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Comparison of N-Terminal Region of Coat Protein in **Zucchini Yellow Mosaic Potyvirus Isolates**

I. TÓBIÁS¹, L. TZEKOVA², L. PALKOVICS² and E. BALÁZS²

¹Plant Protection Institute of Hungarian Academy of Sciences, Budapest, Hungary ²Agricultural Biotechnology Center, Gödöllő, Hungary

Zucchini yellow mosaic potyvirus (ZYMV) was isolated in Hungary from cucumber for the first time in 1995 and now it is widespread causing devastating epidemics in cucurbit crops.

Four isolates were chosen from different cucurbit plants collected in 1995 and 1999 and molecular variability of N-terminal region of coat protein was studied. Hungarian isolates show lower molecular variability within themselves than with ZYMV isolates from other geographical origins. Hungarian isolates have amino acid sequences specific only to them that suggest common ancestor.

Key words: zucchini yellow mosaic virus, potyvirus.

Zucchini yellow mosaic potyvirus (ZYMV) is widely distributed potyvirus that causes important losses in a range of cucurbit crops. The apparently rapid spread of this virus in many parts of the world suggesting a very efficient transmission from plant to plant by several aphid species in a non-persistent manner (Lisa and Lecoq, 1984) and a long distance distribution via infected seeds (Schrijnwerkers et al., 1991).

ZYMV, first described in Italy (Lisa et al., 1981), belongs to the potyviruses, the largest group of plant viruses, characterized by a monopartite, positive-sense, singlestranded RNA encapsidated in flexuous, filamentous particles. The RNA is translated into a single polyprotein cleaved by three viral proteases (Reichmann et al., 1992).

In Hungary zucchini yellow mosaic potyvirus first observed in 1995 causing severe epidemics in cucurbit-growing regions (Tóbiás et al., 1996). Some zucchini yellow mosaic potyvirus were isolated and compared on test plants and ZYMV-10 strain was chosen for further characterization (Tóbiás et al., 1998).

In the past four years, many part of the country cucurbit crops were subject to severe losses due to zucchini yellow mosaic potyvirus. Several isolates have been identified among them some capable of overcoming resistance to the dominant Zym gene (Tóbiás et al., 1998). This emphasize the need for additional sources of resistance derived by either classical breeding or pathogen-derived resistance. Coat protein (CP) mediated resistance could provide an alternative control against very destructive ZYMV. ZYMV CP transformation of melon is in progress. Since the effectiveness of CP resistance would depend on serological relatedness of the attacking virus the variability of ZYMV occurring in Hungary has to be studied. In fact, the N terminus is the only region in the entire coat protein that is unique to potyvirus and contains virus-specific and virus strainspecific epitopes (Shukla et al., 1991). Kundu et al. (1998) compared the coat protein epitopes of ZYMV and found that epitopes in the N-terminal region could differentiate among different ZYMV isolates.

The objectives of this research were to compare sequence data of N-terminal region of CP to characterize the variability of zucchini yellow mosaic potyvirus in Hungarian ecosystem.

Virus isolates were from different cucurbit plants, from different region and different time. ZYMV-10 strain originated from cucumber (Békésszentandrás, 1995), ZYMV-sz6 from oilseed squash (Csabacsüd, 1998), ZYMV-sz3 from melon (Szarvas, 1999) and ZYMV-f1 from zucchini (Fertőd, 1999). The isolates were propagated by mechanical transmission in *Cucumis sativus* cv. Budai csemege. ZYMV-10 strain was previously characterized and partially sequenced (Tóbiás et al., 1998). ZYMV-sz6, ZYMV-sz3 and ZYMV-f1 isolates were PCR amplified and cloned. Total RNA was extracted from leaves with obvious sign of disease by the method White and Kaper (1989).

Degenerate primers ZYMV CP1 (5' GTAATGCTAACCATGGGCACTCAG 3') ZYMV CP3 (5' GGGGATCCGACCTACCCTTTACTG 3') were previously designed to amplify from the second amino acid from the 5' terminal part of the coat protein to the stop codon (first amino acid of CP is constant in ZYMV because of the cleavage site O/S). ZYMV CP3 primed single-stranded cDNA was synthesized from total RNA using the Amersham cDNA synthesis kit. Two microliters from this reaction mixture were used for PCR experiments. Amplification was performed in a volume of 50 µl of PCR buffer (10 mM Tris-HCl pH 9.5, 2.5 mM MgCl₂, 50 mM KCl, 0.1% Triton X-100) containing 100 ng each of dATP, dCTP, dGTP and dTTP, 0.1 nM each upstream and downstream primers and 5 U Taq DNA polymerase (Promega). Forty reaction cycles were perfomed (Perkin Elmer Cetus cycler): template denaturation at 94 °C for 15 s, primer annealing at 55 °C (cycles 1-5) or 60 °C (6-40 cycles) for 30 s and DNA synthesis at 72 °C for 2 min. After electrophoresis in the 1% agarose gel, the amplified DNA fragments were excised and isolated with Prep-A-GeneTM DNA Purification Kit from BIO RAD. Following digestion with Nco I. and BamH I, the PCR products were cloned in the pUC 19 plasmid. Nucleotide sequence of N-terminal region of CP was obtained by using revers primer. Double-stranded DNA sequencing by the dideoxy chain termination method was performed using the T7 DNA polymerase (Pharmacia) according to manufacturer's instruction. Sequence analysis was performed employing University of Wisconsin Genetics Computer Group (GCG) sequence analysis software package version 9.1 and the following Web servers: http://ulrec3.unil.ch/software/BOX_form.html (Boxshade), http://bmerc-www.bu.edu/psa/(PSA), http://www.genome.ad.jp/. Sequence comparisons were performed with EMBL/GenBank databases. Multiple sequence alignments were generated using PILEUP of GCG package. CLUSTALW was used to create an input file for PHYLIP. The tree was displayed using DRAWTREE.

Comparing the amino acids of coat protein of different ZYMV isolates shows that the main differences can be found at the N-terminal part (Fig. 1). Sequences were determined for 129 nucleotides (coding 43 amino acids) of the coat protein coding region. These fragments were chosen for sequencing because the N-terminal part of the coat protein is known to be highly variable and to contain the major virus specific epitopes, due to its localization at the surface of the virion (Shukla et al., 1988).

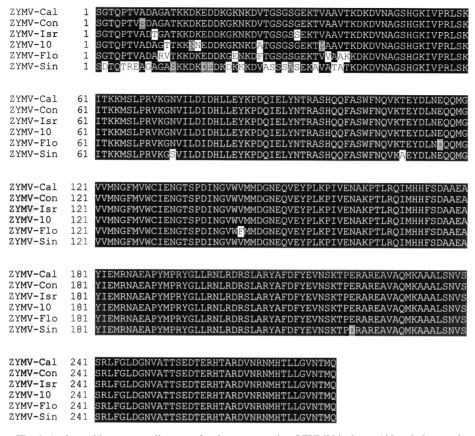


Fig. 1. Amino acid sequence alignment for the coat protein of ZYMV isolates. Abbreviations and respective accession numbers: California strain: ZYMV-Cal: No L 31350, Connecticut strain: ZYMV-Con: No D 00692, Florida strain: ZYMV-Flo: No D 00593, Singapure strain: ZYMV-Sin: No X 62662, Israel strain: ZYMV-Isr: No M 355095 and Hungarian strain: ZYMV-10 AJ251527

ZYMV-f1	1	SGTQPTVADAGATKKN <mark>N</mark> EDDKGKNKD <mark>A</mark> TGSGSGEKTMAAVTKD
ZYMV-sz6	1	SGTQPTVADAGATKKN <mark>N</mark> EDDKGKNKD <mark>A</mark> TGSGSGEKTMAAVTKD
ZYMV-10	1	SGTQPTVADAG <mark>T</mark> TKKN <mark>N</mark> EDDKGKNKD <mark>A</mark> TGSGSGEKTMAAVTKD
ZYMV-sz3	1	SGTQPTVADAGATKKN <mark>N</mark> EDDKGKNRD <mark>A</mark> TGSGSGEKTMA <mark>T</mark> VTKD
ZYMV-Cal	1	SGTQPTVADAGATKKDKEDDKGKNKD <mark>V</mark> TGSGSGEKTVAAVTKD
ZYMV-Con	1	SGTQPTVSDAGATKKDKEDDKGKNKD <mark>V</mark> TGSGSGEKTVAAVTKD
ZYMV-Isr	1	SGTQPTVAD <mark>T</mark> GATKKDKEDDKGKNKD <mark>V</mark> TGSGS <mark>S</mark> EKTVAAVTKD
ZYMV-Flo	1	SGTQPTVADA <mark>RV</mark> TKKDKEDDKG <mark>E</mark> NKD <mark>F</mark> TGSGSGEKTV <mark>VA</mark> AKKD
ZYMV-Sin	1	SDTQTREAGAGASKKDKDEDKDKKKDVASSSASEKAVATATKD

Fig. 2. Amino acid sequence alignment for the N terminal of coat protein of ZYMV isolates. See Fig. 1 for abbreviation and accession numbers

Table 1							
Percentage of nucleotid sequence identity of N-terminal region of coat protein of different zucchini							
yellow mosaic potyvirus isolates. See Fig. 1 for abbreviation and accession numbers							

	ZYMV-Cal	ZYMV-Con	ZYMV-Flo	ZYMV-Isr	ZYMV-Sin	ZYMV-sz3	ZYMV-sz6	ZYMV-f1
ZYMV-10	92	92	86	94	70	97	99	99
ZYMV-Cal		98	90	94	72	90	93	93
ZYMV-Con			89	94	72	90	93	93
ZYMV-Flo				88	69	84	86	86
ZYMV-Isr					74	93	95	95
ZYMV-Sin						71	71	71
ZYMV-sz3							98	98
ZYMV-sz6								100

Table 1 shows the sequence identity percentage for N-terminal part of the coat protein of different ZYMV isolates. The Hungarian isolates had nucleotide sequence identities of 97% to 100%. Isolates from Hungary and ZYMV isolates from different parts of the world have sequence similarities ranging 70 to 94%.

Sequence data indicate that the Hungarian isolates share more homology with the Israel strain than with any other sequenced strains. The ZYMV-Sin isolate has the lowest homology to other ZYMV isolates.

Figure 2 shows amino acid sequence identities of N-terminal region of ZYMV isolates.

Previous studies have shown that a tripeptide sequence DAG located near the N terminus of the CP is required for aphid transmission (Atreya et al., 1995). This motif is highly conserved, but the context in which it is found varies considerable for different potyviruses. ZYMV-10 as an exception to other ZYMV isolates next to the DAG motif has T instead of A. The Hungarian isolates at positions 16 and 17 have uniformly asparagin (N) instead of aspartic acid (D) and lysin (K) at position 27 alanin (A) substitutes valin (V) and at 37 valin (V) changes to methionin comparing to other ZYMV strains. Apart from this ZYMV-sz3 isolate has amino acid changes at 15 and 39 position comparing to others. Hungarian isolates have amino acid sequences specific only to them as indicated in Fig. 2. Although Hungarian isolates are originated from different plants (cucumber, melon, oilseed squash and zucchini), from different regions (Békésszentandrás, Csabacsüd, Szarvas, Fertőd) and isolated at different time (1995, 1998 and 1999) they are very similar according to nucleotide and amino acid sequences. If we analyse phylogenetic relationship of N-terminal region of CP (Fig. 3) we can conclude that the Hungarian isolates clearly form a distinct branch in the phylogenetic tree and share more sequence homology among themselves than with the other strains which suggest their common origin. From this data we can predict that ZYMV-10 CP transformed plants propably will show resistance to all ZYMV isolates occurring in Hungary.

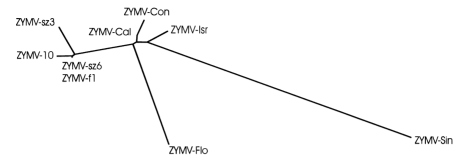


Fig. 3. Phylogenetic tree for N-terminal region of coat protein of ZYMV isolates. See Fig. 1 for abbreviation and accession numbers

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