

Comparative genetic diversity of potato virus Y populations based on coat protein gene

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Summary. – Potato virus Y (PVY) is an important plant pathogen with a wide host range including economically important crops of potato, tobacco, tomato, and pepper. The coat protein gene has been commonly used in studying molecular biology of plant viruses including PVY. In this study, we used a large dataset of CP sequences from isolates collected across the world to assess the detailed molecular evolution of PVY populations with a focus on the Iranian PVY population. Phylogenetic analysis showed that the world PVY population had two major lineages (O:C and N:NTN); each comprising several divergent sublineages. Results showed that the Iranian PVY isolates were distributed across the tree suggesting polyphyletic origin of the Iranian PVY population. Statistical analysis revealed great genetic differences between pairs of the PVY phylogenetic populations. Host populations and also geographical populations of PVY were genetically differentiated. The extent of the genetic diversification among PVY host and geographical populations were mild or moderate. Purifying selection was detected on the CP gene sequences of the PVY populations, suggesting that most of the mutations in the gene were harmful, thereby were eliminated by natural selection. We also detected a variety of recombination patterns to occur along the CP gene of the PVY strains. A significant number of the Iranian PVY isolates were found to be recombinant. Different analyses suggest that Iranian PVY population is highly diverse. In conclusion, results of this study demonstrated that different factors including mutation, host adaptation, geographical distinction and selection pressure shaped the genetic structure of the PVY populations.

Keywords: potato virus Y; genetic variability; genetic differentiation; evolution.

Introduction

Potyviruses are the largest family of plant viruses. They have flexuous filamentous particles containing a single-strand, positive sense RNA molecule of about 9.7 kb (Shukla *et al.*, 1994; Fauquet *et al.*, 2012). Their genome has a viral-encoded VPg protein at its 5' end followed by a single open reading frame (ORF) and a poly-A tail at the 3' end. The viral genome is translated into a single large polyprotein, and then cleaved into at least ten multifunctional proteins by viral proteinases (Boonham *et al.*, 2002; Gibbs and Ohshima 2010; Martin *et al.*, 2010; Fauquet *et al.*, 2012). Moreover, during translation, there is a frame shift in the P3 gene resulting in production of a new short polypeptide (PIPO) (Chung *et al.*, 2008). Host adaptation determines the length of PIPO among potyviruses. Some members of potyviruses have another ORF, named PISPO, which is a suppressor of gene silencing (Clark *et al.*, 2012). However, existence of PISPO in PVY genome is still to be determined.

Potato virus Y (PVY) is the type species of the genus *Potyvirus*. It is responsible for serious diseases in a wide range of plant species, mostly from the family Solanaceae, with worldwide distribution. Like other potyviruses, PVY is transmitted by aphids in a non-persistent manner. Because of the wide host range and prevalence of the virus, PVY is probably the most destructive and widespread plant pathogen across the world (Shukla *et al.*, 1994).

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Abbreviations: PVY = potato virus Y

1 To date, several distinct strains of PVY have been identi-
 2 fied according to their biological, serological and genome
 3 sequence analyses. These strains are designated as PVY^O
 4 (the ordinary strain), PVY^N (the tobacco venial necrosis)
 5 and PVY^C (the stipple streak strain; Moury *et al.*, 2002).
 6 Furthermore, three other strains including of PVY^{NTN},
 7 PVY^{N-Wi} and PVY^{N-O} have been identified as being generated
 8 by recombination between PVY^O and PVY^N (Le Romancer
 9 *et al.*, 1994; Glais *et al.*, 2002). PVY^{NTN} induces potato tuber
 10 necrotic ringspot disease (PTNRD). Since the first report
 11 from Hungary in 1980s, the PVY^{NTN} recombinant has been
 12 frequently identified within the PVY populations worldwide
 13 (Le Romancer *et al.*, 1994; Karasev and Gray, 2013). The
 14 PVY^{NTN} has two main recombination patterns. The first one
 15 (PVY^{NTNa}) contains three recombination junctions (RJs),
 16 while the second structure (PVY^{NTNb}) has one RJ in P1 in
 17 addition to the three RJs in common with PVY^{NTNa} (Chikh
 18 Ali *et al.*, 2010).

19 Molecular evolutionary studies of viruses have shown
 20 the impacts of mutation, recombination, selection pressure
 21 and host adaptation in dynamics of viral populations. These
 22 studies have shed light on the important features of viral
 23 biology such as changes in virulence, geographical spread
 24 and adaptation to new hosts or emergence of a new virus
 25 epidemic. Comparisons of the genetic structure of different
 26 populations of a virus species can determine factors affect-
 27 ing ecology, phylogeny and phylogeographic structure of
 28 the populations across the world (Bermingham and Moritz,
 29 1998). Such studies provide useful information for designing
 30 better strategies aimed at controlling of the viruses (Jones,
 31 2009; Elena *et al.*, 2011). Although increasing number of
 32 studies on the population structure of animal and human
 33 RNA viruses is available, the population structure of plant
 34 viruses is poorly understood (Garcia-Arenal *et al.*, 2001).

35 PVY is reported to be one of the viruses responsible for the
 36 severe yield losses in Iran annually (Pourrahim *et al.*, 2007).
 37 Based on the host range, symptomatology and serological
 38 traits, PVY^O, PVY^{NW}, PVY^C, PVY^N and PVY^{NTN} strains have
 39 been recognized in potato fields of Iran. Furthermore, the
 40 sequence of P1, coat protein (CP) and 3' UTR of some of
 41 the Iranian PVY isolates has been reported (Hosseini *et al.*,
 42 2011).

43 To study the population structure of PVY in Iran, we ana-
 44 lyzed the biological properties and CP sequence variability of
 45 Iranian PVY isolates (including 11 new sequences provided
 46 in this study) in comparison with PVY populations from
 47 other countries. We focused on coat protein gene (CP) be-
 48 cause of its multifunctional properties, sequence variability
 49 and availability of large sequence data for most of the PVY
 50 isolates reported from around the world (Ogawa *et al.*, 2008;
 51 Visser and Bellstedt, 2009; Moury and Simon, 2011).

52 To evaluate, in more detail, the molecular evolution of
 53 PVY populations, we compared the CP gene sequence of

the Iranian PVY isolates with counterparts reported from 54
 different parts of Europe, North and South America, Africa 55
 and Japan (n = 542). 56

57 58 59 60 61 62 63 64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 100 101 102 103 104 105 106

Virus source and serological diagnosis of PVY. A total of 185
 potato leaf samples were collected during July 2013 to July 2014
 from the potato fields of Khorasan Razavi, Northern Khorasan and
 Fars provinces of Iran. The potato plants were sampled randomly
 regardless of their symptoms. Specific polyclonal antibody against
 PVY (DSMZ, Germany) was used to detect the virus in fresh leaf
 crude extracts by DAS-ELISA (Clark and Adams, 1977).

Samples, RT-PCR, cloning and sequencing. Total plant RNA
 was extracted from fresh leaves using Total RNA isolation kit
 (Denazist Asia-Iran) following the manufacturer's instructions.
 The First cDNA strand was synthesized using antisense primer
 (PVY-CPR) and moloney murine leukemia virus (MMuLV)
 reverse transcriptase (Fermentas, Lithuania). 5 µl of the purified
 RNA were mixed with reverse transcription mixture (50 mmol/l
 Tris-HCl pH 8.3, 50 mmol/l KCl, 4 mmol/l MgCl₂, 10 mmol/l
 dithiothritol, 1 mmol/l of each dNTP, 200 U of MMuLV reverse
 transcriptase) and incubated at 42°C for 1 h. The complete length
 of the coat protein (CP) gene was amplified using specific primers
 PVY-CPF (5'-GCTTTCACTGAAATGATGGT-3') and PVY-CPR
 (5'-GTTTTCCCAGTCACGACTTTTTTTTTTTT-3') by Taq DNA
 Polymerase Master Mix Red (Amplicon, Denmark). These primers
 amplify the complete CP gene of the PVY (~800 bp length; Nie and
 Singh, 2003). PCR products were analyzed on 1% agarose gel and
 purified from the gel using the Qiaquick gel extraction kit (Qiagen,
 Germany). The purified products were ligated into pTG19 vector
 (Vivantis, Malaysia) according to the manufacturer's protocol.
 Plasmids were transformed into *Escherichia coli* strain DH5a and
 the recombinant plasmids were purified from the bacterial cells
 using Plasmid DNA Isolation Kit (Denazist-Iran). Finally, the puri-
 fied recombinant plasmids were bi-directionally sequenced using
 pUC-M13 universal primers at MacroGen Inc. (Seoul, South Korea).
 Consensus sequences were verified using the BLAST program in
 NCBI database.

Phylogenetic analysis. Due to space limitation to show the tree
 in a single page, preliminary neighbor joining clustering of isolates
 was carried out using SDT v.1 software. Then 150 representatives
 out of 542 PVY-CP sequences were selected and phylogenetic tree
 was reconstructed. An alignment of the CP gene sequences of 150
 PVY isolates, including 26 isolates from Iran (11 new sequences
 obtained in this study) and 124 isolates from other countries (Table
 1), was generated using the Muscle module in MEGA v.5 (Tamura
et al., 2011). Details of the PVY isolates, their country of origin, and
 GenBank accession numbers are shown in Table 1. MEGA v.5. was
 used for generating phylogenetic tree by the Maximum likelihood
 method with HKY+G4 nucleotide substitution model as the best
 fitted model. Integrity of the evolutionary relationships was assessed

Table 1. Accession numbers and origins of potato virus Y isolates whose coat protein were used in this study

Populations	Origin	Number	Acc. Nos.
Middle East	Iran	26	I23cm,145,156,168,194,211,21,221,52cm,61,EU713856,HM243471,HM243472,HM243473,HM243474,HM243475,HM243476,HM243477,HM243478,HM243479,HM243480,HM243481,HM243482,HM243483,HM243484,IRAN
	Syria	14	AB185831,AB185832,AB185833,AB185833,AB256029,AB270705,AB270705,AB295475,AB295477,AB29547,AB461450,AB461451,AB461452,AB461452
	Iraq	13	JQ026006,JQ026007,JQ026009,JQ026010,JQ026011,JQ026012,JQ026013,JQ026014,JQ026015,JQ026016,JQ026017,JQ026018,JQ026019
	Jordan	6	EU073854,EU073855,EU073856,EU073857,EU073858,EU073859
	India	2	AF118153,JN034046
Far East	Pakistan	1	JQ518267
	Japan	45	AB331515,AB331516,AB331517,AB331518,AB331519,AB331538,AB331539,AB331540,AB331541,AB331542,AB331543,AB331544,AB331545,AB331546,AB331547,AB331548,AB331549,AB331550,AB702945,AB702950,AB702951,AB702952,AB702953,AB702954,AB702955,AB702956,AB711143,AB711144,AB711145,AB711146,AB711147,AB711148,AB711149,AB711150,AB711151,AB711152,AB711153,AB711154,AB711155,AB714134,AB714135,AB719459,D12539,D12570
	China	34	AJ488834,AJ488834,AM931254,EF592514,EF592514,EF592515,EF592515,EF592516,EF592516,EF592521,EF592521,EF592525,EF592525,EF592526,EF592526,EU182576,EU719648,EU719650,FJ423031,FJ423032,GQ200836,FJ766535,HM590405,HM590406,HM590407,HQ603083,HQ631374,JQ663997,JQ673517,JX872404,JX872405,KJ634024,KJ801915,PVU25672
Africa	Vietnam	8	AM411502,AM411503,AM411504,DQ925435,DQ925437,FM200035,FM201468,FM201468
	South Korea	1	EU885418
	Egypt	2	AF522296,AF52229
	South Africa	88	GQ853593,GQ853594,GQ853595,GQ853596,GQ853597,GQ853598,GQ853599,GQ853600,GQ853601,GQ853601,GQ853602,GQ853603,GQ853606,GQ853607,GQ853608,GQ853609,GQ853610,GQ853611,GQ853612,GQ853613,GQ853614,GQ853615,GQ853616,GQ853617,GQ853618,GQ853619,GQ853620,GQ853621,GQ853622,GQ853623,GQ853624,GQ853625,GQ853626,GQ853627,GQ853628,GQ853629,GQ853630,GQ853631,GQ853632,GQ853633,GQ853634,GQ853635,GQ853636,GQ853637,GQ853638,GQ853639,GQ853640,GQ853641,GQ853642,GQ853643,GQ853644,GQ853645,GQ853646,GQ853647,GQ853648,GQ853649,GQ853650,GQ853651,GQ853652,GQ853653,GQ853655,GQ853656,GQ853657,GQ853658,GQ853659,GQ853660,GQ853661,GQ853662,GQ853663,GQ853664,GQ853665,GQ853666,GQ853667,JN936418,JN936420,JN936422,JN936423,JN936425,JN936427,JN936429,JN936430,JN936432,JN936435,JN936437,JN936438,JN936439,JN936440,JN936441
	United Kingdom	41	AF325927,AF325928,AJ390285,AJ390285,AJ390286,AJ390286,AJ390287,AJ390287,AJ390288,AJ390288,AJ390289,AJ390290,AJ390291,.,AJ390292,AJ390294,AJ390295,AJ390296,AJ390297,AJ390301,AJ390302,.,AJ390303,AJ390304,AJ390305,AJ390306,AJ390308,AJ390309,AJ585195,AJ585196,AJ585197,AJ585198,EF016294,JQ954337,JQ954338,JQ954339,KC614702,KC634004,KC634005,KC634006,KC634007,KC634008,KC634009
Europe	Germany	29	AJ889867,AJ889868,AJ890347,AJ890350,AM113988,HE608963,HE608964,JQ954295,JQ954296,JQ954297,JQ954298,JQ954299,JQ954300,JQ954301,JQ954302,JQ954303,JQ954304,JQ954305,JQ954307,JQ954308,JQ954309,JQ954310,JQ954311,JQ954312,JQ954313,JQ954314,JQ954315,JQ954316,JQ954317,JQ954318,JQ954319,JQ954320,JQ954321,JQ954322,JQ954323,JQ954324,JQ954325,JQ954326
	Poland	11	AJ889866,AJ890342,EF558545,EF558545,FJ666337,JF795485,JF804780,JF804781,JF804785,JF804786,JF804787
	Switzerland	9	JQ954331,JQ954332,JQ954333,JQ954334,JQ954335,JQ954336,JQ954339,JQ954394,X97895
	Belgium	8	JQ969033,JQ969034,JQ969035,JQ969036,JQ969037,JQ969039,JQ969040,JQ969041
	Czech Republic	8	JQ954343,JQ954344,JQ954345,JQ954346,JQ954347,JQ954348,JQ954340,JQ954341
	Italy	5	JQ954322,JQ954323,JQ954324,JQ954325,JQ954326
	Slovenia	4	AJ390293,AJ585342,JQ954376,JQ954377
	Netherlands	4	JQ954327,JQ954328,JQ954329,JQ954330
	Hungary	3	AJ390300,M95491,M95491
	France	2	AJ890348,HM991454
Other	Portugal	1	AJ390307
	Denmark	1	AJ390298,AJ390299
	Netherlands	1	EU563512
	Latvia	1	GQ496607
	Greece	1	JQ954321
	Finland	1	JX424837

Table 1. Continue

Populations	Origin	Number	Acc. Nos.			
N. America	USA	108	AJ390309,AY884982,AY884983,AY884984,FJ666337,AY884985,DQ008213,DQ157178,DQ157179,DQ157180,EF026074,EF026075,EF026076,FJ204164,FJ204165,FJ204166,FJ643477,FJ643478,FJ643479,HQ912862,HQ912863,HQ912864,HQ912865,HQ912866,HQ912867,HQ912868,HQ912869,HQ912870,HQ912871,HQ912872,HQ912873,HQ912874,HQ912875,HQ912876,HQ912877,HQ912878,HQ912879,HQ912880,HQ912881,HQ912882,HQ912883,HQ912884,HQ912885,HQ912886,HQ912887,HQ912888,HQ912889,HQ912890,HQ912891,HQ912892,HQ912893,HQ912894,HQ912895,HQ912896,HQ912897,HQ912898,HQ912899,HQ912900,HQ912901,HQ912902,HQ912903,HQ912904,HQ912905,HQ912906,HQ912907,HQ912908,HQ912909,HQ912910,HQ912911,HQ912912,HQ912913,HQ912914,HQ912915,JQ954349,JQ954350,JQ954351,JQ954352,JQ954353,JQ954354,JQ954355,JQ954356,JQ954357,JQ954358,JQ954359,JQ954360,JQ954361,JQ954362,JQ954363,JQ954364,JQ954365,JQ954366,JQ954367,JQ954368,JQ954369,JQ954370,JQ954371,JQ954372,JQ954373,JQ954374,JQ954375,JQ954385,JQ954386,JQ954388,JQ954389,JQ954390,JQ954391,JQ954392,U91747			
			Canada	11	AF126258,AY166866,,AY166867,,AY512655,AY745491,AY745492,HM367075,HM367076,PVU09509,U09508,U09509	
					Mexico	1
			S. America	Brazil	11	AF255659,AF255660,AF525081,AY840082,JF928458,JF928459,JF928460,JQ924285,JQ924286,JQ924287,JQ924288
				New Zealand	2	AM268435,DQ217931

by 1000 bootstrap replicates. Pairwise distance comparisons of the CP gene sequences were computed using the Tamura-Nei model in MEGA v. 5.0.

Population genetics analysis. DnaSP version 5.10.01 (Libardo and Rozas, 2009) was used to estimate the average pairwise nucleotide diversity (π , average distances between pairs of sequences; Tajima 1983), number of polymorphic site (S), total number of mutations (η), average number of nucleotide differences among sequences from the same population (K), haplotype diversity (H_d , number of haplotypes within a sample), and the ratio of non-synonymous to synonymous nucleotide diversity (dN/dS), also known as ω ratio. The nucleotide diversity (π) may range between 0.0 (no variation) and 0.100 (highest variation between sequences). The haplotype diversity may range from 0.0 and 1.000. This program was also used for Tajima's D (Tajima 1989), F_u and Li's D^* and F^* (Fu and Li, 1993) tests of neutrality. Tajima's D test is based on the differences between the numbers of segregating sites and the average number of nucleotide differences. F_u and Li's D^* test is based on the differences between the numbers of mutations that appear in only one sequence and the total numbers of mutations. F_u and Li's F^* test is based on the differences between the numbers of singletons and the average number of nucleotide differences between every pair of sequences.

Tests of population differentiation. Statistical tests of population differentiation including K_{st}^* , Z^* , S_{nn} and F_{ST} , were calculated using DnaSP version 5.10.01. The K_{st}^* test statistic of genetic differentiation (Hudson, 2000) is expected to be near zero if there is no genetic differentiation (null hypothesis). But the null hypothesis is rejected, when K_{st}^* is supported by a small P-value (0.05) (Tsompana *et al.*, 2005). The Z statistic is calculated from ranking distances between all pairs of sequences. The average ranks for those from within two populations are summed, and the sum is weighted. The Z^*

statistic is a logarithmic variant of the Z statistic (Hudson, 2000). Also, small values of Z lead to rejection of the null hypothesis (no genetic differentiation). The nearest neighbor statistic (S_{nn}) was used to evaluate the frequency of the nearest neighbors of sequences found in the same group (Hudson, 2000). S_{nn} statistic may range between 1.0, when populations from distinct groups were genetically different, to 0.5 when a population is panamictic (Tsompana *et al.*, 2005). F_{ST} is the coefficient of gene differentiation or fixation index, which measures the extent of inter-population diversity with values ranging between 0 (indicating no differentiation between the populations) and theoretical maximum of 1 (when the populations are clearly differentiated; Hudson 2000). However, the observed F_{ST} is much less than 1, even in highly differentiated populations. Statistical significance for all three tests (Z^* , S_{nn} and F_{ST}) was established using 1000 permutations test.

Tests of recombination. Alignments of 542 PVY-CP sequences were analyzed for intraspecies recombination events using Recombination Detection Program (RDP v.4.10 beta) with default parameters (highest acceptable probability value = 0.05; Martin *et al.*, 2010). The RDP4 software detects the occurrence of robust recombination events using a suite of methods including Rdp, Geneconv, Bootscan, Maxchi, Chimaera, 3Seq, Siscan, Lard and Phylpro. Bootscan, Rdp and Siscan are phylogenetic methods, Geneconv, Maxchi, Chimaera and Lard are substitution-based methods, and Phylpro is a distance comparison method. Recombination sites detected at least by four methods were considered as "significant recombination events" and those detected by fewer methods were considered as "tentative recombination events" (Heath *et al.*, 2006).

Biological characterization of the PVY isolates. The partial host range of four Iranian PVY isolates were compared by their inoculation to potato (*Solanum tuberosum*), tomato (*Lycopersicon*

1 *esculentum*), tobacco (*Nicotiana tabacum* var Turkish, *Nicotiana*
2 *tabacum* var Xanthi), pepper (*Capsicum annuum*) and *Chenopo-*
3 *dium quinoa* seedlings. To this purpose, potato leaf tissues infected
4 by these PVY isolates were separately ground in 5 volumes of 50
5 mmol/l phosphate buffer at pH 7.5 containing 2% polyvinylpyr-
6 rolidone (PVP) and 0.05% 2-mercaptoethanol. The extracts were
7 mechanically inoculated on carborandum-dusted leaves. Two weeks
8 post inoculation plants were examined for symptoms and tested
9 for PVY infection using RT-PCR as described above.

11 Results

14 Identification and sequencing of CP genes of PVY isolates

16 From a total of 185 plants sampled, PVY was detected by
17 DAS-ELISA in 46 samples collected from Khorasan-Razavi,
18 Northern-Khorasan (North-East), and Fars (South-East)
19 provinces of Iran. PVY was detected in potato plants show-
20 ing symptoms of yellowing, mosaic, leaf crinkling as well as
21 in the asymptomatic plants.

22 The complete nucleotide sequence of the CP gene from
23 11 Iranian PVY isolates were determined and deposited in
24 GenBank (Table 2).

26 Phylogenetic positions of the PVY populations

28 The maximum likelihood phylogenetic tree of the CP gene
29 from 150 PVY isolates, including 26 isolates from Iran (11
30 sequences provided in this study and 15 sequences obtained
31 from GenBank) and 124 isolates from other countries (ob-
32 tained from GenBank), revealed the segregation of PVY iso-
33 lates into two main evolutionary divergent clades (Fig. 1).

34 These two major clades, designated as PVY^{O:C} and
35 PVY^{N:NTN} included 84 and 66 isolates, respectively. The O:C
36 clade was partitioned into two sister clades designated as
37 O (PVY^O) and C (PVY^C).

The PVY^O clade (Ordinary clade) was divided into five dis- 54
tinct subclades designated as PVY^O-O5, PVY^O-OJ1, PVY^O- 55
O1, PVY^O-O2 and PVY^O-O3. The PVY^O-O2 and PVY^O-O3 56
were the predominant genotypes within the PVY^O clade. 57
Furthermore, the PVY^{N:NTN} clade included two subclades 58
termed as N-Europe and N-North America (Fig. 1). The 59
N-Europe subclade could be further segregated into two 60
groups of PVY^{NTNa} and PVY^{NTNb} 61

62 As shown by the phylogenetic tree, PVY isolates were ar-
63 ranged between the two main clades (PVY^{O:C} and PVY^{N:NTN}
64) irrespective of their country of origin. This suggests that
65 geographical origin has no significant effect on the phyloge-
66 netic divergence of PVY isolates into two main clades. The
67 Iranian isolates were also found to be distributed between
68 these two clades. Of 26 isolates from Iran, 15 isolates grouped
69 within the PVY^{O:C} clade and 11 isolates grouped within the
70 PVY^{N:NTN} clade (Fig. 1).

71 The PVY^O-O2 genotype (n = 10) was the most common
72 genotype among Iranian isolates of PVY^O. None of the
73 Iranian PVY^O isolates placed in PVY^O-O5 and PVY^O-OJ1
74 clades. The majority of the Iranian PVY^{N:NTN} genotypes were
75 clustered with PVY^{NTNa} group in N-Europe subclade.

76 The mean sequence distance among isolates of PVY^{O:C} and
77 PVY^{N:NTN} clades was 3.6% and 3.4%, respectively. The mean
78 sequence distance between these two clades was 9.8%.

79 The Iranian PVY isolates in the PVY^{O:C} clade shared
80 96.46±0.52% CP nucleotide sequence identity with each
81 other and 96.01±1.63% with other isolates of this clade. On
82 the other hand, the Iranian PVY isolates in the PVY^{N:NTN}
83 had 93.76±3.39% CP nucleotide sequence identity with each
84 other and 96.13±2.4% with other isolates of the clade.

86 Population characteristics and polymorphism

88 To study the diversity of the PVY population, all PVY CP
89 sequences (determined in this study or obtained from Gen-
90 Bank; n = 542), were categorized into 3 population groups
91

93 Table 2. Characterization of Iranian potato virus Y isolates reported in this study

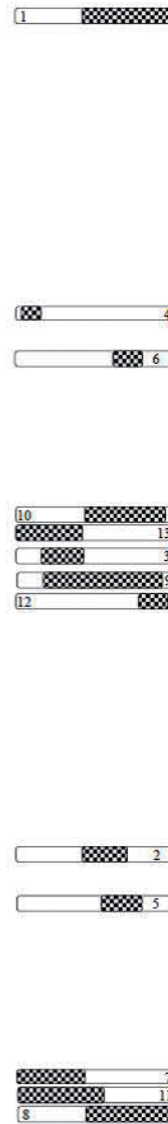
95 Samples	95 Acc. No.	95 Host	95 Strain	95 DAS-ELISA	95 RT-PCR
96 IRAN	96 LN908250	96 potato	96 NTN	96 +	96 +
97 52cm	97 LN907865	97 potato	97 NTN	97 +	97 +
98 123cm	98 LN907866	98 potato	98 NTN	98 +	98 +
99 168	99 LN907864	99 potato	99 NTN	99 +	99 +
100 156	100 LN907860	100 potato	100 N:O	100 +	100 +
101 211	101 LN907861	101 potato	101 SYR-NB/O	101 +	101 +
102 21	102 LN880858	102 potato	102 O	102 +	102 +
103 194	103 LN907859	103 potato	103 O	103 +	103 +
104 221	104 LN907862	104 potato	104 O	104 +	104 +
105 145	105 LN908252	105 potato	105 O	105 +	105 +
106 61	106 LN908251	106 potato	106 O:C	106 +	106 +



Fig. 1

Maximum likelihood phylogenetic analysis of 150 PVY isolates based on the CP gene nucleotides sequences using HKY+G4 nucleotide substitution model, coupled with schematic diagram of PVY recombinants

Recombinant junctions are shown as checker-board box. Iranian isolates are highlighted with green color. Numbers in rectangular represent number of the recombination pattern listed in Table 7.



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1 based on their phylogenetic relationship (4 phylogenetic
2 populations), host adaptation (3 host populations) and geo-
3 graphical origin (6 geographical populations; Table 3).

4 Among 4 PVY phylogenetic populations, strains of PVY^C
5 had the most average number of CP nucleotide differences
6 ($k = 55$ nucleotides) and the highest overall nucleotide di-
7 versity ($\pi = 0.0691$). The largest number of segregating sites
8 ($S = 350$) and mutations within the segregating sites ($\eta = 442$)
9 were observed within the CP nucleotide sequences of the
10 PVY^O lineage (Table 3). The PVY^{NTN} population had the
11 lowest values of k (10.50) and π (0.0131; Table 3). On the
12 other hand, the ratio of nonsynonymous nucleotide diversity
13 to synonymous nucleotide diversity (ω ratio) was <1 for all
14 phylogenetic populations. The highest and the lowest ratios
15 were obtained for the PVY^{NTN} and PVY^N populations with
16 the values of 0.351 and 0.046, respectively. The dN/dS value
17 of the CP gene from different PVY phylogenetic populations
18 ranged between 0.046 and 0.35 (Table 3).

19 PVY host populations included those isolated from po-
20 tato, tobacco and other hosts (e.g. tomato; Table 3). Among
21 3 PVY host populations, the greatest average number of
22 CP nucleotide differences ($k = 61$) and the greatest overall
23 nucleotide diversity ($\pi = 0.0771$) was observed in the PVY
24 population isolated from other hosts (other than potato and
25 tobacco; Table 3). The CP sequence analysis showed that PVY
26 population of potato had the highest number of segregating
27 sites ($S = 442$) and mutations ($\eta = 611$; Table 3). The k and π
28 statistics were the lowest for the PVY population of tobacco
29 with values of 45 and 0.0572, respectively (Table 3). The dN/
30

dS value of the CP gene sequence from the three PVY host 54
populations ranged from 0.08 to 0.1 (Table 3). 55

Whereas the PVY population of tobacco had the lowest ω 56
ratio (0.0836) in the CP gene, the value was highest for PVY 57
strains isolated from hosts other than potato and tomato 58
(other hosts, $\omega = 0.1031$). 59

We also defined 7 geographical populations based on 60
the geographical origin of the PVY isolates (Table 3). The 61
CP Molecular variability revealed that the South American 62
PVY population had the highest number of nucleotide 63
differences ($k = 65$) and the highest overall nucleotide 64
diversity ($\pi = 0.0814$). The largest number of segregating 65
sites ($S = 250$) and mutations within the segregating sites 66
($\eta = 297$) was observed within the Iranian PVY population 67
(Table 3). Moreover, the dN/dS ratio (ω) was <1 for all of 68
the geographical populations except for African population 69
($\omega = 1.03$). In this regard, the largest (1.031) and lowest 70
(0.095) ω were observed in PVY populations of Africa and 71
South America, respectively (Table 3). 72

Genetic differentiation of populations 74

We evaluated the genetic distinction of PVY popula- 76
tions defined in 3 categories: phylogenetic populations, 77
host populations, and geographical populations. In pursue 78
of this goal the PVY-CP gene sequences were subjected to 79
four independent statistical tests of population differentia- 80
tion. The null hypothesis (no genetic differentiation) can be 81
rejected if the statistics Kst*, K, Z, and Snn for the majority 82

Table 3. Genetic characteristics of potato virus Y coat protein from different populations

	Popula- tions	No. seq.	S	η	k	π	SS	NS	dS	dN	dN/dS
Phylogenetic population	All	542	496	722	49.5229	0.0621	179.76	609.24	0.2611	0.0207	0.0794
	O	311	350	442	24.0685	0.0300	182.09	618.91	0.0778	0.0180	0.2323
	C	21	190	211	55.3455	0.0691	182.24	618.76	0.2507	0.0284	0.1133
	N	33	97	103	27.1014	0.0339	180.95	614.05	0.1476	0.0069	0.0468
	NTN	177	190	205	10.5051	0.0131	181.64	619.36	0.0272	0.0095	0.3511
Host popu- lation	Potato	425	442	611	49.039	0.0613	181.04	616.95	0.2314	0.0210	0.0922
	Tobacco	100	255	302	45.678	0.0572	180.82	611.17	0.2176	0.0182	0.0836
	Other host	17	188	208	61.765	0.0771	182.08	618.91	0.2889	0.0298	0.1031
Geographical popu- lation	Iran	26	250	297	58.370	0.0728	182.32	618.67	0.2193	0.0393	0.1792
	Middle East	37	183	203	43.540	0.0551	164.95	588.05	0.0726	0.0549	0.756
	Far East	97	201	226	46.479	0.0584	168.23	608.77	0.0626	0.0620	0.990
	Africa	90	200	217	44.387	0.0556	168.73	608.27	0.0584	0.0602	1.031
	Europe	147	213	251	46.727	0.0584	169.49	610.51	0.0630	0.0620	0.984
	N. America	129	199	214	39.581	0.0494	169.31	613.69	0.0612	0.0498	0.813
S. America	16	177	201	65.275	0.0814	182.11	618.88	0.3162	0.0301	0.095	

51 S: number of segregation sites; η (Eta): total number of mutations; k: average number of nucleotide differences between sequences; π (pi): nucleotide
52 diversity; SS: total number of synonymous sites analyzed; NS: total number of non-synonymous sites analyzed; π (s): synonymous nucleotide diversity;
53 π (a): non-synonymous nucleotide diversity. Bold: indicate the highest value of each test. 104
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106

1 of comparisons were supported by *P*-values less than 0.05
2 (Hudson 2000). (Tables 4 and 5).

3 First, our statistical analysis revealed significant genetic dif-
4 ferences between all pairs of the PVY phylogenetic populations
5 defined in this study (6 pairs; Table 4). The highest (0.722) and
6 the lowest (0.403) F_{ST} value was found when comparing PVY^O
7 population with PVY^{NTN} and PVY^C populations, respectively.
8 This suggested the existence of a great genetic differentiation
9 between all PVY phylogenetic populations.

10 Considering host populations, the null hypothesis of no
11 genetic differentiation was rejected between all pairs (3 pairs)
12 of the PVY host populations (populations of potato, tobacco
13 and other hosts; Table 4). Non-potato PVY populations could
14 also be specified by phylogenetic analysis as they were mostly
15 placed in the PVY^C clade (Fig. 1). The F_{ST} values among PVY
16 host populations ranged between 0.051 and 0.100 (Table 4).
17 This finding concluded a moderate genetic differentiation
18 among PVY host populations.

19 **Table 4. Genetic differentiation estimates for populations of potato virus Y**

Population	Ka/Ks	Ks*	Kst*	Ks*,Kst* P-value	Z*	P-value	Snn	P-value	F_{ST}
O/C	0.097	3.18	0.056	0.000 ***	7.59	0.000***	1.000	0.000***	0.403
O/N	0.061	3.05	0.134	0.000 ***	7.47	0.000 ***	0.991	0.000 ***	0.722
O/NTN	0.098	2.72	0.224	0.000 ***	7.51	0.000 ***	0.993	0.000 ***	0.770
C/N	0.062	3.37	0.144	0.000 ***	5.10	0.000 ***	1.000	0.000 ***	0.581
C/NTN	0.088	2.48	0.198	0.000 ***	6.32	0.000 ***	1.000	0.000 ***	0.625
N/NTN	0.038	2.34	0.196	0.000 ***	6.45	0.000 ***	1.000	0.000 ***	0.592
Potato/tobacco	0.083	3.50	0.014	0.000 ***	10.20	0.000 ***	0.824	0.000 ***	0.051
Potato/other	0.096	3.58	0.005	0.002 **	9.77	0.002 **	0.935	0.001 **	0.075
Tobacco/other	0.088	3.42	0.021	0.000 ***	7.76	0.000 ***	0.899	0.000 ***	0.100

20 Ks, Kst*, Z* and Snn are test statistics of genetic differentiation [30]; F_{ST} examines the extent of genetic differentiation between geographical isolates; *:
21 0.01<P<0.05; **: 0.001<P<0.01; ***: P <0.001; ns: not significant.

22 **Table 5. Genetic differentiation estimates for geographical populations of potato virus Y**

Population	Ks*	Kst*	Ks*,Kst* P-value	Z*	P-value	Snn	P-value	F_{ST}
Iran/Asia	3.619	0.003	0.033 *	9.148	0.0190 *	0.78007	0.9090 ns	0.011
Iran/Middle East	3.528	0.004	0.167 ns	6.477	0.0490 *	0.737	0.0000 ***	-0.005
Iran/Far East	3.486	0.01537	0.0030 **	7.833	0.0010 **	0.837	0.0000 ***	0.009
Iran /Africa	3.699	0.010	0.095 ns	5.46011	0.1000 ns	0.62838	0.1300 ns	0.064
Iran /Europe	3.517	0.007	0.044 *	7.540	0.0210 *	0.81567	0.0000 ***	0.018
Iran /N. America	3.136	0.044	0.000 ***	7.475	0.0000 ***	0.89423	0.0000 ***	0.127
Iran /S. America	3.874	0.004	0.164 ns	5.645	0.1790 ns	0.70417	0.0090 **	0.019
Middle East/ Far East	3.391	0.016	0.002 **	8.033	0.001 **	0.792	0.000 ***	0.049
Middle East /Africa	3.333	0.006	0.045 *	7.949	0.007 **	0.803	0.000 ***	0.020
Middle East / Europe	3.433	0.006	0.020 *	8.707	0.003 **	0.867	0.000 ***	0.036
Middle East / N. America	3.212	0.021	0.001 **	8.458	0.001 **	0.903	0.000 ***	0.073
Middle East / S. America	3.467	0.012	0.085 ns	6.025	0.069 ns	0.751	0.035 *	0.046
Far East/Africa	3.381	0.015	0.000 ***	8.6825	0.000 ***	0.818	0.000 ***	0.016
Far East/ Europe	3.439	0.007	0.005 **	9.257	0.000 ***	0.809	0.000 ***	0.005
Far East/ N. America	3.295	0.040	0.000 ***	9.011	0.000 ***	0.853	0.000 ***	0.137
Far East / S. America	3.453	0.001	0.238 ns	7.674	0.156 ns	0.828	0.201 ns	-0.020
Africa / Europe	3.439	-0.00016	0.371 ns	7.341	0.2830 ns	0.85906	0.0060 **	-0.0004
Africa / N. America	3.012	0.044	0.000 ***	7.287	0.0000 ***	0.87686	0.0010 **	0.288
Africa / S. America	3.756	0.006	0.217 ns	4.992	0.2380 ns	0.60920	0.1520 ns	0.044
Europe / N. America	3.187	0.067	0.000 ***	8.246	0.0000 ***	0.88931	0.0000 ***	0.213
Europe / S. America	3.517	0.007	0.062 ns	7.384	0.055 ns	0.898	0.000 ***	0.019
N. America / S. America	3.103	0.033	0.000 ***	7.356	0.000 ***	0.880	0.000 ***	0.122

23 Ks, Kst*, Z* and Snn are test statistics of genetic differentiation [30]; F_{ST} examines the extent of genetic differentiation between geographical isolates; *:
24 0.01<P<0.05; **: 0.001<P<0.01; ***: P <0.001; ns: not significant.

1 Analysis of CP gene sequences using Kst*, Z and Snn test
 2 statistics revealed that most of the geographical PVY popula-
 3 tions (including Asian, African, European, North American
 4 and South American PVY populations) were genetically dis-
 5 tinct (Table 5). However, the highest F_{ST} values were obtained
 6 when North American PVY population was compared with
 7 African and European populations. The extent of genetic
 8 diversity between most of the geographical population pairs
 9 was little or moderate ($F_{ST} < 0.065$). The exception was North
 10 American PVY population, which had great genetic differ-
 11 ences with most of the other populations. For example the
 12 comparison of Iranian PVY population with the North Ameri-
 13 can population and other geographical populations revealed
 14 F_{ST} values of 0.127 and at most 0.064, respectively (Table 5).

15 Based on these test statistics, geographical isolation may
 16 have played a role in PVY population structure especially
 17 in North America.

18 Considering F_{ST} values, the clear distinction of genetic
 19 populations of PVY can be achieved by phylogenetic analysis,
 20 host relevance and geographical distribution, respectively.

22 *Examination of departure from neutrality*

23
 24 The patterns of nucleotide polymorphism in the CP gene
 25 of PVY isolates were estimated using Tajima's *D*, Fu and
 26 Li's *D** and *F** statistical tests (Table 6). The negative values
 27 of these test statistics would suggest for purifying selec-
 28 tion and/or a recent population expansion (Tajima, 1989;
 29 Tsompana *et al.*, 2005). On the other hand, positive values
 30 would suggest for balancing selection and/or a reduction in
 31 population size.

32 Results showed that Tajima's *D*, Fu and Li's *D** and *F**
 33 statistics were negative for most of the phylogenetic and

54 host populations of PVY. However, these negative values 54
 55 were only significant for PVY^O and PVY^{NTN} phylogenetic 55
 56 populations and PVY population isolated from potato (Ta- 56
 57 ble 6). Moreover, high level of haplotype diversity (Hd) and 57
 58 low rate of nucleotide diversity (π , except for PVY^C) were 58
 59 observed in the phylogenetic PVY populations. These find- 59
 60 ings suggested that these PVY populations may have been 60
 61 influenced by a recent population expansion. 61

62 Except the Asian PVY population, no significant de- 62
 63 parture from neutrality was found for PVY geographical 63
 64 populations according to Tajima's *D*, Fu and Li's *D** and 64
 65 *F** statistics (Table 6). Significantly negative test statistics 65
 66 obtained for the Asian population proposed the occurrence 66
 67 of a recent demographic expansion in this population. Also, 67
 68 high haplotype and high genetic diversities were observed 68
 69 in PVY geographical populations. 69

70
 71 *Recombination events*

72 Searching for evidence of recombination events in the CP 72
 73 gene sequences showed that most of the PVY isolates were 73
 74 recombinants. In this regard 60% (15 out of 25) of the Iranian 74
 75 PVY isolates were found to be recombinants. 75

76 Our statistical analysis identified at least 13 recombina- 76
 77 tion patterns in the CP gene sequences of PVY populations. 77
 78 (Table 7, Fig. 1). 78

79 Ten recombination patterns (events 1-10) appeared to 79
 80 be derived from parents from different PVY lineages (i.e. 80
 81 interlineage recombinants). In contrast, three recombina- 81
 82 tion patterns (events 11-13) were found to be derived from 82
 83 parents within the same PVY lineage (i.e. intralinear re- 83
 84 combinants) (Table 7). 84

85 Nine out of 13 recombination patterns were detected by 85
 86 at least four methods (events 1, 2, 3, 4, 5, 8, 9, 10, 12), hence 86
 87

89
 90 **Table 6. Representation of parameter estimates and test statistics for demographic trends in potato virus Y populations**

Population	Tajima's <i>Da</i>	Fu & Li's <i>D*</i>	Fu & Li's <i>F*</i>	Hd	π
O	-2.441**	-5.713**	-5.132**	0.997	0.030
C	-0.486ns	-0.443ns	-0.528ns	0.985	0.069
N	-0.069ns	-1.034ns	-0.856ns	1.000	0.033
NTN	-2.792***	-6.104**	-5.810	0.984	0.013
Potato	-1.555ns	-7.366**	-5.064**	0.994	0.061
Tobacco	-0.732ns	-2.278ns	-1.921ns	0.998	0.057
Other host	0.016ns	-0.474ns	-0.385ns	0.985	0.077
Iran	-1.072ns	-2.269ns	-2.220ns	1.000	0.072
Middle East	-0.392ns	-2.339ns	-1.965ns	0.995	0.055
Far East	0.197ns	-1.021 ns	-0.593 ns	0.988	0.058
Asia	-1.582ns	-6.566**	-4.837**	0.996	0.063
Africa	0.588ns	0.252ns	0.393ns	0.974	0.058
Europe	-0.0523ns	-1.332ns	-0.959ns	0.991	0.058
N. America	-0.809ns	-1.711ns	-1.602ns	0.954	0.042
S. America	0.337	0.502	0.526	1.000	0.081

53 ns: non-significant; Hd: haplotype diversity; π : nucleotide diversity per site; **: 0.001<P<0.01.

Table 7. Characteristics of recombination events detected in isolates of potato virus Y

Event	Frequency	Recombination site	Recombinant length	Parental sublineage ^a	Recombination detected programs ^b	P-value ^c
1	131	352-801	449	O*N	R.G.B.M.C.S.P.L.Se.	1.17E-08
2	30	331-555	224	O*Nw	R.G.B.M.C.S.P.L.Se.	1.79E-07
3	153	147-342	195	Nw*NO	R.G.B.M.C.S.P.L.Se.	7.55E-03
4	4	16-164	148	C*O	R.G.B.M.C.S.P.L.Se.	5.51E-03
5	53	410-607	197	O*NTN	R.G.B.M.C.S.P.L.Se.	2.51E-02
6	2	496-636	140	C*O	R.G.B.M.C.S.P.L.Se.	1.09E-02
7	5	8-344	336	C*NTN	R.G.B.M.C.S.P.L.Se.	2.05E-02
8	141	345-745	400	O*No	R.G.B.M.C.S.P.L.Se.	1.33E-09
9	103	166-708	542	O*NTN	R.G.B.M.C.S.P.L.Se.	4.65E-07
10	249	351-751	400	O*NTN	R.G.B.M.C.S.P.L.Se.	6.90E-08
11	1	1-439	438	NTN*NTN	R.G.B.M.C.S.P.L.Se.	1.31E-03
12	68	606-791	185	NTN*NTN	R.G.B.M.C.S.P.L.Se.	3.54E-02
13	1	1-345	344	NTN*NTN	R.G.B.M.C.S.P.L.Se.	1.70E-02

(a): Parental sublineage estimated by RDP Program; (b): Abbreviations used for recombination detection programs; R.: RDP, G.: GENECONV, B.: Bootscan, M.: Maxchi, C.: Chimaera, S.: SiSscan, P.: PhylPro, L.: Lard, Se.: 3Seq. Recombination-detecting programs representing significant signal showed in bold; (c): The greatest P-value calculated by the program for the recombination event. The highest reported P-value for the program showed in underlined in RDP4.

considered as “significant recombination events”. The four other recombination patterns were detected by less than four methods (events 6, 7, 11, 13), hence considered as “tentative recombination events” (Table 7).

Interestingly, our analysis revealed that most of the PVY recombinants were classified within the N:NTN clade with no preference between N-Europe and North American sub-clades. Contrary to this, many of the non-recombinant PVY strains were found within the PVY^{O/C} clade (Fig. 1). PVY^O and PVY^C clades contained only one and two recombination patterns, respectively (Table 7). The recombination pattern in the PVY^C lineage seemed to be derived from parents from PVY^O and PVY^C clades. On the other hand, the parents of PVY recombinants in PVY^O were from the same lineage or from PVY^O and PVY^C lineages (Fig. 1).

Six out of 13 recombination patterns (including patterns of 1, 2, 5, 7, 8, 11) were found among the Iranian PVY population. The recombination pattern No. 7 was the most

prevalent pattern within the Iranian PVY population (Fig. 1, Table 7).

Biological characterization of the PVY isolates

In order to compare the symptoms of PVY strains from divergent evolutionary lineages, four Iranian isolates representative of the PVY^O clade (isolates 21 and 221) and PVY^{NTN} clade (isolates 61 and IRAN) were inoculated to tobacco, pepper, tomato and *Chenopodium quinoa* plants.

Two weeks post-inoculation, all isolates induced vein clearing and mottling in *Nicotiana tabacum* cv. Xanthi and mottling and leaf rugosity in *Lycopersicon esculentum*. In addition to mottling, *Capsicum annuum*, developed the symptoms of vein necrosis by isolates 221 and IRAN, vein banding by isolate 21 and rugosity by isolate 61 (Table 8). None of the isolates infected *Chenopodium quinoa* Wild. Except *Chenopodium quinoa*, PVY was detected in all the inoculated plants using RT-PCR analysis.

Table 8. Biological characterization of Iranian potato virus Y isolates reported in this study

Isolate strain	21 PVY _O	221 PVY _O	IRAN PVY _{NTN}	61 PVY _{NTN}
<i>Capsicum annuum</i>	M, VB	M, VN	M, VN	R, M
<i>Nicotiana tabacum</i> cv. Xanthi	VC, M	VC, M	VC, M	VC, M
<i>Lycopersicon esculentum</i>	M, R	M, R	M, R	M, R

M: Mottle, R: rugosity, VC: Vein clearing, VN: vein necrosis, VB: Vein Banding.

Discussion

PVY is a typical example of RNA viruses that benefits the high mutation and recombination rates for adaptation and survival in different hosts and various environments (Boonham *et al.*, 2002; Hu *et al.*, 2009a). Several studies have attempted to determine the genetic structure of PVY populations in Europe, North America and Japan (Glais *et al.*, 2002; Lorenzen *et al.*, 2006; Schubert *et al.*, 2007). In this work, we used the CP gene sequences to compare the

1 genetic structure of PVY populations in Iran with the world
2 populations. We used CP gene, for which a large number of
3 international sequence data exists in GenBank.

4 Phylogenetic analysis using CP gene sequence revealed
5 the polyphyletic relationships of PVY strains reported from
6 34 countries across the world. Results showed that all the
7 PVY isolates could be arranged within 2 main clades desig-
8 nated as PVY^{O:C} and PVY^{N:NTN}. PVY^{O:C} and PVY^{N:NTN} could
9 be further differentiated into 2 subclades designated as PVY^O
10 and PVY^C, and N-Europe and North America, respectively
11 (Glais *et al.*, 2002; Moury *et al.*, 2002; Fanigliulo *et al.*, 2005;
12 Lorenzen *et al.*, 2006). The phylogenetic grouping of the
13 PVY isolates was not fully consistent with their geographical
14 distribution. For example, Iranian PVY strains were found
15 to be distributed throughout the tree, suggesting multiple
16 introductions of PVY virus isolates to Iran. Several PVY
17 strains within PVY^{O:C} and PVY^{N:NTN} clades formed a star-
18 like phylogeny, most probably as the consequence of recent
19 emergences with minimal selection. Similar star phylogenies
20 have been previously reported in the genetic structure of
21 several virus populations including cucumber mosaic virus,
22 pepino mosaic virus, wheat streak mosaic virus and turnip
23 mosaic virus (Roossinck *et al.*, 1999; Pagan *et al.*, 2006;
24 Dwyer *et al.*, 2007; Tomitaka *et al.*, 2007). Also some of the
25 PVY sub-clades in the PVY^{N:NTN} clades had low bootstrap
26 value. This is most probably due to the recombination events
27 as also considered by Ohshima *et al.* (2007).

28 Since the identification of first PVY recombinant strain
29 from Hungary (PVY^{NTN}) in 1980s, several others have been
30 reported in other countries (Le Romancer *et al.*, 1994; Glais
31 *et al.*, 1996; Boonham *et al.*, 2002). In the past decade, many
32 studies have shown the global prevalence of PVY recom-
33 binants (Chrzanowska, 1991; Glais *et al.*, 2002; Lorenzen *et al.*
34 *et al.*, 2006; Hu *et al.*, 2009b). In Europe, the non-recombinant
35 strains of the PVY^O clade have been largely replaced by
36 the recombinant strain of PVY^{NTN} (Boonham *et al.*, 2002).
37 However, in North America, the PVY^O strain has remained
38 the predominant strain in potato (Piche *et al.*, 2004; Bald-
39 auf *et al.*, 2006; Karasev *et al.*, 2008; Gray *et al.*, 2010). Our
40 analysis revealed that PVY^O and PVY^{NTN} were predominant
41 strains in Iran. Comparing results of this study with those
42 published earlier (Pourrahim *et al.*, 2007; Hosseini *et al.*,
43 2011) suggested that during the past decade the prevalence
44 of PVY^{NTN} was significantly increased within Iranian PVY
45 population.

46 The genetic diversity of the CP gene was varied among
47 different clades of the PVY phylogenetic tree, thereby some
48 clades contained more daughter subclades than others. The
49 combination of high haplotype diversity and low genetic di-
50 versity in phylogenetic PVY populations could be explained
51 by a recent population expansion after a genetic bottleneck
52 (Grant and Bowen, 1998; Tsompana *et al.*, 2005). However,
53 relatively long terminal branches of the PVY^C isolates may

suggest that they have accumulated mutations over a long 54
time. All statistical analyses clearly suggested the significant 55
genetic differences between pairs of the main PVY phylo- 56
genetic clades. The extent of this genetic variation between 57
phylogenetic populations was considerable as shown by F_{ST} 58
values. Also, our results revealed that PVY genetic diversity 59
may have resulted from host adaptation, which is in agree- 60
ment with previous reports (Schubert *et al.*, 2007; Ogawa 61
et al., 2008). However, the extent of the genetic variability 62
between pairs of host PVY populations was low ($F_{ST} < 0.1$). 63

Besides host adaptation, some of the PVY genetic diversity 64
was found to have resulted from geographical distribution. In 65
this regard, the majority of the geographical PVY population 66
pairs could be genetically differentiated. However, the extent of 67
genetic differences between most of the geographical popula- 68
tions was low. Contrasting to Cuevas *et al.* (2012), our analysis 69
concluded the distinction of Middle East PVY population from 70
populations of Europe and Far East. According to F_{ST} values, 71
Iranian PVY isolates displayed the highest genetic differentia- 72
tion from isolates of North America ($F_{ST} = 0.127$). This finding 73
parallels the phylogenetic analysis, as no Iranian PVY isolates 74
grouped within the North American clade. Our statistical 75
analysis showed that North American PVY population had 76
the highest genetic distance with other geographical popula- 77
tions ($F_{ST} > 0.07$). Altogether, findings of this study confirmed 78
that geographical distribution and host adaptation have played 79
a role in shaping genetic diversity of the PVY populations. 80

The extent of the selection pressure acting on genes can 81
be evaluated by computing the dN/dS, the ratio of the non- 82
synonymous to the synonymous polymorphisms (Garcia- 83
Arenal *et al.*, 2001). The global dN/dS ratio of the PVY CP 84
gene was less than 1 (~0.08) revealing the exertion of high 85
negative selection pressure on this gene. The negative selec- 86
tion pressure restricts the variability PVY CP gene. Different 87
dN/dS values found on the CP genes of distinct PVY popula- 88
tions may suggest that these populations have experienced 89
different constrains. 90

Analysis of dN/dS revealed that the CP gene sequence 91
of PVY^C population experienced higher negative selection 92
pressure compared to that of PVY^{NTN} and PVY^O strains. This 93
is consistent with low CP gene sequence diversity ($\pi = 0.069$) 94
of the PVY^C population as also considered by others (Schu- 95
bert *et al.*, 2007; Ogawa *et al.*, 2008). The low diversity of CP 96
gene in PVY^C could also be explained by Muller's ratchet 97
phenomenon, i.e. the stochastic loss of genotypes (Garcia- 98
Arenal *et al.*, 2001). 99

In most cases when phylogenetic, host or geographical 100
PVY populations were considered, Tajima's D and Fu and 101
Li's D and *F*-test yielded negative values. The negative values 102
obtained in these tests in combination with high haplotype 103
diversity and overall low nucleotide diversity of CP gene 104
suggested that PVY phylogenetic population has undergone 105
a recent population expansion or background selection. 106

1 High rates of haplotype and genetic diversity in the PVY
 2 host and geographical populations could reduce the risk of
 3 extinction and points to the evolutionary potential of these
 4 populations for adaptation into the diverse environments.
 5 Recombination has been considered as an important
 6 source of genetic variation in potyviruses (Gibbs and
 7 Ohshima, 2010). For example, 76% of isolates of the poty-
 8 virus turnip mosaic virus were reported to be recombinant
 9 (Ohshima *et al.*, 2007). Our analysis also showed the occur-
 10 rence of recombination events in the CP gene of PVY isolates
 11 as also considered by others (Glais *et al.*, 2002; Moury *et al.*,
 12 2002; Lorenzen *et al.*, 2006). Besides the six recombination
 13 patterns identified in the earlier studies (Moury *et al.*, 2002;
 14 Fanigliulo *et al.*, 2005; Lorenzen *et al.*, 2006), we identified
 15 seven new patterns of recombination in the CP gene of the
 16 PVY strains. Our findings also conclude that recombination
 17 in the CP gene is a common and frequent force driving the
 18 PVY evolution.
 19 Biological features of PVY strains have been described in
 20 several previous studies (Chrzanowska, 1991; Le Romancer
 21 *et al.*, 1994; dAquino *et al.*, 1995; Boonham *et al.*, 2002; Bal-
 22 auf *et al.*, 2006). Tomato and tobacco plants can be infected
 23 by most PVY isolates (Stobbs *et al.*, 1994). Commonly, PVY
 24 isolates of pepper cannot infect potato plants systemically
 25 (McDonald and Kristjansson, 1993; dAquino *et al.*, 1995).
 26 Whereas members of the PVY^O and PVY^C populations could
 27 induce mottling in pepper plants (McDonald and Kristjans-
 28 son, 1993; Valkonen *et al.*, 1996), PVY^{NNP} reported from Italy
 29 has been the only isolate inducing vein necrosis in pepper
 30 (Fanigliulo *et al.*, 2005). Our experimental data revealed
 31 that the two Iranian PVY isolates (PVY-Iran and PVY-221)
 32 could also induce vein necrosis symptom in pepper, hence
 33 they were biologically similar to the PVY^{NNP} isolate (dAquino
 34 *et al.*, 1995).
 35 In this study we described the structure and dynamics
 36 of the PVY populations in the world, with a focus on the
 37 PVY isolates from Iran. Evolutionary studies showed that
 38 PVY populations could be represented into discrete line-
 39 ages. Geographical origin, mutation, recombination and
 40 host adaptation were the main sources of genetic variability
 41 to shape the population structure of PVY. Strong negative
 42 selection pressure on the CP gene favors the preservation
 43 of wild type strains in the nature. These findings argue that
 44 the CP gene is an effective indicator to study the genetic
 45 diversity and evolution of PVY populations. Iranian PVY
 46 population displayed high haplotype and nucleotide diver-
 47 sities in the CP gene. The high nucleotide diversity within
 48 Iranian PVY population is probably due to the introduction
 49 of distant strains belonging to PVY^O, PVY^C and N-Europe
 50 phylogenetic populations, as also shown in the phylogenetic
 51 tree. Altogether, phylogenetic, biological and sequence di-
 52 versity analyses of the CP gene suggested that Iranian PVY
 53 isolates constitute a diverse population. Our analysis also

proposes that the PVY^O strains have been largely replaced 54
 by the recombinant strains of PVY^{NTN} within the Iranian 55
 PVY population during the last decade. The genetic vari- 56
 ability of the PVY populations reported in this study may 57
 provide foundation to improve control strategies of PVY in 58
 different crops. 59
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