#### **Research Article**

# Population genetic structure of *Fusarium verticillioides* the causal agent of corn crown and root rot in Ilam province using **Microsatellite markers**

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Abstract: Analysis of genetic diversity of Fusarium verticillioides populations concludes different levels of information in management of crown and root rot disease in corn farms. Simple sequence repeat (SSR) markers were used to determine genetic structure and estimate genetic diversity in 77 F. verticillioides isolates from major producing areas in Ilam province, Iran such as: Dehloran, Mosiyan, Dasht Abas, Mehran, Eivan, Holeylan and Darreh Shahr during 2016 - 2017. Nine microsatellite (SSR) primer pairs revealed that the average number of alleles in populations were 34, the number of alleles in populations varied from 27 alleles in Dehloran and Mosiyan as the lowest to 40 alleles in Darreh shahr as the highest. Observed alleles (Na) number and effective number of alleles (N<sub>e</sub>) were higher in Mehran (N<sub>a</sub> = 1.860; N<sub>e</sub> = 1.463) compared to other populations. The genetic diversity (H<sub>e</sub>) was higher in Holeylan ( $H_e = 0.284$ ) and Shannon's information index (I) was also higher in Mehran (I = 0.436) but lower values were estimated for Mosiyan ( $H_e = 0.195$ ; I = 0.303). The lowest genetic distance was found between Dehloran and Mosiyan (0.013) while the highest genetic distance was revealed between Dehloran and Darehshahr (0.139). Total gene diversity (H<sub>t</sub>) and gene diversities between subpopulations (H<sub>s</sub>) were estimated at 0.292 and 0.249 respectively. Gene diversity attributable to differentiation among populations (G<sub>st</sub>) was 0.147, while gene flow (N<sub>m</sub>) was 2.890. Cluster analysis based on UPGMA showed the lowest genetic distance between Dehloran and Mosivan and then Dasht abas. The dendrogram indicated a high genetic distance between Darehshahr and the six remaining populations. Results from this study will be useful in breeding program of crown and root rot resistant cultivars and developing control methods for this disease.

Keywords: corn, genetic distance, root rot, SSR

#### Introduction

Corn Zae mays L. is the highest yielding produce among cereals and regarding cultivation area is next to wheat and rice, but stands first as forage.

Fusarium verticillioides (Saccardo) Nirenberg [Gibberella fujikuroi (Sawada) Wollenw] is a prevalent pathogen of agriculturally important (Leslie, 1991. 1995). Fusarium crops verticillioides has been associated with human and animal toxicities since it was first described in 1881 (Saccardo, 1881). This species and other anamorphs of G. fujikuroi are the most common fungi associated with corn production in North America and other temperate regions in the world

Handling Editor: Naser Safaie

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(White, 1999). Fusarium species are capable of causing seedling disease, root rots, stalk rots and ear rots of corn as well as damaging stored grain (Mohammadi and Mofrad, 2011). Although yield usually is not much affected, kernel infection by G. fujikuroi is of concern because of the loss of grain and seed quality and the potential occurrence of fumonisins and other mycotoxins (Munkvold and Desjardins, 1997). Members of F. verticillioides and F. proliferatum (Matsushima) Nirenberg (teleomorph: G. intermedia) belong to the Liseola section of Fusarium (Nelson et al., 1983), and are well-known pathogens of corn, causing stalk and ear rot (Nelson et al., 1981; Leslie et al., 1990) and also rice root rot and stunting of crown (Hsieh et al., 1977). Crown and root rot is primarily a seed borne disease. Soil temperature of 35 °C is most favorable for infection (Nyvall, 1999). Spores of this pathogen are distributed by wind between corn fields (Ooka and Kommedahl, 1977). Long distance distribution by infected seed (Mohammadi and Mofrad, 2011). Asexual sporulation is clearly the most successful and impressive reproductive mechanism because of the large number of conidia that is produced by a single colony (Kendrick, 2003). Symptoms of the disease are a white to pink or salmon-colored mold on the ear and brown to black lesions on the crown and root. Infection of developing corn kernels may occur through the silks. Through holes and fissures in the pericarp or at points where the pericarp is torn by the emerging seedling and as a result of systemic infection of the corn plant by F. verticillioides (Glenn et al., 2003). Fusarium verticillioides produces abundant, mostly singlecelled microconidia in long chains (Glenn et al., 2004). For more than 20 years, assessments of fungal pathogens have used multilocus markers to detect population genetics (Milgroom, 1996). DNA-based techniques have increasingly become a tool for understanding the genetic diversity and also phylogenetic relationships of Fusarium spp. Researchers have already worked on molecular variation in Fusarium spp. (O'Donnell, 2000; Datta et al., 2011). Ren et al. (2012) showed high polymorphism and high genetic diversity among F. verticillioides isolated

from corn using SSR markers. Plant pathologists should study the population genetic of plant because pathogens evolve pathogenic fungi, (Momeni and Nazari, 2016). Pathogens population must constantly adapt to changes in their environment to survive (McDonald, 1997). Defining the genetic structure of populations is a logical first step in studies of fungal population genetics because the genetic structure of a population reflects its evolutionary history and its potential evolve (McDonald, to 1997). Knowledge derived from genetic structure of pathogen populations has direct agricultural applications. For instance, the genetic variation maintained within a population indicates the speed at which a pathogen evolves (McDonald and McDermott, 1993). This information may eventually be used to predict how long control measures such as fungicides and resistant cultivars are likely to be effective (McDonald and Linde, 2002). Several studies have recorded the genetic distinction of the mating populations of the G. fujikuroi complex species, including vegetative compatibility (Leslie, 1993), isozyme analysis (Huss et al., 1996) and electrophoretic karyotyping (Xu and Leslie, 1996). Knowledge of genetic structure of F. verticillioides populations is useful for development of effective strategies in controlling the disease (McDonald, 2004). Several molecular techniques available for are investigating genetic variability among plant pathogens. populations of fungal Microsatellites also known as simple sequence repeats (SSRs), which are randomly distributed throughout the genome of fungi and other eukaryotes (Li et al., 2002; Wostemeyer and Kreibich, 2002; Sahran and Naef, 2008), provide a powerful tool for taxonomic and population genetic studies (Britz et al., 2002). Alleles vary according to the number of repeat units present but other mutations have also been shown to be responsible for allele length variation in SSR analysis (Burgess et al., 2001; Slippers et al., 2004). The objective of present study was to collect corn samples with crown and root rot symptoms from corn farms in Ilam province and to determine population genetic structure of F. verticillioides using SSR markers.

#### **Materials and Methods**

#### Samples and fungal isolates

Corn plants in panicle stages with symptoms of brown to black lesions on crown and root were randomly sampled from 35 corn fields (each field  $\geq 1$  ha) in seven different regions as populations in the west of Iran, Ilam province, during 2016 - 2017 growing season (Fig. 1., Table 1). A total of five symptomatic plants were collected per field.

Each sample were cut into 3 - 5 mm long pieces, were surface-sterilized with 0.5% sodium hypochlorite for 1 - 2 min and 70% ethanol for 30 seconds rinsed three times with sterile distilled water and air dried with sterile filter paper. The sterilized samples were placed on a general medium, potato dextrose agar (PDA), and Nash and Snyder medium (Taherkhani *et al.*, 1998; Jo *et al.*, 2008). Samples were incubated 3 - 5 days at 25 °C with a 12 - h photoperiod for production of

conidia. Nash and Snyder is a selective medium for Fusarium species and facilitates the formation of large, easily recognizable colonies (Chen et al., 2007). The fungal isolates were isolated and purified using the hyphal tip and single spore method (Hawker, 1950). Fusarium colonies were observed microscopically, those colonies identified as F. verticillioides, were transferred to carnation-leaf agar (CLA), potassium chloride agar (KCLA) and potato dextrose agar (PDA) Isolates were identified (Merck Company). species morphologically to based on characteristics of the macroconidia, microconidia, phialides, chlamydospores and colony growth traits (Leslie and Summerell, 2006). Obtaining 98 isolates of Fusarium section Lesiola, among which F. verticillioides was the most frequent (78.5% of the population), followed by F. proliferatum (21.5%). Identified isolates were stored for short time on SNA at 4 °C, but for long time stored in tube containing glycerol at 4 °C.



Figure 1 Geographical origins of the seven Fusarium verticillioides populations used in this study.

Table 1 Location and number of isolates in each Fusarium verticillioides population.

Populations	No. of isolates in population	Code of isolates	Location	Province
1	12	FvC1-FvC12	Dehloran	Ilam
2	10	FvC13-FvC22	Mosiyan	Ilam
3	9	FvC23-FvC31	Dasht abbas	Ilam
4	12	FvC32-FvC43	Darreh shahr	Ilam
5	12	FvC44-FvC55	Mehran	Ilam
6	11	FvC56-FvC66	Holeylan	Ilam
7	11	FvC67-FvC77	Eivan	Ilam

#### Pathogenicity test

For inoculation, 100 g of corn seeds soaked for 24 hours in distilled water sterilized at 121 °C for 20 minutes, then inoculated with  $1 \times 2$  cm mycelial block of 7 days-old cultures on PDA plates. The flasks containing the inoculated seeds were shaken daily to prevent clumping of inoculated seeds. Pathogenicity test was carried out on the susceptible local cultivars under greenhouse conditions. Germinated seeds of corn plants were planted in pots containing sterilized soil. The corn seedling was inoculated by adding propagules to the soils around plant roots, covered with white plastic bags to keep high humidity for 24 hours. Control treatments were corn plants that were grown in sterilized soil without inoculation. Twenty-five days after inoculation, plants were observed for the development of disease symptoms and F. verticillioides was reisolated from inoculated plants.

# **DNA extraction**

Liquid cultures were initiated by adding 2-4  $mm^2$  pieces of filter papers containing of F. verticilioides to 250 ml Erlenmeyer flasks containing 100 ml PDB medium (potato dextrose broth plus 2 g yeast extract per liter). Flasks were incubated at room temperature approximately 25 °C and shaken on a rotary shaker for 7 - 14 days. Mycelium was collected by filtration through sterile filter paper with a vacuum funnel. Mycelia were harvested, frozen and stored at -20 °C. DNA was extracted using modified Cetyl Trimethyl Ammonium а Bromide (CTAB) procedure (Doyle and Doyle, 1990). Mycelia were ground in liquid nitrogen and suspended in 2% CTAB extraction buffer (1.4 M NaCl, 0.1 M Tris-HCl, PH 8.0, 20 mM EDTA, 0.2% β-mercaptoethanol). Samples were treated with 5 units RNAse at 37 °C for 30 min and then extracted with chloroform isoamylalcohol 24:1 (v/v). DNA in the supernatant was precipitated with isopropanol, rinsed with ethanol, and adjusted to a final concentration of 20 ng/µl in TE (pH 7.4) and stored at -20 °C (Weising et al., 1991; Bayraktar, 2010).

#### SSR amplification and analysis

Nine SSR primer pairs (Table 2) were selected based on their high PIC as described by Ren et al. (2012). Primer aliquots for each marker were prepared by mixing equimolar amounts 20 pmoles of appropriate forward and reverse primer in  $1 \times TE$  (1 mM EDTA, 10 mM Tris-HCl, pH 8.0). PCR amplification was performed in a 25 µl reaction volume containing 2.5 µl of 10X PCR buffer, 1.5 mM MgCl<sub>2</sub>, 0.2 mM of dNTPs mix (100 mM of each dNTPs), 1 µl of each forward and reversed primer, 0.6 U of Tag polymerase with 25 ng of template DNA. Amplification was performed using Mnotach 60 thermal cycler., PCR conditions for SSR were as follows; one initial denaturation step at 95 °C for 3 min followed by 30 cycles of 94 °C for 60 s, annealing for 60 s (appropriate annealing temperature were used for each primer set, between 52 - 59 °C) and 72 °C for 60 s. The thermal cycles were terminated by a final extension of 5 min at 72 °C. Amplified products were resolved in 3% agarose gel at 60 V using Tris Boric Acid EDTA (1X TBE) buffer and stained with DNA Safe Stain at 0.5mg/ ml and photographed under UV Trans laminator with Gel Doc. Intas. A 1 kbp ladder was used as a molecular size standard.

#### Statistical analysis

Populations were defined as different geographic regions. The clear and intense amplified DNA bands were scored as binary digit code of "0" (for absence) and "1" (for presence), respectively. The pair-wise distance among the isolates was calculated from the binary matrix using the simple mismatch coefficient (Sneath and Sokal, 1973) that is recommended for haploid fungi (Kosman and Leonard, 2005). Genetic distance of the populations was estimated from the SSR data using the UPGMA, clustering method on the basis of Nei's (1978) unbiased genetic distance. All data analyses were performed using POPGENE ver. 1.31 (Yeh et al., 1999), Gen Alex ver. 6.501 (Peakall and Smouse, 2012) and NTYsys ver. 1.07c (Rolhf, 1990).

Genetic similarity between pairs was estimated using Jaccard's similarity coefficient. Similarity coefficients were used for the construction of UPGMA (Unweighted Pair Group Method with Arithmetic Average) dendrogram (Rohlf, 1990). For each primer pair, polymorphic information content (PIC), marker index (MI) was calculated. The polymorphic information content (PIC) was calculated using PICi = 2fi (1 - fi), where i is the information of marker<sub>i</sub>, fi is the frequency of the amplified allele (presence of fragments) and (1 - fi) is the frequency of the null alleles (Roldan-Ruiz et al., 2000). The genetic variation was measured in terms of genetic diversity and was computed by averaging PIC estimates over all loci (Nei and Li, 1979). The marker index (MI) was calculated by MI = PIC\*EMR, where EMR is the "effective multiplex relationship" given by the product of the total number of fragments per primer and the fraction of polymorphic fragments (Varshney et al., 2007). Genotypic diversity (H<sub>t</sub>) among

isolates was estimated from allelic frequencies using the equation  $H = 1 - \Sigma xi2$ , where xi is the frequency of the i<sup>th</sup> allele of a particular locus (Nei, 1973). The coefficient of population subdivision (G<sub>st</sub>) was computed as  $(H_t - H_s)/H_t$ , where,  $H_t$  is the total genetic diversity and H<sub>s</sub> is the average gene diversity over all subgroups (Nei, 1973). The allele frequencies at polymorphic loci, the N<sub>m</sub> values (effective migration rate), and the genetic identity among populations for characterizing genetic variation, observed number of alleles  $(N_a)$ , effective number of alleles  $(N_e)$ , Nei's gene diversity (He) and Shannon's information index (I) were calculated for each locus and population. Mean values of gene diversity in total populations (H<sub>t</sub>), gene diversity between populations (H<sub>s</sub>), proportion of gene diversity attributable to differentiation among populations (G<sub>st</sub>) and estimate of gene flow (N<sub>m</sub>) from G<sub>st</sub> were estimated across loci (McDonald and McDermot, 1993).

Table 2 SSR primers of Fusarium verticillioides used in this study (Ren et al., 2012).

Primer locus	Repeat of cloned allele	Primer Sequences	Expected Size (bp)	Tm (°C)
3H02	(ATG) <sub>11</sub>	F:ATCACCAAAACAGAGCAAAG R:GTGGCTGAGAAAGACAAGAA	181	52
4H18	(TTTC) <sub>6</sub>	F:TGATGCGGTCAAAGAATGG R:ACTGGAGCAGATGAAGAGC	152	56
5H08	(CTTT) <sub>6</sub>	F:ACCAACTAACATCCCGAATC R:CGTAAACTCAAACGCAAGG	410	58
5H12	(GAAA) <sub>7</sub>	F:GGCACCAACATTCCTGACG R:AACCGCCTACAAGCACCA	404	59
6H02	(GTTC) <sub>14</sub>	F:AACGGTATAGCGAGAAACG R:GGACGGAGTCGAATGAAGT	222	55
7H21	(GA) <sub>25</sub>	F:TTATGTCTGCCCAATCGC R:CCAGTTTCGCATCTGTCG	442	57
8H01	(AAG) <sub>9</sub>	F:TGGTATGATGGCTGGTTCG R:ACTAGAGGCTGGGTTGGTGT	485	55
9H05	(ATGG) <sub>11</sub>	F:AGGCACGAGTGGATAAGGC R:TTGGAAGCAAGCGAAGGA	221	56
9H09	(GT) <sub>18</sub>	F:ATCGGTGGTTTCTTGCTGC R:GCTCCCAACTGCCTACCTACA	263	59

### **Results and Discussion**

#### Isolation

Ninety-eight isolates of *Fusarium* in the section Liseola were isolated from 350 corn plants with symptom of crown and root rot disease. Among them 77 isolates were *F. verticillioides*. Previous studies also showed that *F. verticillioides* were predominant in corn fields.

#### Pathogenicity test

Results proved that all *F. verticillioides* isolates were pathogenic on corn. In the pathogenicity test, inoculated roots showed brown to black lesions and all isolates caused discoloration of the corn crowns and roots while no such symptoms were observed in control treatments.

# Distribution of alleles at polymorphic SSR loci

All nine microsatellite loci were polymorphic with a total of 43 alleles detected. Among which3H02, 4H18, 5H08, 5H12, 6H02, 7H21, 8H01, 9H05 and 9H09 showed a total of 5, 6, 5, 5, 4, 4, 5, 4 and 5 alleles per locus, respectively (Table 3, Fig. 2). Moreover, the average number of alleles per locus in populations was the highest (6 alleles) in Mehran and Holeylan populations and the lowest (2 alleles) in Dehloran and Mosiyan population (Table 4).

Table 3 Number of alleles at each locus across the five populations of *Fusarium verticillioides* from Ilam province.

SSR			Ν	Number of alleles			
primer	Dehloran	Mosiyan	Dasht abas	Darreh shahr	Mehran	Holeylan	Eivan
3H02	5	4	4	5	4	4	4
4H18	3	3	3	4	6	6	4
5H08	3	3	3	4	5	4	5
5H12	3	3	3	4	4	4	4
6H02	3	2	3	4	4	3	4
7H21	3	3	3	4	4	4	4
8H01	2	3	3	5	4	5	5
9H05	3	3	3	4	4	4	4
9H09	2	3	4	4	5	4	5



**Figure 2** Amplification profile of *Fusarium verticillioides* isolates using 6H02 SSR primer pairs. Numbers indicate the isolates (31 - 45), M indicates size marker (Ladder 1 kb).

SSR primer Number of alleles		Number of Polymorphic	Percentage of polymorphism	Polymorphic information
		alleles		Content (PIC)
3H02	5	5	100	0.228
4H18	6	6	100	0.281
5H08	5	5	100	0.288
5H12	5	5	100	0.294
6H02	4	4	100	0.325
7H21	4	4	100	0.335
8H01	5	5	100	0.289
9H05	4	4	100	0.313
9H09	5	5	100	0.277

**Table 4** Number of alleles, percentage of polymorphism and polymorphic information content obtained from SSR primers among *Fusarium verticillioides* populations.

### **Primers characteristics**

A summary of characteristics of nine microsatellite loci is given in Table 4. Number of alleles, percentage of polymorphism and Polymorphic Information Content (PIC) of SSR primers were evaluated. The PIC value varied from 0.33 (primer 7H21) to 0.228 (primers 3H02), which reflects the informative content of the primers. The total number of alleles, percentage of polymorphic alleles of all SSR markers were 43 and 100%, respectively. A total of 43 amplified alleles were polymorphic, and average number of polymorphic alleles per primer was 4.77. Percentages of polymorphic alleles were 100%. Total allelic number in populations varied from 27 (Dehloran and Mosiyan) to 40 (Mehran). Allele diversity of populations ranged from 0.195 in Mosiyan to 0.284 in Holeylan (Table 5).

#### Genetic variability of populations

Various population genetic parameters in nine microsatellite loci for all populations are given in Table 6. Observed allele number ( $N_a$ ) and effective numbers ( $N_e$ ) of alleles were higher in Mehran compared to other populations. The value of gene diversity ( $H_e$ ) were higher in Holeylan (H = 0.284) and Shannon's Information index (I) were also higher in Mehran (I = 0.436) but lower values were estimated for Mosiyan ( $H_e = 0.195$ ; I = 0.303).

Based on the SSR data, average genetic distance was calculated between seven

populations. The lowest genetic distance was found between Dehloran and Mosiyan populations (0.013), while the highest was revealed between Darreh shahr and Dehloran populations (0.139), (Table 7).

#### Genetic relationships among populations

Nei's genetic distances were estimated between populations. Cluster analysis based on UPGMA dendrogram to show the genetic relationships among populations (Fig. 3). The lowest genetic dissimilarity was found between Dehloran and Mosiyan and then Dasht abas. Moreover, the dendrogram showed the highest dissimilarity between Darreh shahr population and the six remaining populations. In this study, genetic diversity within and among seven populations of F. verticillioides sampled from Ilam was evaluated using SSR markers. The advantage of microsatellite markers over random amplified polymorphic DNA and polymerase chain reaction restriction fragment length polymorphism markers are their high polymorphism, specificity, high good reproducibility and unambiguous scorability (Tenzer et al., 1999; Sahran and Naef, 2008). These results are in accordance with results from previous studies on F. verticillioides, demonstrating that the population of this fungus is genotypically highly diverse (Danielsen et al., 1998; Chulze et al., 2000). The results showed that allele frequencies were different among nine SSRs loci. The allele frequencies showed that F. verticillioides may spread as

asexual stage in all populations. The sample sizes per population and SSRs primers were too small in this experiment for a statistically powerful test of gametic equilibrium. In this study the genetic similarity detected between isolates within the seven populations was probably due to exchange of corn seeds between sampled regions and geographical closeness as well. SSR results indicated that seven populations of *F. verticillioides* had low degrees of gene diversity, but Mehran maintained higher genetic diversity than the other regions (Table 6). Different research has been carried out on international populations of

F. verticillioides, and a different level of genetic diversity was detected within and between populations (Amoah et al., 1995). High genetic variability among  $F_{\cdot}$ verticillioides isolates from different hosts has been reported by RFLPs of ribosomal DNA and RAPDs analysis in Ghana (Amoah et al., 1995). Mitter et al. (2002) reported high genetic variation among F. moniliforme isolates from different geographical regions in India and different hosts by RAPD markers. In Iranian populations, over 91% of the gene diversity was distributed on a local level within populations (Fig. 4).

Table 5 Number of alleles and mean number of allelic diversity in *Fusarium verticillioides* populations.

Population	Number of alleles	Mean number of Allelic diversity
Dehloran	27	0.200
Mosiyan	27	0.195
Dasht abas	29	0.232
Darreh shahr	38	0.273
Mehran	40	0.282
Holeylan	38	0.284
Eivan	39	0.279

Table 6 Genetic diversity of Fusarium verticillioides populations based on microsatellite loci.

Population	Ν	R%	N <sub>a</sub>	Ne	H <sub>e</sub>	Ι	
Dehloran	12	62.79	1.256	1.330	0.200	0.305	
Mosiyan	10	62.79	1.256	1.310	0.195	0.303	
Dasht abas	9	67.44	1.369	1.387	0.232	0.351	
Dareh shahr	12	88.37	1.767	1.442	0.273	0.421	
Mehran	12	93.02	1.860	1.463	0.282	0.436	
Holeylan	11	88.37	1.767	1.463	0.284	0.435	
Eivan	11	90.70	1.814	1.452	0.279	0.432	

Na = Observed number of alleles

Ne = Effective number of alleles

He = Nei's (1973) gene diversity I = Shannon's information index

R= the percentage of polymorphic loci

Table 7 Genetic distance between	Fusarium	verticillioides	populations.
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Population	Dehloran	Mosiyan	Dasht - abas	Darreh - shahr	Mehran	Holeylan	Eivan
Dehloran	0						
Mosiyan	0.013	0					
Dasht abas	0.020	0.018	0				
Darreh - shahr	0.139	0.136	0.122	0			
Mehran	0.044	0.046	0.037	0.071	0		
Holeylan	0.097	0.091	0.060	0.066	0.048	0	
Eivan	0.109	0.104	0.079	0.052	0.056	0.042	0



Figure 3 Dendrogram constructed with UPGMA based on Jaccard's similarity coefficient among seven populations of *Fusarium verticillioides* originated from Ilam province.



Figure 4 Analysis of molecular variance (AMOVA) in Fusarium verticillioides populations from Ilam province.

However, there was a high degree of genetic similarity among populations separated by short geographical distances like Dehloran and Mosiyan. The low level of gene diversity (G<sub>st</sub>) was detected among all seven populations. Low G<sub>st</sub> value (0.147) indicated low genetic differentiation among the seven populations and showed evidence for geographical subdivision among populations (Bayraktar, 2010). The genetic distances were very small and the geographic distances between populations ranged from 20 to 375 km, this indicates that populations linked by movement of conidia within populations. Gene flow may have occurred among these populations with infected plant debris, infected seeds, agricultural vehicles and fungal spores. Gene flow has a significant role on the genetic diversity of populations. In the absence of gene flow, genetic drifts cause developing different allele frequencies at neutral loci, leading to differentiation in

isolate populations (Keller et al., 1997). The high genetic similarity among populations of F. verticillioides suggests that gene flow has occurred across long distances. N<sub>m</sub> averaged 2.890 in all loci and populations, suggesting a high level of gene flow needed to prevent populations from diverging by genetic drift (Keller et al., 1997). Another factor that has an important role in the genetic diversity of F. verticillioides populations is the mating system. With this assumption if the sexual reproduction of F. verticillioides occurs in nature, the sexual spores (ascospores) may play a major role in population biology. Moreover, infected seed can lead to persistence of genotypes; we consider that infected seed can explain the distribution and diversity of genotypes found at the end of the growing season in natural populations. If the main source of primary inoculums was asexual spores from seeds, it is expected to find some clones that were distributed among different locations in a field. Genetic drift and limit the number selection would of genotypes present in field populations, unless sexual reproduction generated new genotypes and wind dispersal distributed ascospores among populations every year (Shah et al., 1995). Depending on environmental factors, diversity of *F*. genetic verticillioides populations may possess potentially a risk of incidence of severe disease and high loss in corn farms in these regions. These data can help breeders in screening for resistant cultivars based on genetic diversity of F. verticillioides populations for local disease management. Understanding population genetic structure of F. verticillioides in the present study may provide insights into the epidemiology and evolutionary potential of the pathogen and could lead to improved disease. strategies for managing the Moreover, neutral DNA markers such as RAPD and AFLP (McDonald and McDermott, 1993) would generate further results on the genetic structure of F. verticillioides populations in corn.

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ساختار ژنتیکی جمعیتهای Fusarium verticillioides عامل پوسیدگی ریشه و طوقه ذرت در استان ایلام با استفاده از نشانگرهای ریزماهواره

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**چکیدہ**: بررسی تنوع ژنتیکی جمعیتھای قارچ *Fusarium verticillioides* سطوح مختلفی از اطلاعات در مورد مدیریت بیماری پوسیدگی ریشه و طوقه در مزارع ذرت را فراهم می کند. از نـشانگرهای SSR برای تعیین ساختار ژنتیکی و تخمین تنوع ژنتیکی در ۷۷ جدایه F. verticillioides از مناطق کشت عمده ذرت در استان ایلام از جمله: دهلران، موسیان، دشت عباس، مهران، ایوان، هلیلان و درهشهر در سال زراعی ۱۳۹۵–۱۳۹۴ استفاده شد. با به کار گیری نه جفت پرایمر SSR، میانگین تعداد آلـل.هـا در جمعیتها ۳۴ آلل بود که فراوانی آللها در جمعیت دهلران و موسیان با ۲۷ آلل، کمترین و در جمعیت درهشهر با ۴۰ آلل، بیشترین میزان فراوانی را داشتند. تعداد آللهای مشاهده شده (Na) و تعداد آلل-های مؤثر (Ne) در مهران (Na = ۱/۸۶۰, Ne = ۱/۴۶۳) نسبت به دیگر جمعیتها بیشتر بود. بیشترین تنوع ژنتیکی (H<sub>e</sub>) در هلیلان (H<sub>e</sub> = ۰/۲۸۴) و بیشترین شاخص شانون (I) در مهران (I = ۰/۴۳۶) بود اما کم ترین میزان برای موسیان (He = ۰/۱۹۵, I = ۰/۳۰۳) برآورد شد. کم ترین فاصله ژنتیکی بین دهلران و موسیان (۱/۰۱۳) بود درحالی که بیش ترین فاصله ژنتیکی بین دهلران و درهشهر (۰/۱۳۹) وجود داشت. تنوع ژنتیکی کل (H<sub>t</sub>) و تنوع ژنی بین زیر جمعیتها (H<sub>s</sub>) بهترتیب ۰/۲۹۲ و ۲۴۹/۰ تخمین زده شد. میزان تنوع ژنتیکی مؤثر در تمایز بین جمعیتها (Gst) ۰/۱۴۷ و میزان جریان ژنی UPGMA بود. تجزیه کلاستر براساس UPGMA نشان داد که کمترین فاصله ژنتیکی بین دهلران، (ارس) ۲/۸۹۰ موسیان و سپس دشتعباس وجود دارد. این دندروگرام بیشترین فاصله ژنتیکی را بین درهشهر و شش جمعیت دیگر نشان میدهد. نتایج حاصل از این مطالعه در راستای تولید ارقام مقاوم به بیماری پوسیدگی ریشه و طوقه و همچنین توسعه روشهای کنترل بیماری میتواند مورد استفاده قرار گیرد.

واژگان كليدى: ذرت، فاصله ژنتيكى، پوسيدگى ريشه، SSR