Effect on Human Health by Residues of Commonly Used Pesticides in Vegetables Cultivation

Farzana Khalil, Maliyat Tarannum Maruf, Mohammad Tariqu Islam, Mohammad Mahbub Rabbani, S. Mosaddeq Ahmed

Abstract—People in Bangladesh are over scared for toxicity of vegetables & fruits, because the educational level of the farmers is not up to the mark. Farmers apply pesticides randomly on agricultural fields without maintaining pre-harvest intervals and dose. The aim of this work was to study dissipation pattern of some pesticides in some vegetables kept at room temperature which represent market condition and to identify and quantify pesticides applied on some vegetables. By interviewing the farmers, three commonly used pesticides; cypermethrin, chlorpyrifos and fenvarelate were sprayed in the farmer’s fields of the western part of Bangladesh. Samples were extracted by QuEChERS method, cleaned-up by adsorption chromatography technique, and analyzed by GC-ECD technique. Recoveries were found to be between 87-107% with RSD lower than 10% at three spiking levels. Matrix matched calibration curves were linear for all the analytes with \(r^2 \geq 0.99\). LODs were found to be 0.01 mg/kg for cypermethrin and 0.002 mg/kg for both of fenvalerate and chlorpyrifos. The samples of the same cultivar’s variety were purchased from the markets of Savar, Mymensingh and Cumilla. Our finding showed only the presence of cypermethrin in tomato but the values were below MRL and no pesticide residue was found in eggplant.

Index Terms—Pesticide residues, health hazard, tomato, eggplant, pre-harvest intervals.

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I. INTRODUCTION

Food products can be contaminated by several sources and processes through the food chain from production to consumption. This contamination has a harmful effect on the quality of the foodstuff and may posture a health risk. It is essential to ensure contaminant levels not to exceed Maximum Limits (MLs) for the protection of public health. Nowadays concerns regarding contamination in foodstuffs are a major issue in all countries especially in Bangladesh.

Organophosphate insecticides break down very rapidly and hence have very low persistence in the environment. Therefore, they do not cause any long-term hazards. However, these compounds are extensively toxic in nature. Further, due to their instability they must be applied at frequent intervals. Therefore, they are not always economically viable [1].

The most recent group of insecticides is the pyrethroids which have two important advantages, neither persistent nor toxic. It has been observed that metabolites of pyrethroid pesticides are relatively low toxic or non-toxic compared with parent pyrethroid compounds [2]. They were reported to have relatively low toxicity as compared to organophosphorus, organochlorine or carbamate pesticides [3]. Therefore, despite their high cost, they account for about one-third of world insecticides use [4].

Chlorpyrifos (0,0-diethyl 0-3,5,6-trichloro-2-pyridinyl-phosphorothioate) is a broad-spectrum, chlorinated organophosphate (OP) insecticide, acaricide and nematicide [5]. Cypermethrin [(RS)-α-cyano-3-phenoxybenzyl(1RS, 3RS; 1RS, 3RS)-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane carboxylate] and fenvarelate [(RS) -α-cyano -3- phenoxy benzyl (RS) -2- (4-chlorophenyl) -3- methyl butyrate] are the synthetic pyrethroids which have become most important insecticides in wide-scale use. These are effective against the pests which are resistant to organochlorine, organophosphorous, and carbamate insecticides [6]. It has been said that "no pesticide is perfect, but the pyrethroids come close" [7].

Chlorpyrifos may decomposes in plants and produce chlorpyrifos oxon and 3, 5, 6-trichloro-2-pyridinol, which is further degraded to 3, 5, 6-trichloro-methoxypyridine and carbon dioxide [8]. Cypermethrin may photo degrades rapidly and subject to microbial degradation. After photo degradation
it may produce DCVA, 3- phenoxybenzaldehyde and 3-phenoxybenzoic acids [9]. Fenvalerate and its active isomer, esfenvalerate, predominantly undergo ester cleavage through oxidation at the acid and alcohol moieties and conjugation reactions. At pH 9.0 it undergoes ester hydrolysis, resulting in the formation of 2-(4-chlorophenyl)-3-methylbutyric acid (CPIA) as a major degradation product. It is rapidly photo-degraded with the action of UV light to the major product, 2-(3-phenoxyphenyl)-3-(4-chlorophenyl) methylpentanenitrile (decarboxy-fenvalerate) [10].

The vegetables are low in fat, high in dietary fibers; contain water, minerals and vitamins, possessing a very high nutritional density. So, it is advised peoples to consume more vegetables in meals. But in our country farmers have been using pesticides frequently in vegetables especially on bean, eggplant, cauliflower and tomato to get higher yields. The overdose, frequent uses of pesticides and not following the pre-harvest time frame left residues in food which pollute our food and be harmful for our health. Since it is very difficult to monitor and assess the pesticide contamination, pesticides residue is becoming a foremost food safety concern of consumers and government.

Pesticide usage cannot be properly regulated due to ineffective legislation, lack of awareness and technical knowhow among the farming society in Bangladesh. Belgium is an agricultural country. In winter season many vegetables are being cultivated every year more or less in all area of the country. In west and north part of this country vegetables are grown extensively. It has been reported that farmers use pesticides randomly without following the label directions and collect vegetables and sell in market not following the pre-harvest interval. Sometimes they harvest vegetables at the same day or next day after pesticides spraying and these vegetables go to consumers within 2-4 days from pesticides application time. For this reason, vegetables were collected from Jhenidah district (Fig. 1), a western district of Bangladesh after spraying pesticides on vegetables by farmers to see the dissipation pattern of the pesticides kept at room temperature which represent market conditions. The samples of same cultivar’s variety were also purchased from the wholesale & city markets of Savar, Mymensingh and Cumilla district in two different days for each vegetable.

Staff and other facilities of government to perform the necessary monitoring programs are not available. Besides the country is not yet established its own authorized limits for residues and depends upon Codex acceptable limit. So, farmers should be recommended the exact dose to eliminate residual effect of pesticides on consumers. In addition, the pre-harvest interval of each pesticide should be known so that the amount of residual pesticides in vegetables might be lower than the tolerable range. The present study intended to investigate and quantify the residues of some commonly used pesticides on selected vegetables grown under local agro-climatic conditions of Bangladesh.

In 2003, Anastassiades et al. introduced a new method of analysis, namely QuChERS, an acronym for quick, easy, cheap, effective, rugged and safe, covers a variety of sample preparation and clean-up techniques for the analysis of multiple pesticide residues in agricultural matrices. The method uses acetonitrile (MeCN) for extraction (1 mL MeCN/1 g sample) by twister mixing followed by an addition of 4:1 (w/w) anhydrous MgSO₄:NaCl (0.5 g salts per g sample) to separate MeCN extract from the water in the sample. After centrifugation, dispersive solid-phase extraction (dispersive SPE) approach was applied to mix-up 1 mL of the extract with 25 mg primary secondary amine (PSA) sorbent and 150 mg anhydrous MgSO₄. The extract is centrifuged again and transferred to an auto-sampler vial for analysis by gas chromatography/mass spectrometry (GC/MS) and/or other technique. This method is popular due to its simplicity, low cost, less time consuming, reducing glassware and solvent requirement, producing relatively high efficiency results and minimal number of steps it involves [11]. In this study, QuChERS method of extraction and cleanup was carried out by adsorption column using mixture of florisil, alumina and charcoal. Here GC-ECD was used for analysis of pesticide residues.

II. EXPERIMENTAL METHODS

A. Chemicals and Reagents:

Cypermethrin (91% purity), chlorpyrifos (99.5% purity), and fenvalerate (98.5% purity) purchased from Dr. Ehrenstorfer, Germany were used for analysis and related information of these three pesticides are presented in Table 1. Ethyl acetate, acetone, n-hexane, dichloromethane, methanol (pesticide residue grade), anhydrous magnesium sulphate, sodium chloride (analytical grade), and aluminium oxide (alumina) from Merck, Germany, florisil from ACROS organics, USA, charcoal from Uni-Chem, China were used for this analysis. Florisil and alumina were activated by heating at 105 °C for 3 h, charcoal was activated by washing with distilled water, n-hexane, methanol, acetone in Buchner funnel & then dried at 105 °C and all kept in desiccators.

B. Standard Solution Preparation:

100 ppm primary stock solutions of cypermethrin, chlorpyrifos and fenvalerate were prepared by dissolving an appropriate amount in n-hexane. These primary standard solutions were diluted to 20 ppm and 5 ppm middle and
working standard solutions respectively. These solutions were labeled indicating name of the standard, concentration and date of preparation. These solutions were stored in amber bottles (100 mL) in the freezer (-24°C) away from the pesticide residue laboratory.

C. Field Experiment and Sampling:

Eggplant (Solanum melongena var. esculentum) and tomato (Lycopersicon lycopersicum) were collected from the farmer’s fields of Jhenidah district (Figure 1). Cypermethrin & fenvalerate were sprayed by farmers at recommended dose on eggplant in two different fields whereas chlorpyrifos together with cypermethrin were applied on tomato in another field. Before pesticide spraying samples were collected from the respective fields. After two hours samples (nearly 5 kg) were collected and one subsample (1 kg) kept in chill-box (0 day sample) and then all the samples were transferred to the laboratory. Control and 0 day samples were kept in freezer and rest of the samples were kept in normal temperature. These normal temperature subsamples were kept in freezer after 1, 3, 5, 7 & 10 days. Before store in freezer the samples were chopped and homogenized in kitchen blender.

D. Extraction Process:

20 mL ethyl acetate was added to 10 g homogenized sample in 50 mL teflon tube and shaken for 1 minute in hand & vortex for 1 min. 6 g anhydrous MgSO₄ & 1.5 g NaCl were added and vortexed for 1 min and then centrifuged for 5 minutes at 4000 rpm. 10 mL supernatant solution was taken in 100 mL round bottomed flask, evaporated in rotary evaporator and then reconstituted in n-hexane (2 mL).

E. Purification:

A glass column (40 cm long & 12 mm internal diameter) was packed with a 10.5 g mixture of aluminum oxide, florisil and charcoal (10:10:1) in n-hexane. The column was equilibrated with 50 mL n-hexane and then the sample extract in n-hexane (2 mL) was transferred to the column. The column was washed with 20 mL of n-hexane and eluted with 100 mL of dichloromethane at the rate of 1 mL min⁻¹. The eluent was concentrated to dryness on a rotary vacuum evaporator and dissolved the residue in 2 mL of n-hexane and injected to GC-ECD. Extraction and purification methods used in this experiment are depicted in Figure 2 using a flow chart.

F. GC-ECD Analysis:

A gas chromatograph (GC-2010 Shimadzu) equipped with ⁶⁰Ni Electron Capture, (EC) detector was used in this experiment. A non-polar (Rtx-5 MS) or (HP-5 MS) Quartz capillary column (30 m long × 250 μm i.d. × 0.25 μm film thicknesses) from Agilent, USA was used to carry out the separation. Nitrogen was used as both carrier (column flow 1.92 mL/min.) and make up gas. The injector temperature was 220 °C & the detector temperature was 290 °C. The injection volume was 1μL and it was carried out in a split-less mode. The oven temperature was set as - initial temperature of 120 °C held for 2 minutes; increased at 10 °C min⁻¹ to 270 °C; held for 1 min. and then another was increased at 2 °C min⁻¹ to 290 °C; held for 3 min. The column flow was 1.0 mL min⁻¹. Total program time was 31.00 minutes.

G. Quality Assurance:

At first the control vegetables (samples collected before pesticide spraying) matrices were confirmed that there were no pesticides by doing blank experiments. The recovery experiments were done with these control matrices. The recovery experiments were performed in three replicates at 2-3 fortification levels. The fortified samples were left to stand for 2 h to allow for the adsorption of pesticides onto the samples.

<table>
<thead>
<tr>
<th>Name</th>
<th>Chemical structure</th>
<th>Chemical class</th>
<th>LD₅₀ mg/kg</th>
<th>MRL mg/kg</th>
<th>ADI mg/kg b.w.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlorpyrifos</td>
<td><img src="image" alt="Chemical Structure" /></td>
<td>Organophosphorus</td>
<td>Rat (oral) 135-163</td>
<td>0.05</td>
<td>1.0 (tomato)¹¹</td>
</tr>
<tr>
<td>Cypermethrin</td>
<td><img src="image" alt="Chemical Structure" /></td>
<td>Pyrethroid</td>
<td>Rat (oral) 250-4150</td>
<td>0.2 (tomato)¹¹</td>
<td>0.05 (JMPR)</td>
</tr>
<tr>
<td>Fenvalerate</td>
<td><img src="image" alt="Chemical Structure" /></td>
<td>Pyrethroid</td>
<td>Rat (oral) 451</td>
<td>1.0 (eggplant)¹³</td>
<td>0.02 (JMPR)</td>
</tr>
</tbody>
</table>
These samples were then extracted, purified and analyzed following the aforementioned procedures. The pesticide residues in the fortified, field and market samples were quantitatively determined by external standard method using peak area. The linear matrix matched calibration curves for all pesticides over six calibration levels, from 0.025-2.0 mg/L were constructed by the direct injection of calibration standards into the GC and all standard curves were within the acceptable limits of the linearity criterion which are shown in Table 2.

**III. RESULTS AND DISCUSSION**

**A. Matrix Effect:**

In quantitative analysis by gas chromatography matrix effect is defined as a foremost setback. It is regarded as a signal suppression or enhancement of the analytes due to the co-elution of matrix components with the analytes, resulting in poor analytical accuracy, linearity, and reproducibility [13-14]. The signal response of the analytes can vary considerably from matrix to matrix and differs significantly in pure solvent and in matrix. Therefore, it is needed to use the matrix-matched standards to make calibration curve in order to nullify quantitative errors in pesticide residues analysis. This effect was rather mild and not so much significant [15].

**B. Specificity:**

The unnecessary compounds interfering with the analytes were examined by comparing the chromatograms of the standard, blank sample and fortified sample. There were no interference peaks at the retention time of chlorpyrifos, cypermethrin, and fenvalerate observed.

**C. Linearity:**

In this study, calibration curves were prepared in matrices that extracted from control matrix (eggplant and tomato) and the linearities, limits of detection (LODs) and limits of quantification (LOQs) were calculated which are shown in Table 2. Matrix matches calibration curves (fenvalerate in eggplant, cypermethrin & chlorpyrifos in both eggplant & tomato) were constructed in the range of 0.025-2 mg/L. The linearity with a correlation coefficient of $r^2 \geq 0.99$ was excellent. The residual concentrations of cypermethrin, fenvalerate and chlorpyrifos in both the treated and market samples were determined using the matrix matched calibration curves developed herein.

**D. Limits of Detection and Quantification:**

The limit of detection (LOD) defined as the minimum concentration of analyte in the test sample that can be measured with a stated probability that the analyte is present at a concentration above that in the blank sample. On the other hand the limit of quantification (LOQ) implies the minimum concentration of analyte in the test sample that can be determined with acceptable precision (repeatability) and accuracy under the stated conditions of the test [16]. The LOD and LOQ were found 0.01 mg/kg and 0.033 mg/kg for the cypermethrin in tomato, 0.002 mg/kg and 0.0066 mg/kg for

**Table 2 Name, correlation coefficients ($r^2$), LODs, LOQs and recoveries of the tested pesticides**

<table>
<thead>
<tr>
<th>Pesticides &amp; Vegetables</th>
<th>Linearity ($r^2$)</th>
<th>LOD &amp; LOQ (mg/kg)</th>
<th>$^a$Accuracy (% recovery), Precision (% RSD) (Spiking level, mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cypermethrin in tomato</td>
<td>0.997</td>
<td>0.01, 0.033</td>
<td>87, 8.45 (0.05) 102, 5.38 (1.0) 97, 4.64 (0.25)</td>
</tr>
<tr>
<td>Fenvalerate in eggplant</td>
<td>0.995</td>
<td>0.002, 0.0066</td>
<td>98, 3.85 (0.25) 107, 7.34 (0.5) 85, 6.72 (1.0)</td>
</tr>
<tr>
<td>Chlorpyrifos in tomato</td>
<td>0.998</td>
<td>0.002, 0.0066</td>
<td>89, 6.83 (0.025) 95, 9.15 (0.05) 103, 3.61 (0.1)</td>
</tr>
</tbody>
</table>

$^a$Mean of three replicate

The matrix effect (ME) was determined by using a standard solution prepared in sample extract and pure solvent. ME % was calculated via the following equation [12].

$$\text{ME %} = \frac{(\text{Peak area of matrix standard} - \text{Peak area of solvent standard}) \times 100}{\text{Peak area of solvent standard}}.$$
fenvalerate in eggplant, 0.002 mg/kg and 0.0066 mg/kg for chlorpyrifos in eggplant & tomato, respectively (See Table 2).

Table 3 Pesticide residues (Av. ± SD, mg/kg) in vegetables at various days following its application

<table>
<thead>
<tr>
<th>Samples after pesticides spraying</th>
<th>Day after spraying</th>
<th>Cypermethrin in tomato</th>
<th>Fenvalerate in eggplant</th>
<th>Chlorpyrifos in tomato</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>-</td>
<td>BDL</td>
<td>BDL</td>
<td>BDL</td>
</tr>
<tr>
<td>0</td>
<td>0.39 ± 0.04</td>
<td>0.68 ± 0.05</td>
<td>1.73 ± 0.07</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.28 ± 0.05</td>
<td>0.45 ± 0.03</td>
<td>1.31 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>0.15 ± 0.017</td>
<td>0.27 ± 0.07</td>
<td>0.95 ± 0.13</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>0.12 ± 0.006</td>
<td>0.16 ± 0.02</td>
<td>0.73 ± 0.09</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>0.07 ± 0.002</td>
<td>0.09 ± 0.08</td>
<td>0.54 ± 0.04</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>0.03 ± 0.012</td>
<td>0.05 ± 0.01</td>
<td>0.24 ± 0.02</td>
<td></td>
</tr>
</tbody>
</table>

E. Recovery:
The extraction efficiency was assessed by doing recovery experiment. The recovery experiments were done with control samples collected from field before pesticide spraying which were initially confirmed that there were no pesticides. The recovery experiments were performed in three replicates at three fortified concentrations. The recoveries were 87-102% for cypermethrin in tomato, 92-107% for fenvalerate in eggplant and 89-103% for chlorpyrifos in tomato with precision below than 10%.

F. Dissipation of Pesticides in Field Samples:
Farmers in our country are applying pesticides in their field for crop productivity and selling the crops directly into the markets. Concerning in food safety & security, the study was designed to reproduce the way of using pesticides in the real fields and the samples available in the market for the consumers. The amounts of Chlorpyrifos, cypermethrin and fenvalerate residues found in different vegetables after variable days of treatment with pesticides are reported in Table 3.

Discussing with farmers, it was selected the vegetables fields where pesticides were not applied within 10 days. No residues were detected on blank vegetables collected from the same fields before pesticides spraying. The sample collection was started after 2 hours of pesticides spraying. The pesticides were dissipated with time. The maximum residue levels were detected on 0 (zero) day. The residue concentrations of cypermethrin were found 0.39 mg/kg in tomato on 0 (zero) day and 0.03 mg/kg (87% dissipation) in tomato after 10 day. The residue concentrations of fenvalerate were found 0.65 mg/kg on 0 (zero) day and 0.05 mg/kg (86% dissipation) in eggplant after 10 day. Similarly, the residue concentrations of chlorpyrifos were found 1.73 mg/kg in tomato on 0 (zero) day and 0.24 mg/kg (78% dissipation) in tomato after 10 day. This result indicates that the percentage of dissipation of cypermethrin and fenvalerate on different vegetables are similar (86-87%) but chlorpyrifos dissipates slowly (78%) with time. The dissipation patterns with time have been shown in Figure 3 for cypermethrin in tomato, in Figure 4 for fenvalerate in eggplant and in Figure 5 for chlorpyrifos in tomato. The amount of residues gradually decreases with the increase of time elapsed for all the three pesticides. As the amount of chlorpyrifos residue is high on zero day and also the rate of dissipation is lower than other two pesticides studied so it will take longer time to be disappeared from the vegetable completely. These results suggest that chlorpyrifos stays on vegetables for longer time and hence farmers should allow considerable days after spraying to harvest vegetables.

Fig. 3. Dissipation curve of cypermethrin in tomato

Fig. 4. Dissipation curve of fenvalerate in eggplant.

On the context of maximum residue limits (MRL), it can be observed from Table 3 that Cypermethrin residues went below the MRL value at 3 days in tomato (0.2 mg/kg) and...
fenvalerate residues went below MRL value (1.0 mg/kg) at 0 day in eggplant. One the other hand Chlorpyrifos residues went below the MRL value (1.0 mg/kg) at 3 days after spraying in tomato. These results suggest that if Cypermethrin and Chlorpyrifos are sprayed on tomato the cultivators should allow minimum 3 days to harvest the vegetables but in case of fenvalerate spray it is safe anytime.

G. Analysis of Market Samples:

Eggplant and tomato samples of the same cultivar’s variety were purchased from three local markets of Savar, Cumilla and Mymensingh two times. The amounts of pesticide residues found are listed in Table 4. It is noted that Cypermethrin was detected in tomato samples but no pesticides were found in eggplant samples. However, the amount of Cypermethrin was much lower than the MRL value. The vegetables might be picked up after reasonable interval between two consecutive insecticide sprays.

<table>
<thead>
<tr>
<th>Market</th>
<th>Sampling</th>
<th>Eggplant</th>
<th>Tomato</th>
</tr>
</thead>
<tbody>
<tr>
<td>Savar</td>
<td>1st</td>
<td>BDL</td>
<td>BDL</td>
</tr>
<tr>
<td></td>
<td>2nd</td>
<td>BDL</td>
<td>Cp = 0.0065±0.0007</td>
</tr>
<tr>
<td>Cumilla</td>
<td>1st</td>
<td>BDL</td>
<td>Cp = 0.043±0.004</td>
</tr>
<tr>
<td></td>
<td>2nd</td>
<td>BDL</td>
<td>Cp = 0.039±0.002</td>
</tr>
<tr>
<td>Mymensingh</td>
<td>1st</td>
<td>BDL</td>
<td>Cp = 0.056±0.004</td>
</tr>
<tr>
<td></td>
<td>2nd</td>
<td>BDL</td>
<td>Cp = 0.025±0.001</td>
</tr>
</tbody>
</table>

H. Health Risk Assessment:

Estimated daily intake (EDI) of pesticide residues was calculated by multiplying the residual pesticide concentration (mg/kg) by the food consumption rate (0.345 kg/person/day) and dividing by a body weight of 60 kg for an adult people [17]. The Maximum Residue Limit (MRL) for cypermethrin in tomato is 0.2 mg/kg, eggplant is 0.03 mg/kg, according to Codex Alimentarius Commission [18-19] and the Acceptable Daily Intake (ADI) for human is 0.05 mg/kg body weight. The health risk index (HI) was also calculated. The HI showed that cypermethrin residues in all vegetables were found less than 1 (HI<1). From the concentrations of cypermethrin residues obtained, dietary exposure and health risks were calculated for adult. The statistic of daily food intake of an adult person of our country exposes that vegetables are in third position after rice, cereal and fish which give the daily food consumption in weight [20]. When HI is less than 1, the food concerned is considered as acceptable. If the value is greater than 1 the food is considered a risk to the consumer [21-22]. Most of the vegetable samples were not harmful to health as all samples had cypermethrin lower than the MRL values [23]. World Health Organization has settled MRL for pesticide residues in different vegetables. In line with that very recently Bangladesh Food Safety Authority has enacted a regulation titled “Chemical Contaminants, Toxins and Harmful Residues” [24].

IV. CONCLUSION

The method achieved good validation results (good repeatability/RSD and accuracy/recovery) within a short time. Results showed that all pesticides went below the MRL values very quickly. So the applied dose of pesticides by the farmer in the field to control pests in vegetables should be lower or pre-harvest interval should be longer. As a result, all samples were lower the MRL, so the vegetables samples might not competent to serious health hazard.

The most serious alarm about the pesticide use is its hazardous effects on different components of the environment. In Bangladesh context, the vegetables growers have been using the pesticides frequently to have advanced and pest free harvest. But the overdoses of pesticides make the residue problematic, which might contaminate our food and environment. Recently a multiplicity of Good Agricultural Practices (GAP) codes, standards and regulations has been developed by various food industry and producers organizations in first world countries. Bangladesh being a developing country, we need to develop a culture of Good Agricultural Practice to ensure safety and quality of product in the food chain, to capture new market advantages by modifying supply chain governance, to improve natural resources use, workers health and working conditions, and to create new market opportunities for farmers and exporters. We need to train our farmers, retail suppliers and consumers Good Agricultural Practices that address environmental, economic and social sustainability for on-farm processes, and result in safe and quality food.

There is always a time gap between the harvest and availability of vegetables in the local market. It is not enough to deliver manufacturers simply with a manual of food safety guidelines and skillful implementation and documentation. From these studies it is worthwhile for the consumers to keep vegetables at ambient temperature rather than store in a refrigerator for 2-3 days. Keeping the view in mind of the above reports, we want to suggest our Bangladeshi farmers to follow the Good Agriculture Practices of safe and quality food for healthy future generation.

ABBREVIATIONS USED

MRLs = Maximum Residue Limits; GC-ECD = gas chromatography with electron capture detector; i. d. = internal diameter; ND = not detected; SD= standard deviation; RSD = relative standard deviation; RT = retention time; BDL = Below Detection Limit, Cp = Cypermethrin.

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