Characterization of Iranian nonaflatoxigenic strains of *Aspergillus flavus* based on microsatellite-primed PCR

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ABSTRACT

Out of fifty-two Iranian nonaflatoxigenic strains of *Aspergillus flavus*, collected from various substrates (soil and kernel) and sources (peanut, corn and pistachio), fifteen representatives were selected according to their different geographical origins (six provinces: Guilan and Golestan, Ardebil, Fars, Kerman and Semnan) and vegetative compatibility groups (VCGs, IR1 to IR15) for microsatellite-primed PCR analysis. Two inter-simple sequence repeat (ISSR) primers AFMPP and AFM13 were used to determine polymorphism and the relationship among strain isolates. *A. flavus* isolates were identified by their morphologies and their identities were confirmed by PCR amplification using the specific primer pair ITS1 and ITS4. The results revealed variations in the percentages of polymorphisms. In the ISSR analysis, primers AFMPP and AFM13 generated a total of 18 and 23 amplicons among the fungal strains, out of which 12 (66.7%) and 22 (95.7%) were polymorphic, respectively. Cluster analysis of the ISSR data was carried out using 1D DNA gel image analysis. The two dendrograms obtained through these markers showed six different clusterings of testing nonaflatoxigenic *A. flavus* strains, but we noticed that some clusters were different in some cases. The microsatellite-primed PCR data revealed that the Iranian nonaflatoxigenic isolates of *A. flavus* were not clustered according to their origins and sources. This study is the first to characterize Iranian nonaflatoxigenic isolates of *A. flavus* using ISSR markers.

Keywords: Aflatoxin; Molecular marker; Inter-simple sequence repeat; Polymorphism
INTRODUCTION

Aspergillus flavus Link. ex Fries, a haploid organism found worldwide in a variety of crops, including maize, cottonseed, almond, pistachio and peanut, causes substantial and recurrent worldwide economic liabilities [1, 2]. This filamentous fungus produces aflatoxins (AFLs) B1 and B2, which are among the most carcinogenic, acutely hepatotoxic and immunosuppressive compounds found in nature [3-5]. Recent efforts to reduce AFL contamination in crops have focused on the use of nonaflatoxigenic A. flavus isolates as biological control agents.

Taxonomically, A. flavus belongs to the Aspergillus genus of the section Flavi [6, 7]. Molecular biology has offered several insights into the detection and genetic relationships of fungal isolates from their DNA sequences, taxonomy, population structure and the epidemiology associated with them [8]. Various molecular methods have been used for the detection of Aspergillus from environmental and clinical samples [9-11]. Targets for the genus level detection of Aspergillus have included the 18S rRNA gene, mitochondrial DNA, the intergenic spacer region, and the internal transcribed spacer (ITS) regions. Ribosomal RNA (rRNA) genes in ribosomal DNA possess characteristics that are suitable for the detection of pathogens at the species level [12]. These rDNA sequences are highly stable and exhibit a mosaic of conserved and diverse regions within the genome [13]. They also occur in multiple copies with up to 200 copies per haploid genome arranged in tandem repeats [8], each consisting of the 18S small subunit (SSU), the 5.8S, and the 28S large subunit (LSU) genes. ITS primers 1 and 4 have been used to amplify the entire 5.8S rRNA gene, both ITS regions I and II, and a portion of the 18S small-subunit rRNA gene.

In recent years, there has been vast progress in the development of molecular biology tools and technologies [3,13-16]. Inter-simple sequence repeat (ISSR) is based on the amplification of regions (100-3000 bp) between inversely oriented, closely spaced microsatellites by primers (25-30 bp) consisting of several simple sequence repeats [17]. These primers anneal to simple-sequence repeats (microsatellites) that are abundant throughout the eukaryotic genome and evolve rapidly [16,18-19]. Prior knowledge of the DNA sequence of the genome to be analyzed is not required for primer design [20]. However, since there is a lot of diversity among fungi, primers that work for one may not work for another. Hence, ISSR primers need to be optimized for each species [21].

Microsatellite loci with di- to hexanucleotide repeats and 1000-bp flanking sequences were identified from the genome sequence of A. flavus NRRL3357 (http://www.aspergillusflavus.org/) using Tandem Repeats Finder version 4.00 [22]. Molecular typing of A. flavus using microsatellites yields multiple advantages such as high discriminatory power, high reproducibility and easy exchange of the results [14]. It is reported that the ISSR sequences as molecular markers that can lead to the detection of polymorphism, which is a new approach to study SSR distribution and frequency [23].

The specific aims of this work were to: 1. examine genetic relatedness among nonaflatoxigenic isolates of A. flavus belonging to three Iranian pistachio, peanut and maize populations, and 2. assess polymorphisms among nonaflatoxigenic isolates of A. flavus using PCR amplification of ISSR molecular markers.
MATERIALS AND METHODS

**Fungal strain**: Out of fifty-two nonaflatoxigenic isolates of *Aspergillus flavus* from three populations of *A. flavus* (peanut, maize and pistachio) isolated from different geographical origins (Guilan, Golestan, Ardebil, Fars, Kerman and Semnan provinces) and substrates (soil and kernel) (data not shown), 3, 7 and 5 representatives were randomly selected according to their vegetative compatibility groups (VCGs, IR1 to IR15) for microsatellite-primed PCR (MP-PCR) analysis (Table 1). Strain isolates were stored as spore suspensions in 20% glycerol at -20°C. All strains had already been characterized for their aflatoxicogenic ability after the mycelium collection yeast extract sucrose broths were analyzed by HPLC, to confirm AF production. This test is important because AF production is extremely dependent on growth conditions; it was, therefore, important to determine aflatoxicogenic ability under current test conditions. Using specific primer pairs ITS1 and ITS4 as described previously [24], all *A. flavus* strains were identified and confirmed based on amplifications of internal transcribed spacers (ITS) of ribosomal DNA (rDNA) by polymerase chain reaction (PCR) combined with sequencing of the amplicons [25,26].

**DNA extraction**: Total DNA was extracted from the mycelia of fungal isolates obtained from 7-day-old cultures grown in YES liquid media according to Prabha *et al.* (2012) with minor modifications [27]. Briefly, mycelia were collected by vacuum filtration, ground into a fine powder in liquid N2 and stored at -20°C. The frozen powder was then suspended in a 500 μl TES buffer (200 mM Tris-HCl, pH 7.5; 25 mM EDTA and 250 mM NaCl and 0.5% SDS), vortexed for 5 sec and incubated at 65 °C for 10 min. The reaction mixture was centrifuged at 13,000 rpm for 1 min and DNA was extracted with phenol/chloroform (1:1). DNA was then precipitated in 300 μl of cold iso-propanol and incubated for 30 min at -20°C and recovered by centrifugation at 13,000 rpm for 5 min. Afterwards, the pellet was washed with 70% cold ethanol and dried for 15 min at 37°C. Finally, the isolated DNA was resuspended in 50 μl of sterile distilled water and stored at -20°C. DNA concentration was measured spectrophotometrically with a NanoDrop Spectrophotomer ND-1000. DNA quality was also examined by running on 1.2% gel agarose for 75 min at 80 V, after which the gel was exposed to UV light. The presence of a highly resolved high molecular weight band and absence of smear confirmed the good quality of DNA.

**Molecular identification of *Aspergillus flavus***: Identification of *A. flavus* using an Internal transcribed spacer (ITS) was conducted. Primer pairs (ITS1 and ITS4, Table 2) were derived from the ITS1-5.8S-ITS4 region [24]. PCR amplification was carried out in a 25 μl reaction mixture (Table 3) in a Biometra Thermal Cycler (T1 thermocycler; Biometra, Göttingen, Germany). The PCR product was analyzed by electrophoresis in 1.2% agarose gel stained with DNA green viewer dye (greenGel stain,10 mg/ml) and visualized with the UVsolo TS gel documentation system (Biometra).

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Table 1: Monospore isolates used to evaluate polymorphisms in non-aflatoxigenic isolates of *Aspergillus flavus*, substrate, vegetative compatibility group (VCG) and geographical origin

| Strain isolate | Substrate                  | VCG | Geographical origin                  |
|----------------|---------------------------|-----|--------------------------------------|
| IRP-049        | Soil/pistachio orchard    | IR1 | Rafsangan/Kerman province            |
| IRP-107        | Soil/pistachio orchard    | IR2 | Rafsangan/Kerman province            |
| IRP-082        | Soil/pistachio orchard    | IR3 | Damghan/Semnan province              |
| IRP-144        | Soil/pistachio orchard    | IR4 | Damghan/Semnan province              |
| IRG-075        | Soil/Peanut field         | IR5 | Minoodasht,Golestan province         |
| IRG-129        | Soil/ Peanut field         | IR6 | Astane-e Ashrafieh/Guilan province   |
| IRM-074        | Soil/ Maize field          | IR7 | Darab/Fars province                  |
| IRM-193        | Soil/ Maize field          | IR8 | Fasa/Fars province                   |
| IRM-014        | Soil/ Maize field          | IR9 | Pars Abad/Ardebil province           |
| IRM-211        | Soil/ Maize field          | IR10| Pars Abad/Ardebil province           |
| IRP-179        | Kernel/ pistachio orchard  | IR11| Rafsangan/Kerman province            |
| IRG-517        | Kernel/ Peanut field       | IR12| Astane-e Ashrafieh/Guilan province   |
| IRM-031        | Kernel/ Maize field        | IR13| Pars Abad/Ardebil province           |
| IRM-041        | Kernel/ Maize field        | IR14| Darab/Fars province                  |
| IRM-081        | Kernel/ Maize field        | IR15| Darab/Fars province                  |

Table 2: Primer, target gene, sequence and expected PCR product size

| Primers | Region | Primer sequences (5’→3’) | Annealing temp. (°C) | PCR product Size (bp) |
|---------|--------|--------------------------|----------------------|-----------------------|
| ITS1    | ITS    | TCCGTAGGTTGAACTGCGG       | 58                   | 600                   |
| ITS4    | ITS    | TCCCTCGCTTATGATGC         |                      |                       |

Table 3: PCR reaction mixture for amplification of the ITS region

| Final concentrations (volume) | Reaction mixture |
|--------------------------------|------------------|
| 1X (2.5µl)                    | PCR buffer       |
| 50 mM (1 µl)                  | MgCl₂            |
| 10 pmol/µl (1.5 µl)           | Primer           |
| 2.5 mM/µl (2 µl)              | dNTPs            |
| 5 U/µl                        | Taq DNA polymerase|
| 20 ng/µl (2 µl)               | Template DNA     |
| 14.3 µl                       | D.D.W            |
| 25 µl                         | Total            |

Microsatellite-primed PCR and electrophoresis: Two ISSR primers that included AFMPF and AFM13 and showed more polymorphisms in previous studies were used [11, 22, 28,29] (Table 4). The genomic DNA sample was amplified using ISSR primers in a 25 µl reaction mixture containing PCR Buffer 1X, 0.2 µM ISSR primers, 3 mM MgCl₂, 1 unit Taq DNA polymerase and 50 ng of the template DNA sample. The PCR was carried out in a Biometra Thermal Cycler (T1 thermocycler; Biometra, Göttingen, Germany) with the following profile: initial heating at 93°C for 5 min, thirty cycles of denaturation at 93°C for 30 s, annealing at 45°C for 1 min, extension at 72°C for 1.5 min and a final extension period at 72°C for 5 min. The result of each amplification reaction was analyzed on 2% agarose gel in a TBE buffer 1X (pH 8) and run at 80 V. Amplified fragments were then visualized using an ultraviolet transilluminator (UVsolo TS gel documentation system, Biometra) and compared with a 100 bp DNA size marker (Fermentas).

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Table 4: List of ISSR primer sequences and their annealing temperatures ($T_a$)

| Primers  | Repeat motifs | Primer sequences (5' → 3') | $T_a$ (°C) |
|----------|---------------|----------------------------|------------|
| AFMPP    | (GACA)$_4$    | GACAGACAGACAGACA           | 30.7       |
| AFM13    | (GTG)$_5$     | GAGGCTTGGGCGTTCT            | 47.4       |

**Data analysis:** The internal transcribed spacer (ITS) region, ITS 1–5.8S–ITS 2, from nonaflatoxigenic isolates of *A. flavus* were amplified, sequenced, and compared with the reference strain sequence in GenBank. Gel images from ISSR-PCR fingerprint patterns of genomic DNAs were analyzed using 1D DNA gel image analysis software (TotalLab v2, Nonlinear Dynamics, Newcastle upon Tyne, UK) and dendrograms were constructed. The allele size was calculated using Alpha Imager software [30]. The ladder in which all alleles were absent was used as an outgroup for dendrogram rooting.

**RESULTS AND DISCUSSION**

ITS amplicons from *A. flavus* strains were 600 bp in size. Comparison of the reference strain and the GenBank sequence demonstrated that both ITS 1 and ITS 2 regions were needed for the accurate identification of *A. flavus*. ISSR profiles and allele sizes at ISSR markers AFM13 and AFMPP resulting from the analyses of Iranian nonaflatoxigenic strains of *A. flavus* are shown in Figures 1 and 2 and Tables 5 and 6, respectively.

**Figure 1:** ISSR-PCR fingerprint pattern of genomic DNAs isolation from different nonaflatoxigenic isolates of *Aspergillus flavus* generated using primer ISSR M13. Lanes 1-15 were IRP49, IRP107, IRP179, IRP82, IRM41, IRM193, IRM74, IRP144, IRM81, IRM31, IRM14, IRM211, IRG129, IRG517, IRG75, respectively. M: molecular-weight marker (100 bp DNA ladder).
Figure 2: ISSR-PCR fingerprint pattern of genomic DNAs isolation from different nonaflatoxicogenic isolates of *Aspergillus flavus* generated using primer ISSR MPP. Lanes 1-15 were IRP49, IRP107, IRP179, IRP82, IRM41, IRM193, IRM74, IRP144, IRM81, IRM31, IRM14, IRM211, IRG129, IRG157, IRG75, respectively. M: molecular-weight marker (100 bp DNA ladder).

Table 5: Allele size of *Aspergillus flavus* at microsatellite marker AFM13 resulting from the analysis of nonaflatoxicogenic isolates of *Aspergillus flavus* from Iran

| Lanes | Fragment size (bp) |
|-------|-------------------|
| 1     | 953               |
| 2     | 953               |
| 3     | 953               |
| 4     | 953               |
| 5     | 953               |
| 6     | 953               |
| 7     | 953               |
| 8     | 953               |
| 9     | 953               |
| 10    | 953               |
| 11    | 953               |
| 12    | 953               |
| 13    | 953               |
| 14    | 953               |
| 15    | 953               |

Note: Lanes 1-15 were IRP49, IRP107, IRP179, IRP82, IRM41, IRM193, IRM74, IRP144, IRM81, IRM31, IRM14, IRM211, IRG129, IRG157, IRG75, respectively.
ISSR marker polymorphism: The characteristics of ISSR marker polymorphisms are shown in Table 7. The percentages of polymorphic fragments for AFM13 and AFMPP were 95.7% and 66.7%, respectively. At the population level, the percentage of polymorphic bands from AFMPP ranged from 0% to 44.4%, and the average value was 25.9%.

**Table 6:** Allele size of *Aspergillus flavus* at microsatellite marker AFMPP resulting from the analysis of nonaflatoxigenic isolates of *Aspergillus flavus* from Iran

| Lanes | 1 bp | 2 bp | 3 bp | 4 bp | 5 bp | 6 bp | 7 bp | 8 bp | 9 bp | 10 bp | 11 bp | 12 bp | 13 bp | 14 bp | 15 bp |
|-------|------|------|------|------|------|------|------|------|------|-------|------|-------|------|-------|------|
| Fragment size (bp) | - | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 1400 | 1440 |
| 312 | 338 | 410 | 475 | 500 | 563 | 602 | 644 | 667 | 680 | 721 | 778 | 810 | 843 | 880 | 900 |
| 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 900 | 1400 | 1440 |
| 800 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | 880 | - | 1440 |
| 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | 843 | - | - |
| 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | 810 | - | - |
| 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | 778 | - | - |
| 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | 721 | - | - |
| 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | 710 | - | - |
| 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | 667 | - | - |
| 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | 644 | - | - |
| 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | 602 | - | - |
| 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | 563 | - | - |
| - | 520 | 520 | 520 | 520 | 520 | 520 | 520 | 520 | 520 | 520 | 520 | 520 | - | - |
| - | - | - | - | - | - | - | - | - | - | - | - | - | 500 | - |
| 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | 475 | - | - |
| 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | 410 | - | - |
| 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | 338 | - | 338 |
| 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | 312 | - | - |

**Note:** Lanes 1-15 were IRP49, IRP107, IRP179, IRP82, IRM41, IRM193, IRM74, IRP144, IRM81, IRM31, IRM14, IRM211, IRG129, IRG517, IRG75, respectively.

**Table 7:** Characteristics of ISSR marker polymorphisms

| Primer sequence | No. of fragments | No. of polymorphic fragments | % polymorphic fragments | Product size range (bp) |
|-----------------|-----------------|------------------------------|-------------------------|------------------------|
| (GACA)$_4$      | 18              | 12                           | 66.7                    | 312-920                |
| (GTG)$_5$       | 23              | 22                           | 95.7                    | 475-953                |

**Phylogenetic analysis:** Polymorphic fragments were used for the statistical interpretation of phylogenetic relations (Total Lab 120, UVP soft). Figures 3 and 4 show the dendrograms constructed using cluster analysis. Genetic similarities (x-axis) are expressed as 0–1. AFM13 and AFMPP initially split all fifteen nonaflatoxigenic isolates of *A. flavus* into two main groups (I and II) at 3.8% and 3.1% genetic similarities, respectively (Figures 3 and 4).

The larger primer groups AFM13 and AFMPP comprised two and three subgroups, respectively, each splitting further into six smaller groups (I-VI) containing one to four *A. flavus* strain isolate(s) (Figures 3 and 4). Otherwise, using ISSR primers AFM13 and AFMPP, the fifteen strain isolates belonging to three populations of Iranian nonaflatoxicigenic isolates of *A. flavus* from pistachio, maize and peanut were separated and placed into six distinct clusters based on genetic similarities (Figures 3 and 4).
Similarity among isolates

**Figure 3:** Dendrogram showing clustering of Iranian nonaflatoxigenic isolates of *A. flavus* based on genetic similarities in PCR reactions using ISSR primer AFM13. The two main groups formed are shown under I and II. Strain isolates with highest genetic similarities (more than 80%), are indicated by the letters A, B, C and D. (Ladder=Outgroup).

Similarity among isolates

**Figure 4:** Dendrogram showing clustering of Iranian nonaflatoxigenic isolates of *A. flavus* based on genetic similarities in PCR reactions using ISSR primer AFMPP. The two main groups formed are
shown under I and II. Strain isolates with highest genetic similarities (more than 80%), are indicated by the letters A, B, C and D. (Ladder=Outgroup)

In the present study, a method to identify *A. flavus* strain isolates was developed using the 18S and 28S rRNA genes for primer binding sites. rDNA has been utilized by many investigators for species determination in a wide variety of yeasts and fungi [31-33]. In this research, 15 nonaflatoxigenic strains of *A. flavus* belonging to different substrates, geographical regions and VCGs were analyzed to determine the degree of polymorphism. The ISSR marker was identified by PCR amplification of DNA using primer pairs composed of microsatellite sequences that may be anchored at the 3' or 5' end of 2 to 4 arbitrary and often degenerate nucleotides [34,35]. The results indicated that to amplify ISSR sequences in DNA extracted from nonaflatoxigenic isolates of *A. flavus*, ISSR primers AFMPP and AFM13 produced positive results from the PCR trials. ISSR, which is a dominant marker, has greater robustness in repeatability and high variability [36]. The two ISSR primers (AFMPP and AFM13) produced a series of discrete bands of different intensities at annealing temperatures 30.7°C and 47.4°C. Several isolates had similar banding patterns such as those in lanes 1 and 4 (strain isolates IRP49 and IRP82) and lanes 5 to 13 (strain isolates IRM41, IRM193, IRM74, IRP144, IRM81, IRM31, IRM14, IRM211, IRG129) in the AFMPP profile.

Altogether, the ISSR primers AFM13 and AFMPP generated 232 and 135 polymorphic bands ranging from 337 bp to 953 bp and 312 bp to 920 bp across fifteen strain isolates, respectively. Of the 23 and 18 ISSR discernible bands from primers AFM13 and AFMPP, 22 and 12 were polymorphic, respectively.

The Iranian nonaflatoxigenic isolates exhibited a high level of polymorphism, which was reflected in the number and percentage of polymorphic loci. Because of its simple technology and high level of polymorphism, microsatellite-primed PCR has been widely used for population genetic studies [22,37-39]. They produce different numbers of DNA fragments, depending on their simple sequence repeat motifs. In the current study, it was found that the ISSR AFM13 (GTG)_5 tested was more polymorphic among our nonaflatoxigenic isolates. ISSR analysis aims at studying the polymorphism of highly repetitive genome regions [39,40]. The percentage of polymorphic bands from AFM13 ranged from 23.1% to 69.9%, and the average value was 45.5%. Usually, ISSR primers based on di- and tri-nucleotide repeats reveal high polymorphisms [41,42] which was also found to be true for the present study. Hatti *et al.* (2010) reported an average of 9.33 polymorphic bands per ISSR primer [43]. In contrast, Batista *et al.* (2008) showed high genetic variability among strains of *A. flavus* and other species of the *A. flavus* group by using the ISSR marker [28]. They showed that the (GACA)_4 primer yielded a higher polymorphism as compared to (GTG)_5.

In our study, some polymorphic bands appeared more than once across the different strain isolates. Primers based on a repeat sequence and the resulting PCR reaction amplify the sequence between two ISSRs, yielding a multilocus marker system [43]. The dendrogram analysis for AFM13 showed that cluster II was comprised of IRM74, IRM193, IRM144 and IRM81 isolates, while cluster III contained IRM41, IRM31, IRP179 and IRM14 isolates. Clusters IV and V possessed IRP107, IRM211 and IRP49, IRG129, IRG75 isolates, respectively. Strain isolates IRP82 and IRG517 grouped into
clusters I and VI, respectively, showed their separate identities in comparison with other isolates. Although ISSRs are mostly random-type markers, they are thought to be highly useful for genetic diversity and phylogenetic studies [18].

For the ISSR primer AFMPP, cluster II comprised of IRP49, IRP107 and IRP179, while cluster III contained IRM193 and IRP144 isolates. Clusters IV and V possessed IRM41, IRM74, IRM81 and IRP82, IRM14, IRM31, IRM211 isolates, respectively. Strain isolates IRG517 and IRG75, IRG129 grouped into clusters I and VI, respectively, showed their separate identities in comparison with other isolates. Therefore, it can be concluded that ISSR markers could be used to study population structure among A. flavus and related species [12,28,29,44].

The similarity for maize (IRM193, IRM144, IRP41, IRM31, IRM14 and IRP179) and peanut strain isolates (IRG517 and IRG129) reached over 80% for the ISSR primer AFM13. Likewise, the similarity for maize (IRM193, IRP144, IRM41, IRM74 and IRM14) and peanut strain isolates (IRP179, IRG517 and IRG129) reached over 80% for the ISSR primer AFMPP. According to previous studies, ISSR markers have been used to determine similarity and dissimilarity between aflatoxigenic and nonaflatoxigenic isolates of A. flavus [43].

Similar to Yin et al. (2009) who showed that the toxigenic and atoxigenic isolates of A. flavus, collected from peanut fields, were not clustered based on their regions, ability of aflatoxin and sclerotial production [33], in the present study, the analysis of microsatellite-primed PCR data showed that Iranian nonaflatoxigenic isolates of the A. flavus were also not clustered based on their geographical origins and substrates. To the researchers’ knowledge, this is the first study of population analysis of nonaflatoxigenic isolates of A. flavus based on microsatellite-primed PCR in Iran.

Biological variability and the management of genetic variation within a species is a commonly recognized value in natural resources administration. Two primers, AFM13 and AFMPP gave reproducible banding profiles for most Iranian nonaflatoxigenic isolates of A. flavus tested. In this study, the ISSRs exposed significant numbers of polymorphisms, providing indication of A. flavus variability. Each of the two ISSR primers revealed a relatively high intra-species variability among the A. flavus isolates with considerable variation in morphological features. ISSR has an advantage over randomly amplified polymorphic DNA (RAPD) because its primers are longer, allowing for higher annealing temperatures that apparently provide a higher reproducibility of fragments than RAPD. Cluster analysis of the ISSR data divided the isolates of A. flavus to groups. The different subgroups formed by each primer were indicative of intra-species variability. The ecological niche may be used to explain how the several groups of A. flavus strain isolates were formed by the ISSR primers.

The varying similarity ranges within strain isolates of the A. flavus could also be a result of isolates that share a host range and/or ecological niche. Population genetics data can provide valuable information, often unattainable via other approaches, for monitoring species of management, conservation and ecological interest. Our experiments have demonstrated that ISSR analysis is a powerful tool for the identification of polymorphisms in Iranian nonaflatoxigenic isolates of A. flavus. Whilst
this technique gives useful information, several other ISSR primers are needed for more reliable results.

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Conflict of Interest: The authors declare that they have no competing interest.

REFERENCES
1. Frisvad JC, Thrane U, Samson R. In: Dijksterhuis J, Samson RA. (Ed.), Food Mycology: A Multifaceted Approach to Fungi and Food, CRC Press, Taylor and Francis Group, Boca Raton, 2007; pp.135–140.
2. Robens J, Cardwell FK. The cost of mycotoxin managment to the USA: Management of aflatoxin in the United States. J Toxicol 2003;22:139-152.
3. IARC (International Agency for Research on Cancer). World Health Organization, International Agency for Research on Cancer, Lyon, France, 1993; 56:571.
4. Manonmani HK, Anand S, Chandrashekar A, Rati ER. Process Biochem 2005; 40:2859-2864.
5. Oktay HI, Heperkan D, Yelboga E, Karaguler NG. Aspergillus flavus – primary causative agent of aflatoxins in dried figs. Mycotaxon 2011;115:425-433.
6. Egel DS, Cotty PJ, Elias KS. Relationships among isolates of Aspergillus sect. Flavi that vary in aflatoxin production. Phytopathol 1994;84:906-912.
7. Gams W, Christensen, M, Onions AH, Pitt, JI, Samson RA. Infragenetic taxa of Aspergillus. In Advances in Penicillium and Aspergillus Systematics, ed. by Samson RA, Pitt JI, Plenum Press, New York, 1985; pp. 55-62.
8. Paplomatas EG. Molecular diagnostics for soil-borne fungal pathogens. Phytopathol Mediterr 2004;43:213-220.
9. Bretagne S, Costa JM, Marmorat-Khuong A, Poron F, Cordonnier C, Vidaud M, Fleury-Feith J. Detection of Aspergillus species DNA in bronchoalveolar lavage samples by competitive PCR. J Clin Microbiol 1995;33:1164-1168.
10. Einsele H, Hebart H, Roller G, Loffler J, Rothenhofer I, Muller CA, Bowden RA, van Burik J, Engelhard D, Kanz L, Schumacher U. Detection and identification of fungal pathogens in blood by using molecular probes. J Clin Microbiol 1997;35:1353-1360.
11. Yamakami Y, Hashimoto A, Tokimatsu I, Nasu M. PCR detection of DNA specific for Aspergillus species in serum of patients with invasive Aspergillosis. J Clin Microbiol 1996;34:2464-2468.
12. O’Donnell K. Ribosomal DNA internal transcribed spacers are highly divergent in the phytopathogenic ascomycete Fusarium sambucinum (Gibberella pulicaris). Curr Genet 1992;22:213-220.
13. Bruns TD, White TJ, Talqor JW. Fungal molecular systematics. Annu Rev Ecol Sys 1991;22:525-564.

http://mbrc.shirazu.ac.ir
14. Beckmann JS. Oligonucleotide polymorphisms: A new tool for genomic genetics. Biotechnology 1988;6:161-164.
15. Frankham R, Ballou JD, Briscoe DA. Introduction to Conservation Genetics. Cambridge University Press, Cambridge, UK. 2002.
16. Louis M, Louis L, Simor AE. The role of DNA amplification technology in the diagnosis of infectious diseases. Can Medical Assoc J 2000;163:301-309.
17. Reischl U, Lohmann CP. [Polymerase chain reaction (PCR) and its possible applications in diagnosis of infection in ophthalmology.] Klin Monatsbl Augenheilkd 1997;211:227-234. [in German]
18. Morgante M, Hanafery M, Powell W. Microsatellites are preferentially associated with nonrepetitive DNA in plant genomes. Nat Genet 2002;30:194-200.
19. Kurtzman CP, Smiley MJ, Robnett CJ, Wicklow DT. DNA relatedness among wild and domesticated species in the Aspergillus flavus group. Mycologia 1986;78:955-959.
20. Tautz D, Renz M. Simple sequences are ubiquitous repetitive components of eukaryotic genomes. Nucleic Acids Res 1984;12:4127-4138.
21. Klaassen CH. MLST versus microsatellites for typing Aspergillus fumigates isolates. Med Mycol 2009;47:27-33.
22. Balloux F, Lugon-Moulin N. The estimation of population differentiation with microsatellite markers. Mol Ecol 2002;11:155-165.
23. Bornet B, Muller C, Paulus F, Branchard M. Highly informative nature of intersimple sequence repeat (ISSR) sequence amplified using tri- and tetra-nucleotide primers from DNA of cauliflower (Brassica oleracea var. botrytis L.) Genome 2002;45:890-896.
24. White TJ, Bruns T, Lee S, Taylor J. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In: PCR Protocols: a guide to methods and applications. (Innis MA, Gelfand DH, Sninsky JJ, White TJ, eds). Academic Press, New York, USA: 1990; pp. 315-322.
25. Chen Z-Y, Brown RL, Damann KE, Cleveland TE. Identification of unique elevated levels of kernel proteins in aflatoxin-resistant maize genotypes through proteome analysis. Phytopathology 2002;92:1084-1094.
26. Criseo G, Bagnara A, Bisignano G. Differentiation of aflatoxin-producing and non-producing strains of Aspergillus flavus group. Lett Appl Microbiol 2001;33:291-295.
27. Prabha TR, Revathil K, Vinod MS, Shanthakumar SP, Bernard P. A simple method for total genomic DNA extraction from water moulds. Curr Sci 2012;104:345-347.
28. Batista PP, Santos JF, Oliveira NT, Pires APD, Motta CMS, Luna Alves LinaEA. Genetic characterization of Brazilian strains of Aspergillus flavus using DNA markers. Genet Mol Res 2008;7:706-717.
29. Tran-Dinh N, Carter D. Characterization of microsatellite loci in the aflatoxigenic fungi Aspergillus flavus and Aspergillus parasiticus. Mol Ecol 2000;9:2170-2172.
30. Naik AS, Taware SD. Cytogenetic diversity analysis of Coix species using ISSR markers. Biosci.Biotech Res Asia 2009;6:647-652.
31. Farr DF, Castlebury LA, Rossman AY. Morphological and molecular characterization of *Phomopsis vaccinii* and additional isolates of *Phomopsis* from blueberry and cranberry in the eastern US. Mycologia 2002;94:494-504.
32. Rehner SA, Uecker FA. Nuclear ribosomal internal transcribed spacer phylogeny and host diversity in the *Coelomycete Phomopsis*. Can J Bot 1994;72:1666-1674.
33. Yin Y, Lou T, Yan L, Michailides TJ, Ma Z. Molecular characterization of toxigenic and atoxigenic *Aspergillus flavus* isolates, collected from peanut fields in China. J Appl Microbiol 2009;107:857-865.
34. Gupta M, Chyi YS, Romero-Severson J, Owen JL. Amplification of DNA markers from evolutionary diverse genomes using single primers of simple sequence repeats. Theoretical Appl Genet 1994;89:998-1006.
35. Meyer W, Mitchell TG, Freedman EZ, Vilgalys R. Hybridisation probes for conventional DNA fingerprinting used as single primers in the polymerase chain reaction to distinguish strains of *Cryptococcus neoformans*. J Clin Microbiol 1993;31:2274-2280.
36. Wu KS, Jones R, Danneberger L, Scolnik PA. Detection of microsatellite polymorphisms without cloning. Nucleic Acids Res 1994;22:3257-3258.
37. Archibald JK, Crawford DJ, Santos-Guerra A, Mort ME. The utility of automated analysis of inter-simple sequence repeat (ISSR) loci for resolving relationships in the Canary Island species of *Tolpis* (Asteraceae). Am J Bot 2006;93:1154-1162.
38. Clausing G, Vickers K, Kadereit W. Historical biogeography in a linear system: genetic variation of Sea Rocket (*Cakile maritima*) and Sea Holly (*Eryngium maritimum*) along European coasts. Mol Ecol 2001;9:1823-1833.
39. Reddy MP, Sarla N, Siddiq EA. Inter simple sequence repeat (ISSR) polymorphism and its application in plant breeding. Euphytica 2002;128:9-17.
40. Rogers D. What is a genetic marker? Why we care about genetics. 2006; Vol 5. www.grep.ucdavis.edu/projects/GeneticFactsheets
41. Ziekevich E, Rafalski A, Labuda A. Genome fingerprinting by simple sequence repeat (SSR) anchored polymerase chain reaction amplification. Genomics 1994;20:178-183.
42. Joshi SP, Gupta VS, Aggunaol RK, Rangekar PK, Brar DS. Genetic diversity and phylogenetic relationship as revealed by ISSR polymorphism in the genus *Oryza*. Theo Appl Genet 2000;100:1311-1320.
43. Hatti AD, Taware SD, Taware AS, Pangrikar PP. Genetic diversity of toxigenic and non-toxigenic *Aspergillus flavus* strains using ISSR markers. Int J Curr Res 2010;5:61-66.
44. Hadrich I, Makni F, Sellami H, Cheikhrouhou F, Sellami A, Bouaziz H, Hdiij R, Elloumi M, Ayadi A. Invasive aspergillosis: epidemiology and environmental study in haematology patients (Sfax, Tunisia). Mycoses 2010b;53:443-447