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Occurrence and diversity of viruses infecting pepper in Serbia

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Abstract: A two-year investigation (2009–2010) of the presence and distribution of pepper (Capsicum annuum L.) viruses in Serbia revealed that viruses occur each year in open-field production. Disease incidence, as estimated by the number of symptomatic plants in the field, highly varied depending on the year and sampling locality. Disease incidence ranged from 20% to 60%. Four viruses: Cucumber mosaic virus (CMV), Potato virus Y (PVY), Alfalfa mosaic virus (AMV) and Pepper mild mottle virus (PMMoV), of which PVY was predominant, were detected by serological testing of pepper samples collected from many localities in Serbia. Molecular detection of PVY was performed based on amplification of a 975 bp fragment in all tested samples, using the specific primers PVYc/PVYd that amplify the gene for P1 protein. The RT-PCR products derived from the four isolates (PL-28-09, PL-15-09, PL-3-10, PL-108-10) of PVY were sequenced (KC288142, KC288143, KC288144, and KC288144, respectively) and compared with the PVY sequences available in GenBank. Sequence analysis, conducted with MEGA5 software, revealed 99.8-100% nt identity among the four Serbian PVY isolates from pepper. The sequences of PVY isolates from Serbia share the highest nucleotide and amino acid identity with isolates from Slovenia, Croatia, Germany, and tobbaco isolate from Serbia. All of the four Serbian isolates were clustered in sub-group N-1 with other European isolates of necrotic strains.

Keywords: pepper viruses; *Potato virus* Y; serological assay; RT-PCR; molecular characterization.

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Introduction

Pepper (*Capsicum annuum* L.) is among the most important vegetable crops in Serbia. It is the second most widely grown vegetable crop in Serbia, covering the area of about 20,000 ha of open-field and protected crops, with an average yield of 8.3 t/ha. In Serbia, different types of pepper are grown for fresh consumption, processing and the production of spices. The main pepper production region, about 80% of the national production, is concentrated in the Vojvodina Province.

Viral diseases constitute the major limiting factor in pepper production, affecting both yield and quality of pepper fruits (Nono-Womdin, 2001). They caused economic losses of 15 billion dollars annually on worldwide basis (Van Fanbing 1999), particularly in tropical and semitropical regions, but great losses have also been reported in temperate regions (Hull and Davies, 1992). Sixty-eight viruses have been reported to cause damages in pepper crops (Pernezny *et al.*, 2003), among which the most significant and widespread pepper viruses in the Mediterranean Basin are aphid-transmitted, *Potato virus* Y (PVY), *Cucumber mosaic virus* (CMV) and *Alfalfa mosaic virus* (AMV), the thrips-transmitted *Tomato spotted wilt orthotospovirus* (TSWV), as well as seed- or mechanically transmitted tobamoviruses, *Pepper mild mottle virus* (PMMoV), *Tobacco mosaic virus* (TMV), *Tomato mosaic virus* (ToMV), and *Tobacco mild green mosaic virus* (TMGMV) (Moury and Verdin, 2012).

Potato virus Y (PVY: genus, *Potyvirus*, family *Potyviridae*) is dispersed worldwide and it is one of the most common and destructive viruses infecting potato, tomato, tobacco and pepper. PVY strains have been classified into five groups: PVY^{O} (ordinary strain), PVY^{C} (common strain), PVY^{N} (necrotic strain), PVY^{Z} and PVY^{E} based on reaction in potato plant carrying resistance genes *Ny*, *Nc*, and *Nz*, as well as tobacco symptoms (Chikh-Ali *et al.*, 2013). In addition, new recombinant genotype/strains have been identified based on sequence analysis, including PVY^{NTN} and PVY^{NWI} , which have mosaic genomes built of fragments of PVY^{N} and PVY^{O} sequences (Ali *et al.*, 2010). PVY^{NTN} and PVY^{NWI} were first found in Hungary and Poland, respectively, but within a short period of time they have become common in potato fields (Ali *et al.*, 2007). Of the 26 species of the family *Aphididae*, *Myzus persicae* is the most efficient PVY vector (Katis and Gibson, 1985; Woodford, 1992). The virus is not seed-borne (Jones *et al.*, 2014).

Despite the importance of pepper in Serbia, only local and limited information is available about the presence and incidence of viruses (Mijatović *et al.*, 2007; Krstić and Bulajić, 2008). In recent years, an extremely high percentage of early virus infections of pepper have caused severe plant stunting, and leaf and fruit malformation was observed in many pepper crops in the Vojvodina Province (Petrović *et al.*, 2010). A two-year survey was conducted in order to give an insight into the occurrence and distribution of viruses infecting

pepper crops in Serbia, by evaluating their relative incidence and potential importance using DAS-ELISA test. This study also focused on determining the presence of PVY using molecular testing and establishing the genetic relationship of its isolates originating from Serbia with those from other parts of the world.

Materials and Methods

Sample collection

During 2009 and 2010, from June to August, research was conducted to investigate the presence and distribution of pepper viruses in Serbia. A total of 365 samples were collected from open-field production. Samples comprising leaves from plants with symptoms resembling those of virus infection were randomly collected after the visual inspection of 31 different localities. Each sample was placed in a plastic bag and stored at 4°C until testing by ELISA or RT-PCR.

Serological assay

Serological testing was performed by double-antibody sandwich (DAS-ELISA) kits using antisera for detection of 9 economically significant pepper viruses: CMV, PVY, TSWV, AMV, TMV, PMMoV, *Potato virus X* (PVX), *Pepper mottle virus* (PepMoV) and *Pepper veinal mottle virus* (PVMV) according to the manufacture's manual (Loewe Biochemica, Sauerlach, Germany). Extracts from fresh leaves were ground in extraction buffer in a ratio 1:10 (w/v). After addition of the substrate (1 mg/ml of p-nitrophenyl phosphate), the plates were incubated at room temperature for 2 hours and the extinction was measured at 405 nm (A405) using an ELISA plate reader (Multiscan Ascent, Finland). Samples were considered to be positive when the absorbance values were two times the absorbance values of negative controls. Commercial positive and negative controls (Loewe), as well as healthy pepper leaves were included in each ELISA for validation of the test results.

RT-PCR detection and sequence analysis

Presence of PVY in four selected pepper samples was further confirmed by conventional reverse transcription (RT)-PCR. Total RNAs were extracted from 100 mg of freeze-dried leaves of naturally infected plants using RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. RT-PCR was performed using the One-Step RT-PCR Kit (Qiagen) with the PVY specific primer pair PVYc/PVYd (Marie-Jeane Tordo *et al.*, 1995; Glais *et al.*, 1996), yielding a 975-bp fragment corresponding to the partial gene for P1 protein.

The RT-PCR reaction mixture included 400 Mm each of the four dNTPs, 0.6 μ M of the viral sense and complementary sense primer, and 1 μ l extracted RNA in a final volume of 25 μ l. Amplifications were performed in a thermal cycler (Biometra, T-1 Thermocycler) with the following cycling parameters: reverse transcription at 50°C for 30 min and an initial PCR denaturation step at 95°C for 15 min, followed by 35 cycles consisting of a denaturation step of 60 s at 94°C, primer annealing for 60 s at 57°C, and extension for 60 s at 72°C. The final extension was performed at 72°C for 10 min. Total RNAs obtained from a Serbian PVY isolate D7-06 from tobacco (GQ290476) and a healthy pepper plant were used as the positive and the negative control, respectively.

The products amplified from the selected isolates were sequenced in both directions, using the same primers directly after purification with a QIAquick PCR Purification Kit (Oiagen). The nucleotide sequences of the amplification products were deposited in GenBank database and they were assigned accession numbers. Sequences of Serbian virus isolates were compared with the previously PVY reported isolates available in the GenBank (http://www.ncbi.nlm.nih.gov/BLAST/) using the ClustalW program (Thompson et al. 1994) and MEGA5 software (Tamura et al., 2011). A p-distance model was applied for nucleotide (nt) and deduced amino acid (aa) sequence analyses and the divergence of selected sequences was calculated using sequences trimmed to the length of the shortest fragment.

Phylogenetic relationship

The phylogenetic tree was constructed based on the nucleotide sequences of P1 gene of the PVY isolates generated in this study with other PVY (Table 1) isolates from different genotypes and geographic origins available in GenBank databases using maximum parsimony method in MEGA5 software. Genetic diversity within and between groups of host and geographical origin were calculated with Kimura 2-parameter (K2+G) which was chosen as the best-fitting model of nt substitution. To assess the statistical significance of the tree, bootstrap values were calculated on 1000 replicates. All branches with bootstrap value support <50% were collapsed. The isolate of *Pepper mild mottle virus* (PMMoV; Acc.No. M96425) was used as the outgroup sequence.

Isolate name*	Country	Host plant	GenBank			
Isolate name*	Country	Host plant	accession number			
D35-06, D7-06	Serbia	Nicotiana tabacum	GQ290475-76			
Slovenia1	Slovenia	/	AJ315739			
SI 50, SI 64	Slovenia	Solanum tuberosum	AF401603-04			
PVYN-N242	French	Solanum tuberosum	AF248499			
605	Switzerland	/	X97895			
Tu 648	Canada	Solanum tuberosum	AF401610			
v942490	United Kingdom	/	EF016294			
Linda, Satina	Germany	Nicotiana tabacum	AJ890345, 47			
423–3	USA	Solanum tuberosum	AY884982			
N 5Yt	Canada	Nicotiana tabacum	AF401605			
Rusia	United Kingdom	/	AJ315746			
California	California	/	AJ315744			
English	United Kingdom	/	AJ315747			
Tu 619, Tu 660	USA	Solanum tuberosum	AF401608-09			
N266, N394, N	Canada	Solanum tuberosum	AF401600, 01, 06,			
27, N Jg			07			
Canada	Canada	/	AJ315745			
USA	USA	/	AJ315742			
Ukraine	Ukraine	/	AJ315740			
803, Viikki	Finland	Solanum tuberosum	AJ245555-56			
Hungarian	Hungary	/	M95491			
Ditta	Poland	Nicotiana tabacum	AJ890344			
Scotland	United Kingdom	/	AJ315743			
Adgen	French	Solanum tuberosum	AJ890348			
LYE84.2	Spain	Solanum lycopersicum	AJ439545			
SON41	French	Solanum nigrum	AJ439544			

Table 1. PVY isolates with coat protein sequences from GenBank used in the phylogenetic analysis

* - All data are from GenBank

Results and Discussion

Disease incidence and symptomatology in the field

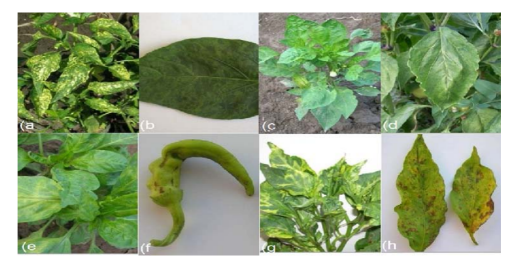
Disease incidence, as estimated by the number of symptomatic plants in the field, highly varied depending on year and sampling locality, ranging from 20% to 60%. Pepper plants exhibited a wide range of symptoms, including stunting, shrubby appearance of plants, yellowish and chlorotic patterns (Figure 1a, 1g), leaf deformation (Figure 1c, 1e), mottling (Figure 1d), mild (Figure 1b) or severe mosaic, veinal necrosis, necrotic spots on the leaves (Figure 1h) or necrotic line patterns, as well as stem and branch necrosis. Fruits of infected plants were

deformed, necrotic (Figure 1f), mottled, or without any symptoms. The observed symptoms were typical of virus infection, as described by many authors (Krstić *et al.* 1996; Echer and Costa 2002; Ozaslan *et al.* 2006; Mijatović *et al.* 2007; Sevik 2011).

Plants can display virus-like symptoms when plants respond to unfavorable weather, nutritional imbalances, infection by other types of pathogens, damage caused by pests or abiotic agents and other factors (van der Want and Dijkstra

2006); therefore the symptoms caused by the viruses have no diagnostic significance.

Figure 1. Symptoms of virus infection on pepper: a) Yellowing (AMV), b) Mosaic (PMMoV), c) Mild mosaic and leaf distortion (CMV), d) Mottling (PVY), e) Mosaic and leaf distortion (CMV+AMV), f) Deformations and necrosis of fruit (PVY+CMV), g) Severe yellowing and leaf distortion (PVY+AMV), h) Necrotic spots (PVY+CMV)



Virus detection and incidence in pepper samples

Serological analyses of collected pepper samples in 2009 and 2010 detected the presence of four viruses: CMV, PVY, AMV and PMMoV. The relative incidence of detected viruses in collected pepper samples is presented in Table 2.

During 2009, PVY was detected in 14 out of 16 inspected localities and was by far the most prevalent virus both in single (25.6%) and mixed infections (25.6%). The second widespread virus was CMV, detected in 13 localities in 37.2% of analyzed samples. AMV and PMMoV were also detected but at lower percentages. The AMV was detected in 21.7% of analyzed samples, while PMMoV was detected in only one locality in 3 analyzed samples. The number of

single infections (56.5%) was considerably higher than the number of mixed infections (26.6%). The most frequent type of mixed infections (18.8%) was infection with two viruses, PVY+CMV (Table 2). Triple infections were found in 1.9% of the tested samples. In 17% of the tested samples viruses were not detected.

In 2010, CMV was detected in 12 out of 15 inspected localities and was the most frequent virus both in single (35.4%) and mixed infections (17.7%). The second most frequent virus was PVY, detected in 15 localities in 43.0% of analyzed samples while AMV was detected in 17.7% of analyzed samples. PMMoV was detected in only one locality in 2 analyzed samples (1.3%). Single infections were also the most frequent infection type (74.7%), while mixed infections was PVY+CMV (14.6%), while triple infections were found in 1.3 % (Table 2). None of the tested viruses was observed in 12 % of the samples.

TMV, TSWV, PVX, PepMoV and PVMV were not detected by DAS-ELISA test in any of the tested samples during the two-year period.

In general, PVY was the most frequent virus, identified in 47.7% of 365 samples collected during the two years of investigation. The second most frequent virus was CMV, which infected 44.1% of tested samples. AMV was detected in 20% of tested samples, while the least frequent was PMMoV (1.4%). Single infections were the most frequent infection type in 2009 and 2010. Most common single infections in total collected samples were those of PVY (25.5%) and they were prevalent only in 2009, with the lower rate in 2010. A combination of two viruses was the most frequent mixed infection type.

The dominant presence of PVY was confirmed in other areas of pepper production in the world (Fanigliulo et al., 2005; Choi et al., 2005b). Also, PVY was the most frequently found virus in 1998 and the second most frequent in 1999 in Turkey (Arli-Sokmen et al., 2005). As opposed to PVY there are several reports displaying the evidence of the incidence of CMV which infects not only common vegetable crops but also some rare ornamental plants. A significant presence of CMV was demonstrated in other areas of pepper production in the world (Sepulveda et al., 2005; Choi et al., 2005a). According to many authors, CMV and PVY seem to be the most damaging viruses (Fakhfakh et al., 1999; Ben Khalifa et al., 2009). These two viruses often occur in mixed infection, which results in increased severity of the disease through synergy (Syller, 2012), which is in agreement with our studies. This study revealed that PVY and CMV were prevalent viruses in pepper crops in Serbia, but their occurrence, distribution, and relative incidence varied during the investigation. However, the dominant presence of PVY has been observed also in tobacco (Stanković et al., 2011), and it was one of the most frequently found virus in individual and mixed infections in tomato production in Serbia (Nikolić et al., 2018). Furthermore, CMV is well established in the tomato (Nikolić et al., 2018), different pumpkin species (Vučurović et al. 2012), tobacco (Stanković et al., 2011) and other hosts which serve as CMV reservoirs.

	Single infection (%)								Mixed in	fection (%)	Total infections (%)				
Year	Locality	No. of tested samples	ΡVΥ	CMV	AMV	PMMoV	PVY +CMV	PVY +AMV	AMV +CMV	PVY +PMMoV	AMV +CMV +PVY	PVY +PMMoV +CMV	ΡVΥ	CMV	AMV	PMMoV
2009	Selenča	10	0	0	50	0	0	0	0	0	0	0	0	0	50	0
	Ravno Selo	15	20	0	33.3	0	13.3	13.3	0	0	6.7	0	53.3	20	53.3	0
	Đurđevo	15	26.7	26.7	0	0	26.7	0	0	0	0	0	53.3	53.3	0	0
	B. Palanka	15	53.3	0	20	0	0	13.3	0	0	0	0	66.7	0	33.3	0
	Veternik	9	22.2	0	0	0	66.7	0	0	0	11.1	0	100	77.8	11.1	0
	Horgoš I	10	20	0	60	0	0	0	0	0	0	0	20	0	60	0
	Horgoš II	5	0	20	0	0	60	0	0	0	20	0	80	100	20	0
	Senta	20	35	15	25	0	0	0	0	0	0	0	35	15	25	0
	Čonoplja	9	66.7	0	0	0	0	0	0	22.2	0	1	100	11.1	0	33.3
	Velika Plana	15	0	46.7	20	0	0	0	13.3	0	0	0	0	60	33.3	0
	Smederevo	15	13.3	46.7	0	0	40	0	0	0	0	0	53.3	86.7	0	0
	Kraljevo	15	53.3	0	0	0/0	26.7	0	0	0	0	0	80	26.7	0	0
	Trstenik	10	40	40	0	0/0	0	0	0	0	0	0	40	40	0	0
	Čačak	15	0	26.7	0	0/0	40	0	0	0	0	0	40	66.7	0	0
	Inđija	15	33.3	0	0	0/0	26.7	26.7	0	0	0	0	86.7	26.7	26.7	0
	Aleksinac	14	14.3	14.3	35.7	0/0	28.6	0	0	0	0	0	42.9	42.9	35.7	0
Subtotal		207	25.6	15.5	15.5	0	18.8	3.85	0.97	0.97	1.45	0.5	51.2	37.2	21.7	1.5

Table 2. Incidence of AMV, CMV, PVY, and PMMoV in single, mixed and total infections in pepper crops in Serbia

	Single infection (%)						Mixed infection (%)							Total infections (%)			
Year	Locality	No. of tested samples	PVY	CMV	AMV	PMMoV	PVY +CMV	PVY +AMV	AMV +CMV	PVY +PMMoV	AMV +CMV +PVY	PVY +PMMoV +CMV	РVҮ	CMV	AMV	PMMoV	
2010	Đurđevo	7	57.1	0	14.3	14.3	0	0	0	0	0	0	57.1	0	14.3	14.3	
	Bački Jarak	10	20	40	0	0	20	0	10	0	0	0	40	70	10	0	
	Bukovac	10	50	10	0	0	30	0	0	0	0	0	80	40	0	0	
	Rimski Šančevi	10	20	30	20	0	20	0	10	0	0	0	40	60	30	0	
	Bačka Topola	15	0	40	26.7	0	20	0	0	0	0	0	20	60	26.7	0	
	Ruma	15	13.3	60	0	0	0	0	0	0	0	0	13.3	60	0	0	
	Vašica	7	14.3	14.3	14.3	0	51.1	0	0	0	0	0	71.4	71.4	14.3	0	
	Šabac	10	40	0	20	0	10	10	0	0	0	0	60	10	30	0	
	Horgoš	15	13.3	53.3	6.7	0	20	0	0	0	6.7	0	40	80	13.3	0	
	Kikinda	7	14.3	71.4	0	0	0	0	0	0	0	0	14,3	71.4	0	0	
	Čonopla	15	46.7	71.4	0	0	13.3	0	0	0	0	6.7	66.7	53.3	0	6.7	
	Smederevo	5	100	0	0	0	0	0	0	0	0	0	100	0	0	0	
	Užice	10	0	0	30	0	0	20	0	0	0	0	20	0	50	0	
	Aleksinac	12	16.7	58.3	8.3	0	0	0	8.3	0	0	0	16.7	66.7	16.7	0	
	Valjevo	10	20	30	20	0	10	0	0	0	0	0	40	40	20	0	
Subtotal		158	25.3	35.4	13.3	0.6	14.6	1.9	1.9	0	0.6	0.6	43.0	53.2	17.7	1.3	
Total		365	25.5	24.1	14.5	0.3	17.0	3.0	1.4	0.6	1.1	2.0	47.7	44.1	20	1.4	

Difficulties in controlling PVY and CMV infections are caused by the great variability of the virus, their polyphagous nature and numerous aphid vectors which transmit the virus with different efficiencies.

Molecular detection and sequence identity analyses of PVY isolates

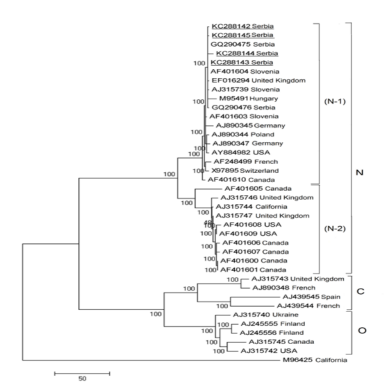
Specific primers (PVYc/PVYd) for the detection of PVY were able to amplify target cDNA fragments of predicted size and successfully detect the presence of the virus in all selected ELISA-positive samples. No amplification product was observed in the healthy pepper control. After purification, the identities of obtained amplicons were confirmed by sequencing.

The nucleotide sequences of the fragment of the P1 gene derived from the isolates PL-3-10 (KC288144), PL-28-09 (KC288142), PL-15-09 (KC288143) and PL-108-10 (KC288144) shared nucleotide identity from 99.1 to 99.9% (100 % aa identity). Isolates PL-3-10 exhibited the highest nucleotide identity (99.2%) with Slovenian isolate (AF401604), Croatian isolate (JF927749), German isolate (JF927756) and tobacco isolate (GQ290476) from Serbia. PL-15-09, PL-28-09 and PL-108-10 also exhibited the highest nucleotide identity (100%, 100% and 99.8%, respectively) with the above isolates.

The phylogenetic tree constructed using the maximum parsimony method based on sequences of the P1 gene of PVY showed the separation of selected isolates into three consistent lineages as reported earlier: N, C, and O (Moury et al., 2002; Fanigliulo et al., 2005; Glais et al., 2002, 2005; Lorenzen et al., 2006; Ogawa et al., 2008, 2012). N lineage was split into two sublineages (N-1 and N-2, previously described as N-Europe and N-North America sublineages, respectively), as reported in an earlier study by Ogawa et al. (2008). Genetic diversity among three molecular groups of isolates ranged from 0.087±0.010 to 0.374 ± 0.026 , whereas within each group the range was: 0.010 ± 0.002 (N-1), 0.019±0.003 (N-2), 0.141±0.011 (C) and 0.046±0.005 (O). All of the four Serbian isolates were clustered in sublineage N-1 with other European isolates of necrotic strains. The lowest genetic diversity was between sublineages (0.087 ± 0.010) , while the highest genetic diversity (0.374 ± 0.026) was between sublineages N-1 and lineages C, and sublineages N-2 and lineages C. Some exceptions were observed, such as placing of some isolates from Canada and the USA in the group of European necrotic isolates, thus confirming their European origin and probable introduction through potato import. The opposite situation was also observed, such as placing of isolates from Great Britain in the group with North American necrotic isolates, which confirms the "gene flow" between Europe and North America, enabled by the intensive exchange of plant material (Đekić, 2010).

Analysis of P1 gene sequences of four selected PVY isolates did not determine the heterogencity of this virus population which originated from pepper in Serbia. All tested PVY isolates originating from pepper were grouped into subcluster of necrotic isolates from Europe, as were the previously characterized PVY isolates from Serbia originating from tobacco (Đekić *et al.*, 2007). However, recent research has indicated that recombinations occur not only between strains, but also between different phylogenetic groups of the same strain, as well as between recombinants (Schubert *et al.*, 2007; Chikh Ali *et al.*, 2010; Kerlan *et al.*, 2011). Therefore, further examination of pepper PVY isolates from Serbia should be focused on sequencing of the whole genome to obtain a clear overview of variability and composition of the virus population in Serbia, as well as an insight into the main sources which spread PVY infection to susceptible crops.

Figure 2. Maximum parsimony tree based on nucleotide sequences of partial gene of P1 protein of KC288144, KC288142, KC288145 and KC288143 isolates of *Potato virus Y* (PVY) using M96425 isolate as out-group sequence. The four Serbian isolates are underlined



Conclusions

There is a need to develop resistant pepper varieties to overcome this disease threat. Knowledge of the pathogen and its variants or strains is a vital prerequisite for reliable breeding for resistance. This investigation provides essential basic information useful for pepper virus control in Serbia. An important option for virus disease control would be to enhance growers' acquaintance with control measures against pepper viruses and their vectors, especially those directed to limit early virus infections. The information about phylogenetic and genetic diversity obtained in this study will provide the first insight into the genetic structure of PVY isolates from pepper in Serbia.

Acknowledgements

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PRISUSTVO I RASPROSTRANJENOST VIRUSA PAPRIKE U SRBIJI

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Rezime

Dvogodišnjim poučavanjima (2009–2010) prisustva i rasprostranjenosti virusa u usevu paprike u Srbiji utvrđeno je da se virusi javljaju svake godine u proizvodnji paprike na otvorenom polju. Intenzitet zaraze bio je različit i kretao se od 20% do 60%, u zavisnosti od ispitivane godine i lokaliteta gajenja paprike. Serološkim testiranjem uzoraka paprike prikupljenih u više lokaliteta u Srbiji, tokom 2009 i 2010. godine detektovani su Cucumber mosaic virus (CMV), Potato virus Y (PVY), Alfalfa mosaic virus (AMV) and Pepper mild mottle virus (PMMoV), pri čemu je PVY bio dominantan. Specifičnim prajmerima PVYc/PVYd uz upotrebu RT-PCR metode umnožen je deo genoma od oko 975 bp koji kodira P1 protein. Amplifikovani fragmenti su sekvencirani i prijavljeni u GenBank bazu podataka, gde su im dodeljeni pristupni brojevi PL-28-09 (KC288142), PL-15-09 (KC288143), PL-3-10 (KC288144), PL-108-10 (KC288144). Proračunom genetičke sličnosti sekvenci izolata dobijenih u ovom radu utvrđen je visok stepen nukleotidne sličnosti, koji se kretao od 99,8-100%. Ispitivane sekvence PVY izolata iz Srbije dele najveću nukleotidnu i aminokiselinsku sličnost sa izolatima iz Slovenije, Hrvatske, Nemačke i izolatom duvana iz Srbije. Svi ispitivani izolati sa paprike poreklom iz Srbije grupisani su u subklaster nekrotičnih izolata poreklom iz Evrope.

Ključne reči: virusi paprike, serološka analiza, RT-PCR test, molekularna karakterizacija.