

# SEQUENCE ANALYSIS OF COAT PROTEIN GENE AND 3' NON-CODING REGION OF SWEET POTATO FEATHERY MOTTLE VIRUS (SPFMV) ISOLATED FROM EGYPT

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**B**efore the advances of data obtained by sequencing of potyviruses, species and strains of viruses were distinguished on the bases of host range, symptoms on indicator plants and serological relationships. Currently, availability of coat protein and the full genome sequences established the criteria that can be used in particular to distinguish sequences representing closely-related virus species from those strains of the same species (Shukla and Ward, 1988; Adams *et al.*, 2005). Additionally, comparisons showed that nucleotide sequences of the 3' untranslated region of the potyviral genome can serve as an aid to identify and classify potyviruses (Frenkel *et al.*, 1989). Increased knowledge of conserved viral sequences allowed the design of oligonucleotide primers for their use in reverse transcriptional polymerase chain reaction (RT-PCR) amplification and rapid identification of uncharacterized potyviruses (Colint *et al.*, 1994; Langeveld *et al.*, 1991; Pappu *et al.*, 1993).

Sweet potato feathery mottle virus (SPFMV) belongs to potyviruses, the largest group of plant positive sense

single stranded RNA viruses (Ward and Shukla, 1991). It has long been recognized to infect sweet potatoes worldwide (Moyer and Salazar, 1989). The virus is transmitted mechanically and by several genera of aphids including the cotton aphid (*Aphis gossypii*) and the green peach aphid (*Myzus persicae*) in a non-persistent manner (Brunt *et al.*, 1996). SPFMV is a major component of sweet potato virus disease that is drastically affecting the yield of sweet potatoes worldwide (Gibson *et al.*, 1998; Gibson and Aritua, 2000). Several isolates and strains of SPFMV have been characterized in different parts of the world, the most important ones being the ordinary strain "O" (Usugi *et al.*, 1991), the russet crack strain "RC" (Moyer and Salazar, 1989), the common strain "C" (Cali and Moyer, 1981) and severe strain "S" (Mori *et al.*, 1995). The first full-length sequence of SPFMV was published earlier (Sakai *et al.*, 1997) and showed that it is the largest genome among potyviruses with a MW of 10,820 nucleotides excluding the poly (A) tail, which is incapsidated into a 35 kDa coat protein subunits. Several attempts were carried out to identify SPFMV on the basis of RT-PCR and sequence analysis

(Colinet *et al.*, 1998; Ryu *et al.*, 1998; Hanada *et al.*, 2000; Sakai, 2000). Recently, several partial sequences of SPFMV coat protein gene and 3' non coding region from different geographic areas are made available on the databases (NCIB database).

In this study, we present the sequence of coat protein gene and the 3' non-coding region of SPFMV isolated from Egypt with its homology to the already available sequences on the database.

## MATERIALS AND METHODS

Sweet potato (*Ipomoea batatas* var. Abees) carrying the Egyptian isolate of SPFMV (Ashoub *et al.*, 2007) was used to isolate RNA and amplifying the coat protein gene and the 3'end of the virus as following:

Total RNA of SPFMV-sweet potato infected plants was extracted following the method described by Ashoub *et al.* (2006) with further purification using Promega SV-40 total RNA extraction kit (Promega, USA) according to the manufacturer's recommendations.

One  $\mu\text{g}$  RNA was denatured at 70°C for 5 min. in the presence of 100 pmol primer Oligo-dT (5'-TTTTTTTTTTTTTTTTTYI-3'), where Y is A, G or C residue and I is an inosin residue. Samples were chilled on ice for 2 min. RNA was reverse-transcribed for 1 h. at 42°C in the presence of 1  $\times$  reverse

transcriptase buffer, 20 mM dNTPS, and 200 units M-MuLV reverse transcriptase (Promega, USA). After incubation, PCR was carried out using 5  $\mu\text{l}$  cDNA in 1  $\times$  PCR buffer, 50 pmol primer Oligo-dT, 10 pmol primer P-SPFMV-CP3'-F (5'-CTTCAGTGACGTTGCTGAAGC-3'), 1.5 mM  $\text{MgCl}_2$ , 10 mM dNTPs, and 2 units *Taq* DNA polymerase (Fermentas, USA). Samples were subjected to 30 cycles of PCR with 15 sec. of denaturing at 94°C, 15 sec. of annealing at 50°C, and 1 min. of extension at 72°C. The series of cycles was preceded by 3 min. initial denaturing at 94°C. Five  $\mu\text{l}$  of the PCR products were analysed on a 1% agarose gel in TAE buffer, visualized by ethidium bromide staining and photographed (Ausubel *et al.*, 1995). PCR product was purified using Wizard SV gel and PCR clean-up system (Promega, USA) following the manufacturer's instructions. After purification, PCR was introduced into pGEM-T Easy vector system (Promega, USA) following the manufacturer instructions to produce pSPFMV-CP3'. Vector was transformed into *E. coli* JM109. Positives were selected and plasmid DNA was isolated using Wizard plus SV minipreps DNA purification System (Promega, USA) following the manufacturer's instructions.

To amplify the rest of coat protein gene, the same described above approach was carried out except that primer P-SPFMV-CP5'-R (5'-AAGAGGTTATGTATATTTCTAGTAA-3') was used to generate the cDNA in combination with SPFMV-CP5'-F (5'-

GAGTACAACCTAGCGCTGACGATC TCAGC-3') to generate the PCR product. Additionally, the concentration of PSPFMV-CP-R and SPFMV-CP-F was 10 pmol each for the PCR reaction and the annealing temperature was 60°C. After ligation as mentioned above, the generated vector was named pSPFMV-CP5`.

Vectors were subjected to sequence analysis using T7 and SP6 promoter primers (Promega, USA), sequence reactions were carried out using the 310 sequencer (Applied bio systems) and analyzed using DNA star program.

## RESULTS AND DISCUSSION

In this study, the coat protein gene and the 3`non coding region were reverse transcribed, amplified as PCR products, cloned and subjected to sequence analysis in two steps. In the first step, reverse transcription reaction was carried out based on the Oligo-d-T primer since all potyviruses have a polyadenilation 3' terminal. In PCR amplification reaction, a forward primer was designed from a conserved region for all SPFMV published in the database of NCIB (Ashoub *et al.*, 2007). The amplified PCR product (Fig. 1-A) has a molecular weight size of 550 bp. When pSPFMV-CP3' was subjected to sequencing reaction, it read 544 nucleotides, excluding the polyadinitation tail and representing 222 nucleotides of 3' non coding region of the virus in addition to 322 nucleotides of the coat protein gene C-terminus. Comparing SPFMV sequence obtained from the

Egyptian isolate with sequences published on the database, it revealed that the similarity was 98% to the South Africa strain C (accession number AY459601). For the second step to amplify the rest of the coat protein gene, primer SPFMV-CP5' -F was synthesized based on the sequence of the South Africa strain C located 104 nucleotides upstream of the coat protein gene start codon combined with primer PSPFMV-CP5`-R designed on the bases of the available sequence of SPFMV Egyptian isolate obtained from the first step. The amplified PCR product (Fig. 1-B) has a molecular weight of 1024 bp. The sequence of pSPFMV-CP5', excluding the 104 nucleotides upstream of the CP gene, in combination with pSPFMV-CP3' represent the complete coat protein gene sequence of the SPFMV Egyptian isolate in addition to the 3' non-coding region with a molecular size of 1164 nucleotides (Fig. 2-A). The comparison analysis with other published SPFMV coat protein gene and 3`non coding region sequences indicated the highest similarity (98%) of the Egyptian isolate with South Africa strain C. Strains used in the comparison, their accession number, geographical location and degree of similarity to the Egyptian isolate are indicated in Table (1). Results of the phylogenetic tree showed the similarity between the Egyptian isolate and the other isolates used in the comparison, which indicated that the Egyptian isolate falls within the group of strain C of SPFMV (Fig. 3). Deduced amino acid analysis of the coat protein gene of SPFMV from Egypt indicated that the protein codes for

313 amino acids with a calculated molecular weight of 35 kDa (Fig. 2-B) which is in agreement with Sakai *et al.* (1997). It has 98.4% similarity with the South Africa strain C when both were compared based on their deduced amino acids. Moreover, the 3' non coding region of the Egyptian isolate was 98.2% similar to the South Africa isolates, while the coat protein gene was 98% similar to the South Africa isolate C.

The availability of sequence information for the coat protein gene is a necessity for downstream applications like molecular and serological diagnostic tools and improving sweet potato crop by introducing resistance *via* established gene transformation strategies.

### SUMMARY

The coat protein gene and the 3' non-coding region of sweet potato feathery mottle virus, SPFMV, isolated from Egypt was subjected to reverse transcriptional polymerase chain reaction, RT-PCR, cloned and applied to sequence analysis. The 1164 nucleotides sequenced representing the full-length coat protein gene were found to code for 330 amino acids with MW of 35 kDa. In addition, 222 nucleotides of 3' non-coding region excluding the poly-A tail were sequenced. Data comparison to the published sequences revealed that the nucleic acid sequence of the Egyptian SPFMV isolate has 98% homology with the common strain, C, isolated from South Africa and 98.4% homology of the deduced amino acids. The phylogenetic analysis indicated

that the Egyptian isolate occurs within the C strains of identified SPFMV.

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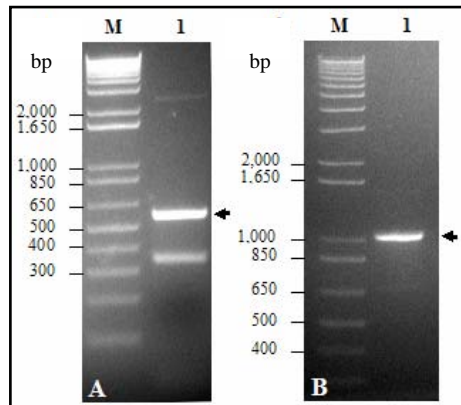
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Table (1): Accession numbers of SPFMV strains found in the NCBI database and their percentage of similarity in comparison to the Egyptian isolate.

| Accession Number | SPFMV Isolates and Strains      | Similarity to SPFMV from Egypt |
|------------------|---------------------------------|--------------------------------|
| AY459601         | South Africa strain C           | 98%                            |
| AY459594         | Kenya: Kisumu strain C          | 95.7%                          |
| AY459596         | Uganda: Namulonge strain C      | 94.5%                          |
| AY459591         | Kenya: Kakamega strain C        | 93.8%                          |
| S43451           | USA: Strain C                   | 88.5%                          |
| AY459592         | Kenya: Kakamega strain EA       | 74.1%                          |
| AY459600         | Spain, Canary Islands strain EA | 74.1%                          |
| AF015541         | Korean strain 2                 | 74%                            |
| AY459599         | Portugal strain EA              | 74%                            |
| AY459593         | Kenya: Kisii strain EA          | 73.9%                          |
| AY459597         | Madagascar strain EA            | 73.8%                          |
| D16664           | ordinary strain, SPFMV-O        | 73.7%                          |
| AY459598         | Tanzania strain RC              | 73.7%                          |
| D38543           | Strain Severe                   | 73.5%                          |
| AY459602         | China strain RC                 | 73.3%                          |
| AF015540         | Korean strain 1                 | 73.2%                          |
| AY459595         | Uganda: Arua strain O           | 73.2%                          |
| S43450           | Strain RC                       | 73.2%                          |

Fig. (1): RT-PCR amplifications of the SPFMV coat protein gene and 3' non-coding region. Amplification of the 3' part using Oligo-dT as reverse primer and P-SPFMV-CP3'-F as forward primer (A), and amplification of the 5' part using PSPFMV-CP5'-R as reverse primer and P-SPFMV-CP5'-F as forward primer (B). PCR products are indicated by an arrow in comparison to the molecular weight marker, M.



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TCTGGTAACCCCCCTGAATTTAAAGATGCAGGTGCGAACCCACCAGCACCAAAGCCAAAAG
GGCCATATGTAGCACCAGAAATCACAGAAGTTACTGATCCCCGAGGATCCCAAACAAGCTGC
GCTCCGCGAAGCTAGACAGAAAACAACCTGCTGTACACCCGAATCATACGGTAGAGATACA
GGTGAGAAAACCTATGCGCTCTGTTTTCCACCAAAGGGTGAAAGACAAAGGATGTCAATGTTG
GTACGACAGGTACATTTTTTAGTACCACGAGTTAAGCTTCATACTAGTAAAATGCGCCAACC
GAGAGTCAATGGAGTCTCCGTAGTAAACTTACAACACCTTGCAACCTATGAACCTGAGCAA
CATAACATTGGGAATACACGCTCAACTCAGGAACAGTTTTCAAGCATGGTACGAAGGTGTCA
AGGGTGATTATGGTGTGATGACGCTGGAATGGCAATCTTGTTGAATGGATTGATGGTATG
GTGCATAGAGAATGGAACATCTCCAAATATAAATGGCGTTTTGGACGATGATGGACGGGGAT
GAACAAGTGACGTACCCCATAAAGCCATTACTGGATCATGCAGTGCCTACTTTTAGGCAAAA
TCATGACACACTTCAGTGACGTTGCTGAAGCGTATATAGAGATGAGGAATCGCACCAAGGC
ATATATGCCTAGGTATGGATTACAACGTAATTTGACTGATATGAGTCTTGCGCGATATGCA
TTTGATTTCTATGAGCTGCACTCAACAACACCTACACGTGCTAAAGAAGCACACATGCAGA
TGAAAGCAGCTGCACTTAAGAATGCGCATAATCGGTTGTTTGGTTTGGATGGAAACGTCTC
CACGCAAGAAGAGGATACGGAGAGGCACACTGCGACTGATGTTACTAGAAATATACATAAC
CTCTTAGGAATGAGGGGTGTGCACTAGGTGAAACTTTGCACTGTATTTATTTACTTATGTG
GTTTTTAGTATGCCTTTATTTAAATTCGTGTTCTTCAGTCCCGACAGAAATGGTTGGGTGT
ATCGACAAAGTGGGCTTTTAGCCTGGTCCATACACTTGAGAAGTTTCTGGTCTATTACGTA
TCATAAGGGACTCTTAAAGTGGAGTACCTCGTAAGAAAAGCCTTTTTGGTTCGTGATC
GAGCA (N)

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Fig. (2A): SPFMV Egyptian isolate coat protein and 3' non coding region nucleotide sequence. The TAG stop codon of the coat protein gene is underlined and the (N) indicates an unidentified number of the adenosine residue at the 3' poly-A tail.

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SGNPPEFKDAGANPPAPKPKGPYVAPEITEVTDPEDPKQAALREARQKQPAVTPESYGRDT
GEKPMRSVSPQRVKDKDQVNVGTTGTFLVPRVKLHTSKMRQPRVNGVSVVNLQHLATYEPEQ
HNIGNTRSTQEQQAWYEGVKGDYGVDDAGMAILLNGLMVWCIENGTSPNINGVWTTMMDGD
EQVTYPIKPLLDHAVPTFRQIMTHFSDVAEAYIEMRNRTKAYMPRYGLQRNLTDMSLARYA
FDFYELHSTTPTRAKEAHMQMKAALKNAHNRLFGLDGNVSTQEEDTERHTATDVTRNIHN
LLGMRGVH*

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Fig. (2B): Translation of SPFMV coat protein gene. The asterisk indicates the termination codon of the coat protein.

Fig. (3): Phylogenetic tree of SPFMV isolates used in this study and their relation to the Egyptian isolate.

