Evolutionary Characterization and Genetic Structure of Iranian Isolates of *Beet Necrotic Yellow Vein Virus* Population Based on p25 Protein

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**ABSTRACT**

To study molecular evolutionary characteristics and genetics of beet necrotic yellow vein virus (BNYVV) isolates population from Iran, nucleotide sequences of p25 and coat protein (CP) were determined and the amino acids sequences thus deduced were analyzed using phylogenetic and population genetics methods. A survey of BNYVV in Iran indicated the infection of 288 collected samples out of 392 samples in most beet growing areas and that most of the isolates (92%) were of the *A*-type and the rest of isolates (8%) were *P*-type. Our molecular evolutionary analysis showed that CP was highly conserved but allowed to assign all isolates to three distinct groups. Different parts of p25 coding regions were under different evolutionary constraints. The most positive selection was detected at the position 68, the second amino acid of the tetrad motif. Iranian isolates were found to cluster with European isolates into three distinct clusters based on p25 sequences. Population genetics analysis revealed that BNYVV populations have low differentiation (*Kt*= 3.97145) and low diversity (*π*= 0.006, *Hd*= 0.860) with frequent gene flow indicating lack of phylogeographic structure between populations.

**Keywords**: Evolution, Phylogenetic analysis, Plant virus, Positive selection.

**INTRODUCTION**

*Beet necrotic yellow vein virus* (BNYVV) causes an economically important disease of sugar beets called rhizomania, which is distributed worldwide (Tamada and Baba, 1973). It was initially reported from Italy (Canova, 1959) and is transmitted by the soil-borne plasmodiophorid *Polymixa betae* (Beemster and de Heij, 1987). Rhizomania typical symptoms include massive lateral root proliferation resulting in a beard-like appearance, constriction of the main taproot and stunting of the infected plant. Leaves may become flabby and wilt without discoloration. Sometimes, there is a proliferation of smaller leaves generated at the crown. Because infected roots are inefficient in water and nutrient uptake, general foliar symptoms are similar to water stress or nitrogen deficiency. The disease causes economic loss of sugar beet by reducing yield. It can cause a decrease in root yield of 30-90% and reduce sugar yield up to 70% (Yardimci and Çulal Kiliç, 2011).

BNYVV is the type species of a rod-shaped particle genus of *Benyvirus*. It contains four or five plus-sense single-stranded genomic RNAs with 10 to 11 open reading frames (ORFs) and 5′-cap and a 3′-poly (A) tail (Tamada and Abe, 1989; King *et al.*, 2012). RNA 1 encodes the RNA-dependent RNA polymerase; RNA 2
encodes coat protein (CP), the triple block involved in cell-to-cell movement and a 14 KDa protein, which is a silencing suppressor; RNA 3 has three ORFs encoding p25 and two other small proteins (King et al., 2012). P25 is responsible for symptom development and previous studies have shown that recently occurring resistance-breaking isolates possess increased p25 variability (Thiel and Varrelmann, 2009). Also, previous studies suggested that p25 is involved in this inhibition of virus translocation in the taproots of the resistant cultivar Rizor (Tamada et al., 1999). Other studies demonstrated the role of RNA3-encoded p25 in the exacerbation of virus-induced symptoms on inoculated leaves (Jupin et al., 1992; Tamada et al., 1999; Klein et al., 2007) and the role of its subcellular localization (Vetter et al., 2004). RNAs 4 and 5 each contain one ORF; RNA 4 encodes p31 that is involved in efficient vector transmission, symptom severity and RNA silencing suppression; RNA 5 encodes p26 and is a pathogenicity factor (Link et al., 2005; King et al., 2012). RNA-5-containing isolates have been reported from France (Koenig et al., 1997), Kazakhstan (Koenig, 2000), China (Li et al., 1999), the UK (Harju et al., 2002) and Japan (Tamada et al., 1996).

Based on molecular characteristics, including RNA composition and nucleotide sequence divergence (Koenig et al., 1995), BNYVV has been classified in three major pathotypes designated as A, B and P. Type A is widespread in most European countries, USA, Iran, China and Japan whereas the B type is more restricted and mainly found in Germany, France and in some cases in Sweden, China and Japan (Schirmer et al., 2005; Pavli et al., 2011). P-type isolates have been reported from France, UK, Kazakhstan (Schirmer et al., 2005) and recently from Iran (Mehrvar et al., 2009).

Molecular diversity of BNYVV was studied for the first time by single-strand conformation polymorphism (SSCP) and restriction fragment polymorphism (RFLP) analyses that confirmed the existence of three major groups of BNYVV isolates (Kruse et al., 1994; Koenig et al., 1995). Subsequent studies were based on amino acid sequences so that aligned amino acid sequences of p25 showed some key residue substitutions in different types of BNYVV isolates (Miyanishi et al., 1999). Phylogenetic analysis of CP, p25 and p26 amino acid sequences collected from worldwide specimens have shown high variability in p25, especially within amino acid residues 67-70 (Schirmer et al., 2005). A survey detected BNYVV infection in six different regions of China and revealed the presence of A- and B-type isolates similar to European isolates (Li et al., 2008).

Rhizomania disease symptoms were first reported in sugar beet fields in Fars province of Iran in 1996 (Izadpanah et al., 1996), where subsequent soil bioassays indicated the widespread occurrence of *P. betae* (Kamran et al., 2000). In other researches, the main Iranian sugar beet cultivation areas were surveyed for the presence of BNYVV and two other soil born viruses (*Beet virus Q* and *Beet black scorch virus*) and their soil borne vector, *P. betae* using serological methods (Farzadfar et al., 2007) and RT-PCR (Shahnejat-Bushehri et al., 2006). In addition, some researchers tried to find and tag rhizomania-resistant genes (Nouhi et al., 2008; Salari et al., 2008). In another study, sequence analyses were conducted on the coat protein gene (CP), and parts of RNA3 and RNA4 of an Iranian strain of BNYVV from the Fars province. Sequence alignments of CP with other isolates showed closed similarities at nucleotide and amino acid levels with BNYVV pathotype A isolates; S from Japan, and YU2 from Yugoslavia (Sohi and Maleki, 2004). Recently studies revealed that RNA-5 was not found in Iranian isolates (Mehrvar et al., 2009). Knowledge of evolutionary history and the genetic structure of virus populations provides different levels of information about ecology and epidemiology that are important in the selection of control strategies. Virus-resistant plants offer the most cost effective strategies for rhizomania disease control. Therefore, the effective use of BNYVV-resistant cultivars and management of rhizomania requires a better understanding of the genetic structure and evolutionary trajectories of the virus.
population (Richards et al., 2005). Population diversity suggests that this virus is capable of rapid evolution and adaptation in response to changing agricultural practices. DNA sequence data can be used to infer the process of population diversity and to identify evolutionary forces that have shaped the genetic structure of virus population (Canova, 1959). Also, evolutionary characterization of the viruses is essential to develop better strategies for controlling all infections. Since the first report of BNYVV in 1952, different studies were performed on various aspects of rhizomania disease and BNYVV. However, there is little information about evolutionary characteristics, population genetic structure and parameters of BNYVV populations. This study hopes to fill some of these gaps.

**MATERIALS AND METHODS**

**Virus Isolation and Sequences Sources**

Sugar beet roots with rhizomania-like symptoms of 392 samples of sugar beet were collected from most regions of Iran, which are listed as follows: Ardabil, North and South Azarbeijan, Khorasan Razavi and Shomali, Semnan, Ghazvin, Zanjan, Ilam, Kermanshah, Hamadan, Fars and Kerman Provinces (Figure 1). Detection of BNYVV and amplification of the p25 coding region of all isolates were carried out by means of RT-PCR as described by Meunier et al. (2003). Also, the CP coding region of an isolate was amplified by the same method. The nucleotide sequences of the PCR products were obtained by using an ABI377 sequencer (Applied Biosystems, USA). The nucleotide sequences of p25 coding region were submitted to the GenBank under Accession Nos. AM745439-AM745638, FJ230957-FJ230958 and FM210563-FM210625; and the nucleotide sequence of the CP coding region of IR-GR1 were catalogued in the GenBank under Accession No. FM210683 and were compared with other 45 sequences obtained from GenBank including two Iranian Isolates (AF522462 and AY277887; Jahromi et al., 2005; Sohi and Maleki, 2004) and CP sequences of other
countries. The sequences of other isolates used in this study were obtained from GenBank and their accession numbers are presented in Figure 2.

Multiple Sequence Alignment and Phylogenetic Analysis

Multiple sequence alignments were performed using Muscle (Edgar, 2004) with the default parameters. The most appropriate model was determined using the Bayesian Information Criterion (BIC) implemented in MEGA version 5.0 program (Tamura et al., 2011). The alignments were used to calculate genetic distance by T92+G method (Tamura, 1992), which was selected by the lowest BIC. Phylogenetic analyses were done by the neighbor-joining methods using MEGA 5.0 under the assumption of the substitution models proposed. The robustness of inferred evolutionary relationships was assessed by 1,000 bootstrap replicates.

Molecular evolution of p25 neutrality test, namely Tajima’s D test was run using DnaSP 4.0 (Rozas et al., 2003). Tajima’s D was used to test if the sequence variant distribution followed the model of neutral evolution. Also, using the MEGA 5.0 program (Tamura et al., 2011), positive selection was inferred by the counting method described by Nei and Gojobori and, later on, by Suzuki and Gojobori (Nei and Gojobori, 1986; Suzuki and Gojobori, 1986).

Figure 2. Phylogenetic analysis of BNYVV based on CP (A) and p25 coding region (B) using neighbor-joining method and T92+G model with 1,000 bootstraps (using MEGA 5.0 software). Black square, circles and triangles correspond to A-, B- and P-type isolates, respectively. All Iranian isolates were shown as underlined. The tetrad motifs of p25 are indicated in parentheses.
According to these methods, the phylogenetic tree of sequences analyzed was used. Selective pressures on each codon were evaluated using the difference between nonsynonymous ($d_N$) and synonymous ($d_S$) substitution rates per codon. Values of $d_N - d_S < 0$, $= 0$ or $> 0$ indicate negative selection, neutral evolution and positive selection, respectively. Estimates of the difference in substitution rates were made from a phylogenetic tree inferred using the Neighbor-Joining algorithm with distances corrected under the T92+G model.

**Population Parameter Estimation**

The total population of Iranian BNYVV isolates was divided into 10 subpopulations based on different sampling regions. Within and between populations, diversities were estimated according to Nei (Nei, 1987) based on the T92+G model. Evolutionary analyses were conducted in MEGA version 5.0 (Tamura et al., 2011). The numbers of base substitution per site from averaging over all sequence pairs within each population and between populations were calculated and standard errors were obtained by a bootstrap procedure (500 replicates). The rate variation among sites was modeled with a gamma distribution (Shape parameter = 0.2). The analysis involved 262 nucleotide sequences. Codon positions included were 1st+2nd+3rd and all positions containing gaps and missing data were eliminated so that there were a total of 660 positions in the final dataset. Phylogenetic relationships among different BNYVV populations from Iran were constructed using Neighbor-Joining (NJ) method based on between population distances. Genetic differentiation between populations was examined by Z test which represents a sequence-based statistical test for genetic differentiation (Hudson et al., 1992). The extent of genetic differentiation or the level of gene flow between populations was measured by estimating $F_{st}$ (the interpopulational component of genetic variation or the standardized variance in allele frequencies across populations). $F_{st}$ ranges from 0 to 1 for undifferentiated to fully differentiated populations, respectively. Normally, an absolute value of $F_{st}$= 0.33 suggests infrequent gene flow. The statistical tests for genetic differentiation, estimation of $F_{st}$, number of haplotypes (h) and haplotype diversity ($H_d$) were performed by DnaSP 4.0 (Rozas et al., 2003). Potential recombination events between nucleotide sequences of p25 coding regions were assayed with SimPlot 3.5.1 (Lole et al., 1999).

**RESULTS**

**Phylogenetic Analysis**

The BNYVV was detected in 288 out of 392 sugar beet root tested samples. Most of these isolates were identified as $A$-type (92%) and the rest of the isolates were $P$-type. Although an Iranian isolate, M1ar (Jahromi et al., 2005) was clustered in $B$-type group, the $B$-type isolate was not observed among our 288 samples. The phylogenetic tree based on CP sequences displayed three main lineages associated with the three types. M1ar, Fars and IR-GR1 isolates were clustered to $B$, $A$ and $P$-type groups, respectively. Although the M1ar isolate was grouped in the $B$-type cluster, it differed by an S 62T substitution (T62N103F172). The $B$-type group was more diverse and included all $B$-type isolates from Europe and Asia (Iran, China and Japan). East Asian $B$-type isolates differed by an N126H substitution that was not seen in other $B$-type isolates. The Fars Isolate was grouped in the $A$-type group, which was more conserved. All CP (28 sequences) sequences within the $A$-type group in this study were identical. In addition, the $P$-type isolates were conservative and the IR-GR1 isolate was identical in sequences to the European $P$-type isolates. These results indicated the occurrence of three types of BNYVV isolates in Iran.
Phylogenetic analysis of p25 revealed high variability within a four-amino acid sequence that is referred to as the tetrad in position 67-70, downstream of the nuclear localization signal motif (Vetter et al., 2004) and upstream of the zinc finger motif (Jupin et al., 1992). The phylogenetic tree of the p25 showed two main groups with four subgroups (Figure 2-B). Iranian isolates mostly belong to subgroup p25-Ia together with common European isolates. Some isolates from west of Iran (Kermanshah and Ilam) were clustered with those of Kazakhstan, the Netherlands and Italy, which all had ALHG motif (Figure 2-B). The Nahavand1 isolate was similar to the Spanish isolate with VCHG tetrad but had the tetrad VLHG that has not been seen in other regions of Iran. An isolate from Khorasan Razavi (Mas2) with the tetrad AHHG stayed together with isolates M1ar from Hamadan and the Chinese isolate Jiu. Totally, the widespread tetrad AXHG was found to occur most commonly including the isolates of Fars, Kermanshah2, Nyshabour1 (ACHG), Ghazvin1 (AFHG), Ardabil1, Bisotoon1, Barsir1, Biz2, Kangavr4 (AYHG) which were grouped with European and American isolates. Isolate Miandoab3 (AFHR) was similar to NL3 isolate from the Netherlands. The p25-Ib cluster was associated with East Asian isolates and both of the subgroups, p25-Ia and p25-Ib, were related to the A-type isolates except the isolate M1ar which was grouped in B-type and belongs to the p25-Ia cluster. The distinct cluster p25-IIa matched completely with the P-type isolates with the conserved motif SYHG including two isolates from the East (IR-GR1) and the West (Pars1) of Iran. Finally, the p25-IIb cluster was found to be related to B-type.

Molecular Evolution Analysis of p25

The nucleotide substitution pattern for p25 coding region showed that the rates of different transitional substitutions varied from 11.46 to 23.95, and those of transversional substitutions varied from 3.93 to 6.75. The nucleotide frequencies were: 0.229 (A), 0.35 (T/U), 0.203 (C), and 0.217 (G). The transition/transversion rate ratios were \( k_1 = 2.734 \) (purines) and \( k_2 = 3.547 \) (pyrimidines). The overall transition/transversion bias was \( R = 1.497 \), where \( R = \frac{[AGk_1 + TCk_2]}{[(A+G)(T+C)]} \).

Mean (relative) evolutionary rates are shown for each site next to the site number (Figure 3). These rates are scaled such that the average evolutionary rate across all sites is 1. This means that sites showing a rate< 1 are evolving more slowly than average whereas those with a rate> 1 are evolving faster than average. These relative rates were estimated using the Jones-Taylor-Thornton (1992) model (+G) (Jones et al., 1992). A discrete Gamma (+G) distribution was used to model evolutionary rate differences.

![Figure 3. Mean (relative) evolutionary rates over p25 amino acid sequence.](image-url)
among sites (5 categories). The ML estimate of the gamma shape parameter was 0.2561. The maximum Log likelihood for this computation was -792.779. The analysis involved 262 amino acid sequences. No recombination events were detected in the p25 coding region of Iranian BNYVV isolates.

Positive Selection

The high genetic stability of viruses can be attributed to negative or purifying selection to maintain the functional integrity of the viral genome. The degree of negative selection in genes, or the degree of functional constraint for the maintenance of the encoded protein sequence, can be estimated by two standard approaches. The first approach, which is based on Tajima’s D method, tests if the number of segregating sites in the sample significantly departs from the neutral expectation. The test suggests that p25 is evolving neutrally in Iran but this test showed a strong positive selection for worldwide samples. These results may be due to sampling error. Therefore, it is convenient to use an alternative test. To overcome this drawback of Tajima’s D, the second approach was to evaluate the neutrality by the index of the nucleotide diversities by differences between nonsynonymous and synonymous positions (dN-dS). The dN-dS index below zero is consistent with negative selection against protein change. In contrast, a dN-dS index above zero may be an indication that adaptive or positive selection is driving gene divergence. In this study, comparisons of all Iranian p25 coding regions showed heterogeneous mean dN-dS values (from -2.94 to 3.63). The positions 68 and 4 showed the highest and the lowest dN-dS values, and were under highly positive and negative selection, respectively (Figure 4).

Population Genetic Structure

All sequenced p25 collected from Iran were grouped into 10 subpopulations based on geographical regions (Table 1). Totally 27 haplotypes out of 262 samples were detected with haplotype diversity ranging from 0.466 to 1.0 and nucleotide diversity from 0.001 to 0.009. Estimation of within population distance (Dw) and Π showed that the population of Ardabil and the populations of Fars and Kerman represented the most and the lowest population diversities in Iran, respectively. However, the lowest value of haplotype diversity was for Ardabil population indicating the presence of a few haplotypes with high nucleotide differences. Estimation of evolutionary divergence over sequence pairs between groups showed the highest distance between the populations from Azarbaeijan Gharbi and Hamadan and that on the whole Azarbaeijan Gharbi was more divergent compared to other populations (data not shown). The number of base substitutions per site obtained by averaging overall sequence pairs was 0.006±0.002. The mean evolutionary diversity within subpopulations was 0.005±0.001. Mean interpopulational evolutionary diversity was 0.005. The coefficient of evolutionary differentiation was 0.246±.041. Result of Z test strongly supported the hypothesis that all subpopulations were genetically undifferentiated (P= 0). Measurement of the coefficient Fst also showed the existence of gene flow as follows: the values of Fst between the subpopulation from Ardabil and
other subpopulations, between Fars, Kerman and Khorasan Razavi, and between Khorasan shomali, Semnan and Azarbaeijan Sharghi were all > 0.33 (data not shown), which is an indication of infrequent gene flow. The absolute values of $Fst$ for all other subpopulation pairs were < 0.33, suggesting frequent gene flow.

**DISCUSSION**

This study provides an overview of the relationships between BNYVV isolates collected from Iran and isolates from other countries. We compared molecular evolutionary characteristics of BNYVV based on CP and p25 sequences. Our analyses support the following conclusions:

First, BNYVV isolates were widely distributed in all beet growing regions in Iran. Second, our data showed that the isolates collected from Iran were very diverse including $A$ and $P$-types in addition to at least one previously reported isolate (M1ar), which was similar to $B$-type. Third, phylogenetic analysis of CP amino acid sequences collected worldwide revealed that this ORF was highly conserved but allowed to assign isolates to three distinct groups (Figure 2) consisting of $A$-type ($K_{17}T_{62}V_{102}S_{103}L_{172}$), $B$-type ($K_{17}S_{62}V_{102}N_{103}F_{172}$) and $P$-type ($R_{17}T_{62}I_{102}S_{103}L_{172}$) (Kruse et al., 1994, Lennefors et al., 2005). The high conservation in CP sequences collected worldwide showed that coat protein was required for basic function(s); stability or reproduction in BNYVV as shown for other conserved proteins (Boyko et al., 1992, Morozov and Solovyev, 2003, Ghazala et al., 2008) while most other plant viruses have variable CPs. Such evolutionary conservation occurs because mutations of these amino acids were harmful to protein function, and were lost over time. Also, low variability of CP could separate three distinctive lineages with different pathogenicities. Amino acid comparison of $A$- and $B$-type revealed that the three conserved amino acids in $A$-type isolates ($K_{17}T_{62}V_{102}S_{103}L_{172}$) were replaced by functionally similar amino acids in $B$-type isolates ($K_{17}S_{62}V_{102}N_{103}F_{172}$). Both $S$ and $T$ in position 62 were carbohydrate linking amino acid, $N$ and $S$ in position 103 were carbohydrate $N$-link and $O$-link, respectively. Although, $L$ and $F$ in position 172 were both hydrophobic, the residue $F$ was more so (Wimley and White, 1996).
Furthermore, P-type isolates had two conserved amino acids in comparison with the A-type; Furthermore, P-type isolates had two conserved amino acid substitutions in comparison with A-type: the R17K and I102V so that each pairs were functionally similar to each other (basic hydrophilic and hydrophobic, respectively). Therefore, we propose that these amino acid substitutions cannot be important for differences in pathogenicity among A-, B- and P-types. Although some substitutions had occurred along coat protein, they did not affect their function. The isolate M1ar which was clustered in B-type group and differed in position 62, had an S62T substitution, which was named pseudo-B-type group due to function similarity between S and T. According to the sequence comparison and phylogenetic analysis of p25, the diversity of BNYVV p25 in Iran is reported to be the highest in the world. Two distinct main groups were determined based on p25 sequence analysis. The p25-I included A-type isolates which was divided into two subgroups; p25-Ia that can be divided into some other subgroups based on their tetrad motif. A-type isolates were widespread and have been reported from Asia, Europe and America (Schirmer et al., 2005). Ten of the thirteen different known tetrad have been found in the Iranian isolates (Mehrvar et al., 2009), no VCHG, TYHG and ASHG tetrad were found in Iran, while the tetrad were found only in Europe, UK and China, respectively (Li et al., 2008). The tetrad AHYG (for Ir3 isolate) and VHHG (for Ir12 isolate) have not been reported previously in other countries (Schirmer et al., 2005, Li et al., 2008). The tetrad profile in this group was [AV][LCHFY][GR] for which the residue H69 was very conserved in all isolates, therefore, it seems to be involved in the active site of p25. The hydrophobic residues A and V were aliphatic neutral amino acids, therefore their replacement with each other could not change protein functionally but the previously reported V67A substitution results in higher virus accumulation (Koenig et al., 2009). The position 68 was the most divergent position; the residues L, C, F and Y were hydrophobic but H was hydrophilic and could change protein structure and function. Existence of H residue at position 68 is rare and is only observed in Iranian, Chinese and Dutch isolates. The residue G in position 70 was conserved and rarely replaced with R in the Miandoab3 isolate. The residue G was neutral but R was a polar hydrophilic positively charged residue whose replacement may affect protein function significantly. The subgroup p25-Ib included East Asian A-type isolates with the profile A [SHY]HG. The group p25-II was divided into two subgroups; p25-IIa included P-type isolates which were more conserved than other subgroups with a unique tetrad SYHG. These findings were in agreement with those that showed the high degree sequence identity among the P-type isolates from Iran and those of Kazakhstan (Mehrvar et al., 2009). Existence of S at the position 67 was important evidence. The residue S67 in P-type isolates was replaced by [AV]67 in A-type isolates since A was a hydrophobic amino acid and S was a polar hydrophilic amino acid which can be covalently linked with carbohydrates. This property may be the reason for P-type high pathogenicity. These results were in agreement with those of Chiba et al. (2008) who found that the p25 was an avirulence factor. Another group, p25-Ib, was completely matched with B-type isolates having the tetrad profile A[FHY]HR and did not include Iranian isolates even the isolate M1ar which grouped in B-type based on CP. Also, some Chinese isolates which were grouped into A- (Wu3) and B-type isolates (Bao and Hoh2) based on CP, were grouped into p25-Ib, p25-Ib and p25-Ia, respectively. These results showed reassortment events in mixed infections by both A- and B-type isolates. Finally, based on p25 phylogenetic analysis it was supposed that the origin of the Iranian
BNYVV was more likely from European isolates and was different from South Eastern Asian.

Estimating transitional and transversional substitutions showed that high transitional substitution events along p25 coding region caused a slow evolution rate. Molecular population genetics analysis based on Tajima’s test suggested that demographic change had occurred. In other words, negative Tajima’s D suggested that demographic forces were acting on the population and there had been a demographic expansion of BNYVV subpopulations resulting in low frequency polymorphism. A total of 31 segregating sites were identified along 405 positions representing more than 7.6% of the total number of sequence bases, indicating low frequency of segregating sites.

Estimation of mean evolutionary rate for each site showed that the positions 68, 67 and 70 (three positions of tetrad) had the highest evolutionary rates, respectively, which was confirmed by strongly positive selection. Therefore, these amino acids evolve more than those in the other positions. The lowest rates were related to the positions 89 (in the zinc finger motif), 109, 110 and 121. These rates for the nuclear localization signal motif (amino acids 57-62) and the zinc finger motif (amino acids 73-90) were very low (0.3-0.6) which indicate the important role of these motifs for virus evolution. The evolutionary rates for the first 13 N-terminal amino acids were very low that might be having major unknown functions. These results were confirmed by the index $d_{N}-d_{S}$, because a strongly positive selection was observed on tetrad and negative selection was detected on the third and fourth nucleotide, the zinc finger motif means any changes in this region might result in virus extinction. There was no positive or negative selection on the NLS site and it had neutral evolution with low evolutionary rate. We assume that the zinc finger site and conserved region in its N-terminal were more important than NLS. Furthermore, the most positive pressure on position 68 indicated the role of this position for virus survival and flexibility against evolutionary forces (Figure 4).

A good understanding of population genetic structure can support the design of an effective disease management program. Lack of phylogeographic structure (Figure 5) between all populations, except for Ardabil and Azarbaidjan Gharbi, indicated high gene flow among these populations that was supported by $F_{st}$ estimation. There was no gene flow between Ardabil population and other populations, between Fars, Kerman and Khorasan Razavi and between Khorasan Shomali and Azarbaidjan Sharghi. The last two pairs of population were geographically far away from the others but Ardabil population had no gene flow with its neighbor populations. Low differentiation ($K<4$) and high gene flow ($F_{st}>0.33$) among all populations except Ardabil and Azarbaidjan Gharbi, were usually interpreted as a signature of recent range expansion from a single source population.

The population also exhibited an overall low genetic diversity, averaging...
0.0047±0.0017 (Table 1). Similar genetic structure and diversity have been found in populations of most plant viruses studied so far (Li et al., 2008). Among populations of Iranian BNYVV isolates analyzed here, the population of Ardabil and Azarbeijan Gharbi showed significantly higher genetic diversity (π= 0.00932 and π= 0.00810, respectively) whereas other populations exhibited low diversity (Table 1). These results in combination with those of population distances indicated that all populations except Ardabil and Azarbeijan Gharbi were closely related and could be considered as one population. These data suggest that these populations most likely were derived via a founder event from a common ancestor and they only recently colonized and spread within the area of Iran. However, we cannot rule out the possibility that the low diversity (πT= 0.00602) and close evolutionary relationship among isolates could be due to selection by the host plant, because all the populations were collected from beet, or due to need of preserve the function of viral protein (García-Arenal et al., 2003).

Gene flow in the region suggests a significant potential risk for the spread of novel strains that might contribute to breakdown of resistance genes. Approaches such as using RNA silencing for breeding should be applied to those parts of genes with low variability and negative selection. Since p25 is a pathogenicity factor and is responsible for the production of rhizomania symptoms (Chiba et al., 2008), based on our findings on evolutionary rates and selective constraints along p25, it is proposed that breeding efforts could examine the possibility of the use of the zinc finger site and the p25 N-terminal for RNA silencing targeting.

REFERENCES


تئیین خصوصیات تکاملی و ساختار زنیک جمعیت جدایی‌های ایرانی ویروس زردی 

p25

نکوریک رگبرگ چندندر بر اساس پروتئین

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چکیده

به منظور مطالعه خصوصیات تکاملی و زنیک جمعیت جدایی‌های ایرانی ویروس زردی نکوریک رگبرگ چندندر (BNYVV) توایی نوکلئوتیدی 25 و پروتئین پوشنی (CP) آن تئیین شده و توایی آمیتوسیدی حاصل از آن با استفاده از روشهای فیلوژنیکی و زنیک جمعیت تجزیه و تحلیل شد. غربالگری در ایران آلودگی BNYVV 288 جدایی از 326 نمونه را نشان داد که حاکی از آلودگی بیشتر مناطق رشد چندندر است. بیشتر جدایی‌ها (92 درصد) از تیب P و ماپی (Hd= 0.860) و نتایج دادن گردید. که در این پژوهش 88 مشاهده گردید که دوین اسید آمینه از یک موئیف چهارتایی است. بر اساس توالی 25 جدایی‌های ایرانی با جدایی‌های اروپایی در سه گروه مشخص دسته‌بندی شدند. بررسی زنیک جمعیت نشان داد که السعوديةهای جمعیت BNYVV فرق کم (K= 3.97145) نتوانست با یکدیگر (H= 0.860) تفاوت بیانی باعث کنند و

جریان زنی با آبی دارند که نشان می‌دهد میان جمعیت‌ها ساختار فیلوژنیکی و گرافیکی وجود ندارد.