COAT PROTEIN GENE OF NEW ISOLATE OF CUCUMBER MOSAIC VIRUS INFECTING BANANA IN EGYPT

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doi: 10.15414/jmbfs.2015.5.2.177-181

ABSTRACT

Banana plants showing typical mosaic and yellow stripes on leaves as symptoms were collected from Assiut Governorate in Egypt. The causal agent was identified as Cucumber mosaic virus (CMV) on the basis of symptoms, transmission, serology, transmission electron microscopy (TEM) and reverse transcription polymerase chain reaction (RT-PCR). Coat protein (CP) gene was amplified using gene specific primer, followed by cloning into desired cloning vector and sequencing. In this study the CMV was transmitted into propagation host either by aphid or mechanically. The transmission was confirmed through direct antigen coating enzyme linked immunosorbent assay (DAC-ELISA). Analysis of the 120 deduced amino acid sequence of the coat protein gene revealed that the Egyptian isolate of CMV shared from 97.50 to 98.33% with those strains belonging to subgroup IA. The cluster analysis grouped the Egyptian isolate with strains Fny and R8 belonging sub-group IA. It appears that there occurs a high incidence of CMV infecting banana belonging to IA subgroup in most parts of Egypt.

Keywords: Banana, CMV, transmission, CP gene, RT-PCR

INTRODUCTION

Banana is considered the most important fruit in Egypt. Many clones are grown for the export market, but many indigenous cultivars are produced and consumed locally. Bananas are affected by five viruses (Diekmann and Putter 1996) these are Banana bunchy top virus (BBTV) genus Nanaviruses, Banana streak virus (BSV) genus Badnavirus, Cucumber mosaic virus (CMV) genus Geminiviruses, Banana bract mosaic virus (BBBrMV) genus Potyviruses, and Banana mild mosaic virus (BMMV), (Thomas et al., 2000; Gambley and Thomas 2001). Banana mosaic virus has been shown to be widespread in banana and also been noted in monocotyledons and a great number of dicotyledons (Chen et al., 2006). CMV isolates have been divided into two subgroups (named I and II) based on different methods such as: serological data, amino acids of the coat protein, nucleic acid hybridization and nucleotide sequence similarities. Roossinck et al. (1999) suggested further division of subgroup I strains into IA and IB, on the basis of the results of phylogenetic analyses. Previously, Daniels (1992) had proposed that CMV subgroup I be subdivided into Ia and Ib on the basis of host reaction and dsRNA patterns. Aim of our study illustrates detection and cloning of CP gene of CMV isolate to correctly identify the disease and to assess similarity or variability among isolates of CMV infecting banana in Egypt as well as other CMV isolates from rest of the world.

MATERIAL AND METHODS

The present investigations were undertaken to identify the causal virus based on biological, serological and molecular testing. The research work has been carried out at Botany Department, Faculty of Sciences, Al-Azhar University, Assiut and Genetic Department, Faculty of Agriculture, Sohag University, Sohag, Egypt.

Plant samples

Banana plants (Musa sp. Cv. Williams) showing leaf mosaic, yellow stripes, leaf distortion and stunting of plant were collected from some farms in Assiut, Egypt. Virus cultures were maintained under insect proof glass house condition.

Viral isolation and mechanical transmission

Naturally infected banana leaves were used for preparing inoculum with 0.1 M phosphate buffer pH 7.0. Pure culture was maintained by obtaining single lesion from Chenopodium amaranticolor Coste & Reyn and inoculating it on to Nicotiana glutinosa L. The virus was further multiplied on the hosts such as N. tabacum cv. White Burly and healthy cv. Williams banana virus-free plants. These hosts were grown in an insect proof glass house. The inoculated plants were serologically tested using CMV antiserum.
Aphid transmission

Individuals of the aphids *Aphis gossypii* collected from cotton plants and *A. craccivora* collected from cowpea. Insects of aphids *Aphis gossypii* and *A. craccivora*. New generation of aphids were transferred to other healthy plants of cotton and cowpea growing in insect proof cages and the aphids were left for reproduction. Several virus free adults of each of the above mentioned aphids were starved for two hours.

They were allowed to feed for 20 min (acquisition period) on *N. tabacum* plants. After that the aphids were transferred to healthy *Nicotiana glutinosa* and *Cucumis sativus* allowed to feed for 10-15 min (inoculation period), and then the insects were sprayed by insecticide (Malathion). The plants were maintained in insect proof cage at 25-30°C for 30 days and the plants were inspected daily for symptoms development. The aphid inoculated plants were serologically tested using CMV antisera.

CMV detection by enzyme-linked immunosorbsent assay (ELISA)

Direct antigen coating-ELISA (DAC-ELISA) test was performed as described by Clark and Joseph (1984) to test the presence of CMV in collected or inoculated plants using CMV antisera. One g. of each leaf sample collected and ground in liquid nitrogen then, one ml of 0.2M potassium phosphate buffer with 0.5% sodium sulphate and 1% polyvinyl pyrolidone (PVP) was added to each sample. The sample squeezed gently with absorbent cotton and equal volume of chloroform was added and vortexed thoroughly. The homogenate was centrifuged at 5000 rpm for 5 min and the supernatant was used for coating the ELISA plates.

Cucumber Mosaic Virus antisera was provided by Dr. Rehab Dawood, Agriculture Reaearch Center, Dokki, Giza, Egypt was used for serodiagnosis study. Plates were coated with 200 μl of prepared antigen sample and incubated at 37°C for 1 hr or over night at 4°C. After incubation, the well’s contents were discarded and washed with PBS-T buffer thrice (flooding with PBS-T for 3 min every time). The blocking solution was added (200 μl / well) and incubated for 1 hr at 37°C. After washing the plate (thrice with PBS-T), 200μl of diluted rabbit antisera (1: 500) was added in the wells and incubated at 37°C for 1½ hrs. The plate was washed and 200 μl of enzyme conjugate (anti- rabbit alkaline phosphatase) was added and incubated at 37°C for 1½ hrs. After incubation, the plates were washed with PBS-T. After final washing, 200μl of substrate (p-Nitro phenyl phosphate-0.6 mg/ml) was added and allowed for color development at room temperature by avoiding exposure to light. Then the reaction was stopped by adding 3M NaOH and the color intensity was read in ELISA reader at 405 nm.

Electron microscopic study

Banana leaves showing mosaic and yellow stripes resulted from mechanical transmission were collected for transmission electron microscopic studies. Samples were fixed in modified Karnovsky fluid’s (David et al., 1973) buffered with 0.1 M Sodium Phosphate Buffer (pH 7.4). Fixation was done for 2 hrs in 1% Osmium tetroxide in the same buffer at 40°C. After several washes in 0.1 M Sodium Phosphate Buffer, the specimens were dehydrated in graded acetone solution and embedded in Epon 812. Ultrathin section of 60-80 nm thickness were cut using an ultra cut ultramicrotome and the sections were stained in alcoholic uracil acetic acid (10 min) and lead citrate (10 min), before examining corresponding grids under transmission electron microscope operated at 60-80 Kv.

RNA extraction and RT-PCR of coat protein (CP) gene

Approximately 100 mg infected dry leaves of banana was frozen in liquid nitrogen and ground to powder with a mortar and pestle. Total RNA was extracted with 1 ml TRIzol reagent, according to the instructions of the supplier using the method of Girin et al. (2007). After ethanol precipitation, the RNA was resuspended in 30 μl RNase free water and treated with RNase free DNase (Ambion), which was then inactivated according to the instructions of the supplier. Approximately 5 μg of total RNA were reverse transcribed using random hexamers (in vitro) and ReverTra-α reverse transcriptase without RNaseH activity (Fermentas) in a final volume of 20 μl. For RT-PCR, template was provided as 2 μl of a 50-fold dilution of cDNA in water in a total reaction volume of 20 μl. The CMV specific primers 5′-GCC GTA AGC TGG ATG GAC- AA-3′ and 5′-TAT GAT AAG AAG CTT GTT TCG CG-3′ designed by Wylie et al. (1993) were used for amplification a part of the CP gene and part of the 3′-noncoding region of CMV RNA3 and were designed to produce amplicons of different sizes to distinguish CMV isolates belonging to subgroups I or II (Singh et al., 2010). Amplification was done using Phusion DNA polymerase (Promega), with initial denaturation at 98°C for 2 min followed by 35 thermal cycles of 98°C for 15 s, 60°C for 30 s, and 72°C for 10 s and followed by final extension at 72°C for 10 min.

Cloning and sequencing of CP gene

A cDNA fragment of about 487 bp was cloned into pCR-Blunt-II-TOPO cloning vector (3519 bp, invitrogen), according to the instructions of the supplier. Colony PCR from selected clones were carried out using the same specific primers of RT-PCR for the conformation of insertion product of CP gene. The DNA sequencing of 3 independent positive clones was carried out using the Sanger method with fluorescent chain-terminating inhibitors di-deoxynucleotide (Smith et al., 1986). The BigDye® Terminator v3.1 Cycle Sequencing Kit (Life Technologies) was used to sequence plasmid DNA with the universal primers M13 forward and reverse on a Applied Biosystems 3100 System according to the indications of the supplier.

Sequences analysis

The partial coat protein (CP) gene sequence of Egypt isolate was submitted to National Center of Biotechnology Institute (NCBI) Database (KP324751). Multiple alignments of amino acid sequences and phylogenetic tree were obtained by using the default options of the Clustal W program version 2.0 (Larkin et al., 2007). Search homologies with proteins from the BLAST basic local alignment search tool of NCBI was done with the FASTA (Pearson and Lipman 1988), and TBLASTN program 2.2.22+ (Altschul et al., 1990). The partial sequence of amino acids corresponding to 363 bp fragment of coding sequences (CDS) of CP gene was compared with some other corresponding known CMV isolates as following: Tn (Y16926), C7-2 (D4279), IA (AB412294), Lx (U20219), N9 (D28780), Fny (D10538), R284ALM183119, Y (D12499), Leg (D16405), LS (A127976), Q (M21464) and Trk7 (L15336).

RESULTS AND DISCUSSION

Transmission and serological diagnosis

Twenty five plants of *Chenopodium amaranticolor, Nicotiana. tabacum, N. glutinosa, Cucumis sativus* and *Banana* were sap inoculated by using naturally infected banana extract (Figure 1A) and carborundum powder. The results revealed that tobacco plants expressed the CMV symptom after 22 days, while banana plants after 48 days of inoculation. Transmission of CMV causes severe mosaic and leaf deformations in inoculated plants, which was confirmed through DAC-ELISA. On another hand, *C. amaranticolor* expressed local lesion symptoms. Vector transmission studies using two aphid species *A. craccivora* and *A. gossypii*, both species are efficiently transmitted the virus in a non-persistent manner within 20 min of acquisition and inoculation of 10 to 15 min. Forty eight plants of Banana and Tobacco were inoculated with infective *A. gossypii* and *A. craccivora*. Inoculated plants showed the symptoms of CMV after 13-22 days of inoculation. These results revealed that the CMV is aphid transmissible and agreed with the data recorded by Rao (1980) who reported. Successful transmission of CMV to test plants by *A. gossypii* in this respect, Mali and Rajegore (1980) who indicated that CMV isolated from banana was transmitted by *A. craccivora* to healthy plants. His method is very efficient even in detecting symptom less plants. The results revealed that positive control recorded a mean absorbance of 0.382. The negative control recorded a mean absorbance of 0.030. The present investigation results has clearly revealed the occurrence and association of cucumber mosaic virus with banana chlorosis was identified based on mechanical transmission, insect vector transmission, host range, serology DAC-ELISA detection. The virus was able to transmit mechanically on the plants used in the study very efficiently and produced the systemic and local symptoms characteristic of CMV. In our present study both the aphid species (*A. gossypii* and *A. craccivora*) transmitted the virus in non persistent manner. Serology is the most reliable and quick method of detection of CMV infection in banana (Rajasulochana et al., 2008). Dheepa and Parannithi (2010) reported that CMV from banana could be transmitted mechanically and by aphid inoculation. CMV has the broadest host range among the plant viruses (Roossinck 2002) and has been reported infecting banana and causes banana chlorosis disease.
The variation in the phylogenetic trees obtained from the nucleotide and amino acid sequences is of course due to the redundancy of the genetic code but may also be due to the fact that only the 3’ coat protein coding region was translated. While the E eg strain identification of banana CMV, in agreement with the previous grouping of CMV strains into subgroups IA, IB and II was based on the phylogeny estimations with full CP open reading frame and rearrangements in the 5’ non-translated region of RNA 3 (Roossinck et al., 1999). In this study the partial 3’ coat protein coding region sequences analyzed and also sufficiently for divided CMV strains into subgroups, these results in agreement with those obtained by Eni et al. (2008).

**Electron microscope studies**

Transmission electron microscopy of infected banana leaves revealed presence of a polyhedral virus particle of ~28-29 nm diameters (Figure IB). The shape and size of the virus particle were identical with those of CMV particle obtained from other sources.

**RT-PCR and cloning of coat protein gene**

PCR amplification of the CP gene using gene specific primers resulted in the amplification of 487 bp fragment (Figure 2). After purification, PCR products were successfully cloned into pCRR-Blunt-II-TOPOR cloning vector. RT-PCR based detection systems significantly improves monitoring and forecasting of banana mosaic epidemics. Data presented in this study clearly indicate that CP region is sufficient to provide a simple and reliable method for detection and strain identification of banana CMV, in agreement with Khan et al. (2011).

**Table 1** Comparisons of nucleotide sequence identities of deduced amino acids between Cucumber mosaic virus isolate (EG-A) and other CMV strains

<table>
<thead>
<tr>
<th>Strain</th>
<th>Sub-group</th>
<th>Amino acid Identity (%)</th>
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<tbody>
<tr>
<td>Tfn</td>
<td>IB</td>
<td>97.50</td>
</tr>
<tr>
<td>C7-2</td>
<td>IB</td>
<td>96.67</td>
</tr>
<tr>
<td>SD</td>
<td>IB</td>
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</tr>
<tr>
<td>IA</td>
<td>IB</td>
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</tr>
<tr>
<td>Tr</td>
<td>IB</td>
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</tr>
<tr>
<td>N9</td>
<td>IB</td>
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</tr>
<tr>
<td>K8</td>
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<tr>
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</tr>
<tr>
<td>Trk7</td>
<td>II</td>
<td>85.00</td>
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**Figure 2** Gel photograph of RT-PCR amplicons: Lane 1, Amplified CMV CP-Eg-A sample, lane 2, negative control and M, 100 bp ladder.

**Sequences analysis**

Analysis of the 120 deduced amino acid sequence of the coat protein gene revealed that the Eg-A strain of CMV had 95.83 to 98.33% homology with subgroups IA and IB strains in this region (Table 1 and Figure 3). The Egyptian isolate shared from 97.50 to 98.33% with those strains belonging to subgroup IA, and from 95.83 to 97.50% with those belonging to subgroup IB. While the Eg-A isolate revealed low level of similarity (85.00 - 87.50%) with those belonging to sub-group II. These results confirmed the placement of Eg-A isolate in subgroup I. The cluster analysis grouped the Egyptian isolate with strains Fny and Ri8 belonging subgroup IA and II belonging subgroup IA. Palukaitis and Zaitlin (1997), and Roossinck (2002) documented that CMV subgroup I is further divided into IA and IB on the basis of sequences and phylogenetic relationships. Strains IA and II of this virus are distributed worldwide, whereas the strains of IB subgroup are mainly restricted to Asia (Roossinck 2002; Koundal et al., 2010). The multiple sequence alignment (Figure 3) showed that the Egyptian isolate differs from all isolates used for comparison in the residue 39 and 58 of the CP which exhibits Serine (S) and Threonine (T), respectively, whereas all other isolates have Alanine (A) at these sites. Two strains (Fny and Ri8) of the four subgroup IA, had 98.33% identity with the Eg-A strain in the 120 amino acids of the coat protein gene and differs with them in these two residues only. In this direction, three out of the six subgroup IB strains used for comparison had three amino acid substitutions at various points (1, 39 and 58) and had 97.5% amino acid identity with the Eg-A strain (Table 1 and Figure 3). It has been reported that substitution of this last amino acid affects symptom expression, inducing chlorosis instead of necrosis in infected tobacco plants (Mochizuki and Ohki, 2011). All the three strains belonging subgroup II used for comparison had 85% and 87.5% amino acid homology with the Eg-A strain and formed separately a distinct clade (Table 1 and Figure 4).

The Shape (B) of the banana mosaic virus shows a polyhedral virus particle of ~28 nm diameters (Figure 1). Electron microscope studies (see arrows). Transmission electron microscopy of infected banana leaves revealed presence of a polyhedral virus particle of ~28-29 nm diameters (Figure 1B). The shape and size of the virus particle were identical with those of CMV particle obtained from other sources.

![Figure 1](image1.png)

**Figure 1** Cucumber Mosaic Virus (CMV) infected banana showed mosaic and yellow stripes in leaves (A) as naturally symptoms and Electron micrograph of a thin section of infected leaf sample of banana showing aggregates viral particle (B) (see arrows).

![Figure 2](image2.png)

![Table 1](image3.png)
CONCLUSION

Cucumber mosaic disease caused by CMV has attained a serious status in most of the banana growing areas. Importance of the disease stems from the fact that it is responsible for losses to 40%. Aim of our study illustrates detection and cloning of CP gene of CMV isolate to correctly identify the disease and to assess similarity or variability among isolates of CMV infecting banana in Egypt as well as other CMV isolates from rest of the world. For this, coat protein gene of CMV isolate collected from Assiut Governorate was cloned and sequenced. High degree of nucleotide sequence identity (97-98%) between isolates belonging to subgroup IA. These results were confirmed by electron microscopy and transmission. Transmission was confirmed through DAC-ELISA technique. It has been concluded that the transmission of CMV was more successful to *N. tabacum* and *N. glutinosa* by mechanically and through aphids.

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