt as described by Bennett et. al., 1994.

2.5 Quantification of lettucenin A

From the replicated TLC plate, fluorescing band containing lettucenin A which had the same Rf value with the inhibition zone was marked by pencil, scraped out and eluted with 100% methanol. The tubes were spun at 12,000 rpm using a centrifuge for five minutes. Supernatant containing lettucenin A was collected and subjected to spectrophotometry and HPLC. For estimation of lettucenin A concentration, a Cary 50 Bio UV-Visible Spectrophotometer was used and set to 446 nm wavelength, which is the maximum wavelength absorbance of lettucenin A (Takasugi et. al., 1985).

2.6 Antibacterial activity of lettucenin A

Different concentrations of lettucenin A (67, 200, 333, 533 and 667 µg ml⁻¹), based on the amount recovered from TLC plates were prepared. Single colony of *Xanthomonas campestris* was transferred into petri dishes containing Nutrient Broth (NB). Initial optical density (OD) of the bacteria was measured by spectrophotometer at 600 nm. The bacteria were incubated until the OD₆₀₀ reading reached the range of 0.4-0.6 (Bacteria was in exponential stage). The bacteria within this range was tested throughout the study with different concentrations of lettucenin A in separate flask, and shaken at 220 rpm using a rotary shaker. One ml of each different lettucenin A concentration was added into each flask with three replicates for each concentration tested. After incubated for half an hour, OD₆₀₀ for each flask was measured and the procedures were repeated for the next one hour, 1.5 hours, two hours, 2.5 hours and three hours of incubation period. For control, lettucenin A was replaced by distilled water.

3. Results and discussion

3.1 Detection of lettucenin A

Lettucenin A accumulated after elicitation with CuSO₄ and AgNO₃, and it activities against *Xanthomonas campestris*, were studied. Lettucenin A gave a bright yellow fluorescence under UV radiation. In assessments of antifungal activity,
lettucenin A was proven to inhibit the growth of *Aspergillus niger* at R<sub>f</sub> 0.45. In addition, there were two other additional inhibition zones with the retention factor R<sub>f</sub> 0.80 and 0.90. Nevertheless, small and slightly inhibition zone was also observed in the control (Figure 1 (b)). Identification of lettucenin A was double confirmed after sprayed with the reagent 2,4-dinitrophenylhydrazine (2,4-DNPH). Lettucenin A displayed a pink coloration after sprayed with this reagent as illustrated in Figure 2. These bands had the retention value of 0.41 and they were not significantly different compared to the retention value that gave bright yellow fluorescence and strong antifungal activity in figure 1. Thus, this compound has been subsequently confirmed as lettucenin A. UV-spectrophotometer was used to scan the maximum absorbance. The maximum absorbance of lettucenin A in methanolic solution is at 446 nm (Figure 3) as described by Takasugi et al., 1985. The presence of lettucenin A was confirmed with HPLC based on the retention time (R<sub>t</sub>) as described by Bennett et al. 1994. Test was repeated twice with two different injection volumes, 10 µl and 50 µl. Both chromatograms had sharp peaks at R<sub>t</sub> approximately 5.3 min although the second injection volume was five fold higher (Figure 4). An arrow indicated that the peaks correspond to lettucenin A.

### 3.2 Antibacterial Activity of Lettucenin A

*X. campestris* showed different responses after exposure to lettucenin A. Bacteria growth rates were significant lower for those exposed to lettucenin A in comparison to the control experiment. *X. campestris* were sensitive to all concentrations of lettucenin A, at least, at the first 1.5 hours of incubation period. However, growth rates of treated bacteria were not significantly different compared to the control experiment when the duration of the experiments was extended to 2.0, 2.5 and 3.0 hours incubation period (Figure 5). For bacteria treated with 67 µg ml<sup>-1</sup> of lettucenin A, the growth rate was significantly lower in comparison to the control experiment at 0.5 and 1.5 hours, where the growth rate was 0.157 and 0.192, respectively. For other concentrations, the bacteria growth rate was significantly lower compared to control experiment. However, the growth rates of treated with lettucenin A were not significant in comparison to the control experiment at 2.0, 2.5 and 3.0 hours of incubation period, except for concentration of 333, 533 and 667 µg ml<sup>-1</sup> at 2.5 hours.

*In vitro* study showed a negative correlation between the concentrations of lettucenin A and the growth rate of pathogen *X. campestris*. Higher concentration of lettucenin A had a better inhibitory effect to the growth of *X. campestris*. But the bacteria growth was also inhibited in the lowest concentration tested; 67 µg ml<sup>-1</sup>. However, growth of *X. campestris* was more effectively restricted by higher concentration of lettucenin A (333, 533 and 667 µg ml<sup>-1</sup>). The effect of lettucenin A against bacteria at the concentration of 333, 533 and 667 µg ml<sup>-1</sup> was not significant among each other.

In other words, concentration of 333 µg ml<sup>-1</sup> maybe strong enough to restrict the bacteria growth. In most cases, higher concentrations of lettucenin A (333, 533 and 667 µg ml<sup>-1</sup>) were more effective against the bacteria. Different concentrations of lettucenin A had different effects on the growth of *X. campestris*. Thus, differences in the resistance and susceptibility of lettuce to *X. campestris* may associate with different concentrations of lettucenin A accumulated, as well as the speed of accumulation around the invading tissue. Localization of lettucenin A where this compound was concentrated in dead and infection sites provides good evidence in indicating the role of lettucenin A against pathogen *X. campestris*.

Besides that, the onset of hypersensitive reaction (HR) is always associated with rapid phytoalexins production. Phytoalexins were found localized to cells that had undergone HR. Fungal or bacterial invasion are then restricted in the HR cells. Accumulation of phytoalexins at the right time and place is very important parameter to cause cessation and restriction to microbial growth. After two hours of incubation, the growth of bacteria was not significant to control. This phenomenon occurred most probably because *X. campestris* may have the ability to detoxify the lettucenin A after the first 1.5 hours. Lettucenin A may be degraded by the bacteria into less toxic compounds. The ability of the bacteria to detoxify host’s phytoalexin is an important determinant of pathogenicity (Kuc, 1995; Mansfield, 2000; Purkayastha, 1995; vanEtten et. al., 1989). In other words, the antibacterial action of lettucenin A is considered as bacteriostatic.

Other examples of phytoalexin detoxification were proposed in the *in vitro* relationship of *Botrytis cinerea*, *B. fabae* with metabolism of wyerone, wyerone acid and wyerone epoxide of broad bean (*Vicia faba*). Detoxification of wyerone and wyerone epoxide was found preceding the onset of germ-tube growth of both *B. cinerea* and *B. fabae*; whereas production of secondary germ tubes from surviving conidia and sub-apical was occurred without comparable metabolism of wyerone acid. The concentration of wyerone was unlikely to decrease during the fungal-phytoalexin interaction (Rossall and Mansfield, 1984; Rossall et. al., 1980)).

Detoxification of phytoalexin by bacteria was also described in the *in vivo* relationship between cotton (*Gossypium* spp.) and bacteria *Xanthomonas campestris* pv. *malvacearum*. Cotton phytoalexins, 2,7-dihydroxyxycadaleine (DHC) was decomposed to lacinilene C (LC) while desoxyhemigossypol (dHG) was decomposed to hemigossypol (HG). Inhibition or toxicity occurred only for the first two hours of exposure to the phytoalexins. Growth rates of *X. campestris* pv. *malvacearum* were similar to the control in the following incubation period (Abraham et. al., 1999).
The amount of phytoalexin accumulated depends on the rate and duration of phytoalexin synthesis, which will be affected by the speed where cells are killed by the invasion pathogens and the ability to tolerate and degrade the phytoalexin (Mansfield, 2000). Thus, in vivo, the invasion of lettuce by *X. campestris* is regulated by the balance between rates of synthesis of lettenenin A and the accumulation in the plant and the degradation of that compound by the bacteria within infection tissues (Bennett et. al., 1994). Low concentration of lettenenin A would be produced at the beginning of the interaction. If *X. campestris* is tolerant and possess the enzymic capacity to detoxify lettenenin A, it will able to grow and detoxify higher concentration of lettenenin A and kill more cells. In this case, *X. campestris* is said to be pathogenic and the invaded lettuce is susceptible to it. If the bacterium is sensitive to lettenenin A, higher concentration of lettenenin A may produce at the infection site and thus inhibiting the further invasion of the bacterium. In this case, *X. campestris* is said to be non-pathogenic and the lettuce is tolerant to it (Bennett et. al., 1994).

4. Conclusion

*In vitro* activity showed that the bacteria was more sensitive to higher concentration of lettenenin A such as 333, 533 or 667 µg ml⁻¹ than lower concentration such as 67 µg ml⁻¹. There was a negative correlation between the concentrations of lettenenin A and the growth rate of the bacteria. However, after two hours of incubation period, the bacteria is suspected to have the ability to detoxify lettenenin A because the growth of bacteria was continuous. Lettenenin A is believed to have a role in the resistance of lettuce to microbial colonization in a sufficient concentration and at the right time. However, the exact concentration of lettenenin A that would inhibit the growth of pathogen *Xanthomonas campestris* under natural condition needs further study.

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References


Figure 1. Chromatograms of leaf extracts developed in hexane: ethyl acetate (1:1, v/v) as the solvent system. Lane 1: lettuce elicited with 5% CuSO₄; Lane 2: lettuce elicited with 1% AgNO₃ and Lane 3: control. (a) Observation under UV 365 nm wavelength (b) bioassay with *Aspergillus niger*.
Figure 2. Chromatogram of lettuce extracts after elicited with (a) 5% of CuSO₄ (w/v) and (b) 1% AgNO₃ and sprayed with reagent 2,4-dinitrophenylhydrazine (2,4-DNPH).

Figure 3. Ultraviolet absorption spectrum oflettucenin A range from 200-500 nm wavelengths in methanolic solution.

Figure 4. HPLC chromatograms at 446 nm excitation showing the presence oflettucenin A in (a) 10 µl and (b) 50 µl volume of injection.
Figure 5. Comparison of growth rate of *X. campestris* in five different concentrations of lettucein A for three hours of incubation.