Antimicrobial Potential of Vapour Phase Acetic Acid in Combination with Ethyl Alcohol against *Salmonella* Typhimurium Contaminated on Bird Eye Chili (*Capsicum frutescens* L.)

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**Abstract.** Bacterial contamination is the main cause of food poisoning. The currently decontamination methods arise to find novel, effective and safe approach. Acetic acid (AC) in both aqueous and vapour forms have been conducted on its effectiveness against several bacterial contaminations. The use in high concentration can cause strong smell and corrosives that have an effect on the qualities of food. This study aims to assess the potential of Mechanically Vapourized AC solution in combination with ethyl alcohol (EA) (MVA-E) on the reduction of *Salmonella enterica* serotype Typhimurium ATCC 13311 contaminated on both microbiological media and on fresh Bird Eye Chili (*Capsicum frutescens* L.). In vitro surface inhibitions of *S.* Typhimurium at low population, ca. 1.0 CFU/cm², and high population, ca. 10.0 CFU/cm² were examined on Tryptic Soya Agar (TSA). MVA-E at the concentration of 10:95 (AC:EA) presented the absolutely inhibited *S.* Typhimurium within 20 min at 27±2°C. At the concentration of 10:75 (AC:EA), the absolutely inactivated *S.* Typhimurium was observed within 10 min at 50±2°C. For the evaluation of antimicrobial activity of MVA-E over time, the results indicated that ca. 8.00 Log₁₀ reductions were observed within 20, 25 and 25 min at the concentration of AC:EA ratio as 10:95, 10:75, and 10:45, respectively at 27±2°C. The effectiveness of MVA-E increased when the temperature of MVA-E process increased. The reduction of *S.* Typhimurium contaminated on fresh Bird Eye Chili by MVA-E was also determined at 4°C, 27±2°C and 50°C. The efficiency of MVA-E on the reduction of *S.* Typhimurium depended on the concentration of EA in MVA-E process, the fumigation time and also the temperature. To the best of knowledge this is the first time a combination of AC and EA in vapour phase has been tested as a preservative method prevent microorganism proliferation.

1 Introduction

Chilies (*Capsicum* sp.) have been cultivated in many regions of the world [1]. One of the most consumed chili in Thailand is Bird Eye Chili, (*Capsicum frutescens* L.), a wide range of chili fruits is available in ASEAN’s markets [2]. It was used as pungent ingredients in several culinary practices [3]. Also it is green ingredient used to prepare several meals, some people consume as fresh fruit. Chilies, such as Bird Eye Chili are often contaminated by microorganisms on the surface and can cause food poisoning illness [4-7]. The outbreaks caused by consumptions of contaminated those with foodborne pathogens as *Salmonella* sp., *Listeria monocytogenes*, *Vibrio* sp. or others [8-10]. As a consequence of such situation, contaminations of *Salmonella* sp. in vegetables become the greatest for public health concerns. Postharvest handling such as washing or spraying with sanitizer is most important procedure in view of the fact that removed, eliminated or reduced the surface microbiological contaminant. The US-FDA suggested the use of 200 mgL⁻¹ chlorine, at a pH of 6.0-7.5, for 1-2 min as disinfection processes for fresh produce. Several reports had indicated that chlorine at that level lacks the effectiveness to eliminate spoilage microorganisms. In addition, chlorine presented the potential to form carcinogenic compounds [11]. Several researchers have projected other sanitization processes [12-13]. Among of those, AC presented the special interest [14-15]. Volatile antimicrobial have become popular in research and have also been exhibited antimicrobial properties in many food product [14, 16-18]. The use of acetic AC in vapour phase was reported about their potential to control the growth of microorganism. However, the use of AC alone in vapour phase caused the strong vinegar odour and affected to the appearance quality of vapourized fresh produces. The use of volatile antimicrobial substances such as EA or in combination with another should be the good alternative technique. Thus, in this study the efficacy of mechanically vapourized AC (MVA) in combination with mechanically vapourized EA (MVE) as mechanically vapouized AC-EA (MVA-E) on the
reduction of S. Typhimurium in both in vitro study and contaminated on model fresh Bird Eye Chili was investigated.

2 Materials and Methods

2.1 Chemical and microbiological media

Tryptic Soya Agar (TSA), Tryptic Soya Broth (TSB) and Peptone were purchased from Difco (Dico, USA). AC was purchased from QRëC (QRëC, New Zealand). 95% Peptone were purchased from Difco (Dico, USA). AC Tryptic Soya Agar (TSA), Tryptic Soya Broth (TSB) and contaminated on model fresh Bird Eye Chili was purchased from QRëC (QRëC, New Zealand). 95% Peptone were purchased from Difco (Dico, USA). AC Tryptic Soya Agar (TSA), Tryptic Soya Broth (TSB) and contaminated on model fresh Bird Eye Chili was kindly provided by The Liquor Distillery Organization of Thailand.

2.2 Preparation of Test Organism

S. Typhimurium, was kindly provided by Department of Food Science and Technology, Thammasat University (Rangsit Centre). Period of expose, S. Typhimurium was sub-culture twice in TSB at 37°C for 18 h before use as inocula.

2.3 In vitro susceptibility of S. Typhimurium to the MVA-E

2.3.1 Agar overlay method

The susceptibility of S. Typhimurium to MVA-E was determined in vitro using modified agar overlay method. Briefly, The S. Typhimurium at ca. 7.0 Log_{10} CFU/mL was used as high inoculums and at ca.4.00 Log_{10} CFU/mL as low inoculum. Contaminated Petri dishes without cover were aseptically placed in fumigation chamber as presented in Fig. 1. AC solution at the concentration of 10.0% in combination with EA at the concentration of 0%, 30.0%, 45.0%, 75.0% and 95.0% was individually placed and directly connected with air pump.

Fig. 1. Schematic illustration represented the fumigation chamber

Plates were contacted with MVA-E and withdraw at the interval time as 0, 5, 10, 15, 20 and 10 min. The reduction ratio was calculated after incubation. The impact of temperature was determined at 4°C and 50°C. The rate of vapour production was calculated.

2.3.2 Time killing analysis

For time killing analysis, the susceptibility of S. Typhimurium to MVA-E was determined in vitro. S. Typhimurium at ca. 8.00 Log_{10} CFU/mL was prepared in 50 mL TSB. Fumigation tube connected with MVA-E was aseptically placed into suspension. MVA-E was generated. At the interval time as 0, 5, 10, 15, 20 and 25 min the microbial populations were enumerated by spread plate technique on TSA. As described above, the impact of temperature was also determined.

2.4 MVA-E effects on fresh Bird Eye Chili inoculated with S. Typhimurium

2.4.1 S. Typhimurium suspension preparation

S. Typhimurium was sub-cultured twice in TSB. Cells were harvested by centrifugation at 1000xg for 15 min at 4°C. The inocula was adjusted to the final concentration at ca 7.00-8.00 Log_{10} CFU/mL

2.4.2 Challenging Test

Bird Eye Chilies were purchased from local markets near Thammasat University. Each fruit was sanitized in 1000 ppm chlorinated water. The fruits were dry under UV lamp. 0.1 mL of prepared S. Typhimurium was contaminated on each to obtain 6.00 Log_{10} CFU/g. The contaminated those were then exposed to MVA-E in the fumigation chamber at 65±2% RH. Each fumigated fruit was then examined for evidence of survival of S. Typhimurium by spread plate technique. The population of organism was calculated as Log_{10} CFU/g. The impact of temperature was also determined.

3 Results and Discussions

3.1 Susceptibility of S. Typhimurium to MVA-E

3.1.1 In vitro agar overlay method

Efficacies of MVA-E against S. Typhimurium on the surface of TSA are demonstrated in Fig. 2. The results presented the inhibitory effect of MVA-E against S. Typhimurium at 95.0% and 75.0% of EA in combination with 10.0% AC at 27±2°C within 20 and 25 min, respectively. At the low concentration, the complete destructive effect was not detected. These inhibitory properties were detected in both low and high inoculums level. As shown, the temperature affected the antimicrobial properties of MVA-E. At low temperature (4°C), the efficacy of MVA-E was decreased compared with room temperature (27±2°C), and increased when exposed with high temperature as 50°C. According to Fig 3, the increasing rate of pH and weight loss of both AC and EA under vaporized process were the main factors affecting the antimicrobial properties. These increasing were obviously presented when those were exposed to high temperature. Hence, pH of test solution and weight loss of AC and EA increased along with the rising of temperature, the antimicrobial properties of MVA-E also increased when the temperature increased. As mentioned in the previous research, the inhibition potential of AC was acted upon the pKa and pH [20,21]. The bacteriostatic and bactericidal of AC and the other
derivatives have been attributed to the lower pH below

The other possibility of the impact of temperature on the
antimicrobial properties of MVA-E is that the upsurge in
The temperature demonstrated the effect on antimicrobial properties of MVA-E. At the higher temperature, the antimicrobial activity was greater than at the low temperature. The time killing analysis is equal to the inhibition curve, known as the ‘killing curve’. In generally, this studies showed that the survivors of those was decreased by concentration of EA was increased, and completely inhibited at the concentration more than 45.0%. Moreover, it could be indicated that the longer the fumigation time the lower the necessary concentration of the MVA-E. The Time-Dependent characteristic was observed according to this study.

3.2 MVA-E effects on fresh Bird Eye Chili inoculated with S. Typhimurium

3.2.1 Challenging Test

The initial amount of contaminated S. Typhimurium on Bird Eye Chili at ca. 6.00 Log10 CFU/g were exposed to MVA-E at the concentration of AC at 10.0% in combination with the different concentration of EA along different fumigation time.
The effect of operating temperature was also determined. The results were shown in Fig. 5. The natural contaminated Bird Eye Chilies were also tested. The numbers of Total Bacteria Count (TBC) were determined as describe above. The results of TBC were presented in Fig 6. Fumigation with MVA-E at room temperature (27±2°C) and 50°C at all concentration of EA demonstrated the lethal effect. At the concentration of AC at 10.0% in combination with 95.0% EA the lethal effect was presented within 25 min. When the temperature decreased to 4°C the complete destruction was not detected along the period of fumigation. The downward trend of fumigation time was observed when the temperature was rising. Weak acid always demonstrate the microbial inhibition properties [19-20].

Mechanically vapourized AC in combination with EA (MVA-E) presented the inhibitory potential to eliminate and decontaminate the number of S. Typhimurium in both in vitro and model food as contaminated on fresh Bird Eye Chilies. It is free of carcinogenic risk. The antimicrobial properties depended on the concentration, fumigation time and the temperature. When these key factors increase, the antimicrobial properties are also increased.

![Fig. 6. Reduction of Total bacteria count contaminated on Bird Eye Chili as affect by MVA-E at different EA concentration.](image)

Consideration on the effective of MVA-E against natural contaminated microorganism on Bird Eye Chili, it could be indicated that the complete elimination was not detected within the period of exposure.

![Fig. 7. Schematic illustration of the possible mechanism of MVA-E against S. Typhimurium.](image)

As presented in Fig. 7, the possible antimicrobial mechanism of MVA-E was demonstrated. It seems that MVA-E decontaminates surfaceborne microorganism and thus sterilizes vegetable surfaces [24]. In this research, the results demonstrated that MVA-E presented the inhibitory effect on both S. Typhimurium in vitro and in model food as Bird Eye Chilis. The mechanically vapourized process allowed the AC in the form of dissociated to become the undissociated molecule in vapour phase. For this reason, the MVA-E presented more inhibitory properties. Another advantage, the gas phase can diffuse much more than liquid phase so the rate of cell membrane penetration become more rapid compared to liquid phase.

4 Conclusions

References