Characterization of seed storage protein patterns of four Iranian Pistachios using SDS-PAGE

Ali Akbar Ehsanpour*, Behrokh Shojaie, Fatemeh Rostami

Department of Biology, Faculty of Science, University of Isfahan, Isfahan, Iran; *Corresponding Author: ehsanpou@yahoo.com

Received 6 March 2010; revised 10 April 2010; accepted 15 April 2010.

ABSTRACT
We used SDS-PAGE to evaluate and characterize the protein patterns of seed storage proteins in four pistachios cultivars (Akbari, Ahmad Aghaei, Fandoghi, and Kaleghouchi). Total protein content of pistachio seeds in all cultivars did not show any significant difference. Results of SDS PAGE pattern of a few protein bands were up regulated whereas some other bands showed down regulation. The identified protein patterns may be used protein marker for pistachio cultivars.

Keywords: Pistachio; Protein Marker; SDS PAGE

1. INTRODUCTION
Pistacia vera L. (pistachio) (2n = 32) is belong to Anacardiaceae family. Iran is one of the largest producer and exporter of pistachio (300,000 tones per year) in the world [1]. Pistachio seed is a very high source of protein, lipid, and vitamins such as vitamin A, B1, B2C, and Niacin. It has also a high percentage of potassium, calcium and phosphorous. Iranian pistachios are different in sizes and shape and they are divided into four major groups including round (Fandoghi), jumbo (Kaleghouchi) and long (Akbari, Ahmad Aghaei) (http://en.wikipedia.org/wiki/Pistachio and http://www.sahravi.com/pistachio/iranian-pistachio.htm).

Plant storage proteins can be classified into two classes; seed storage proteins (SSPs) and vegetative storage proteins (VSPs). SSPs are a set of proteins that accumulate at high levels in seeds during the late stages of seed development, whereas VSPs are proteins that accumulate in vegetative tissues such as leaves, stems and tubers, depending on the plant species. SSP genes were classic targets for work on plant molecular biology. Their abundant expression in seeds allowed for easy detection of the gene transcripts and cDNA cloning during research on plant molecular biology in late 70’s to early 80’s. Characterization of germplasm using biochemical fingerprinting has got special attention due to its increased used in crop improvement and the selection of desirable genotypes for breeding crops. The use of genetic markers and protein profiling has also been successfully used to resolve the taxonomic and evolutionary problems of several crop plants [2-6]. The seed storage protein analyses helps in identification and characterization of diversity in crop varieties, cultivars and their wild varieties and also provides information on phylogenetic relationship of the accessions. It is also known that variation in protein bands provide information on the relationship among the used seeds collected from various geographical regions [6-8]. There are different amounts of storage proteins in all plant seeds. They play two main roles including nitrogen and energy source and defense against insects and pathogens such as bacteria and fungi.

Since, seed storage protein analysis can be a useful tool for identification of species, varieties and cultivars, in this study we investigate the protein pattern in four Iranian pistachios seeds (Akbari, Ahmad Aghaei, Fandoghi, Kaleghouchi) in order to find protein bands as markers for cultivar characterization.

2. MATERIALS AND METHODS

2.1. Plant Material
Fresh mature seeds of Pistachio cultivars including Akbari, Ahmad Aghaei, Fandoghi, and Kaleghouchi were harvested from pistachio garden in Ardestan, Isfahan, Iran.

2.2. Extraction of Seed Proteins
Seed coats from fresh ten seeds from each cultivar were removed and kernels were then grounded in liquid nitrogen with a mortar and pestle. The seed storage proteins from each cultivar were extracted with cold acetone.
by stirring the mixture at 50 rpm at 4°C for 48 h, and subsequently the defatted powder of each cultivar of pistachio was air dried at room temperature for 8 h. Next, the dry powder was suspended in 1:20 (w/v) of 1 ml of 50 mM Tris-HCl buffer containing 1 mM DTT, 2 mM EDTA, 2 mM 2-Mercaptoethanol, pH 7.5. The suspension was stirred at 50 rpm at 4°C overnight and was centrifuged for 25 min at 14000 rpm at 4°C. The precipitate was discarded and the supernatant was used for total soluble protein (mg g⁻¹-dp or defatted powder) assay according to modified Bradford method [9] described by Olson and Markwell [10] using bovine serum albumin as standard protein and SDS-PAGE analysis. SDS-PAGE was performed using 12% separating and 5% stacking gels [11]. After electrophoresis at 120 V, protein bands were stained using silver nitrate and finally the relative density of protein bands were analyzed by ImageJ program (Figure 3). Five protein bands (1, 2, 3, 4 and 5) with approximate MW 45, 33, 32, 27 and 20 kDa respectively, showed maximum expression level in cultivar AA. Protein bands of 6 and 7 with approximate MW 16 and 15 kDa showed maximum expression level in cultivar A, respectively. However, bands 3 and 4 were not detectable in cultivars K and A, while they were detected in the other cultivars. Protein bands 2, 5 and 6 had lower expression level in cultivar F. The minimum level of protein band 1 was observed in cultivar A and protein band 7 in cultivar K.

3. RESULTS

To investigate variations among four Iranian pistachio cultivars, seed storage proteins from cultivar Akbari (A), Ahmad Aghaei (AA), Fandoghi (F) and Kaleghouchi (K) were analyzed. As shown in Figure 1, no significant difference in total protein content was observed in four pistachio cultivars.

The SDS-PAGE protein patterns of four pistachio cultivars showed changes in seven protein bands (Figure 2). Subsequently, the relative levels of protein concentration of these seven protein bands were analyzed by Image J program (Figure 3). Five protein bands (1, 2, 3, 4 and 5) with approximate MW 45, 33, 32, 27 and 20 kDa respectively, showed maximum expression level in cultivar AA. Protein bands of 6 and 7 with approximate MW 16 and 15 kDa showed maximum expression level in cultivar A, respectively. However, bands 3 and 4 were not detectable in cultivars K and A, while they were detected in the other cultivars. Protein bands 2, 5 and 6 had lower expression level in cultivar F. The minimum level of protein band 1 was observed in cultivar A and protein band 7 in cultivar K.

Figure 1. Total soluble protein in four pistachio cultivars Akbari (A), Ahmad Aghaei (AA), Fandoghi (F) and Kaleghouchi (K), defatted powder (dp). Values are the means ± SE. Similar letters (a) indicate no significant difference (P < 0.05) based on Duncan test in four pistachio cultivars.

Figure 2. SDS-PAGE pattern of four pistachio cultivars Akbari (A), Ahmad Aghaei (AA), Fandoghi (F) and Kaleghouchi (K) protein marker (M).

Figure 3. Relative levels of protein expression of pistachio cultivars Akbari (A), Ahmad Aghaei (AA), Fandoghi (F) and Kaleghouchi (K). Values are the means ± SE of proteins bands from three independent experiments.
4. DISCUSSION

Electrophoresis of proteins is a powerful tool for identification of genetic diversity and the SDS-PAGE is particularly considered as a reliable technology because seed storage proteins are highly independent of environmental fluctuations [12,13]. Seed protein patterns can also be used as a promising tool for distinguishing cultivars of particular crop species [14,15]. However, only a few studies indicated that cultivar identification was not possible with the SDS-PAGE method [16]. The SDS-PAGE is considered to be a practical and reliable method for species identification [17].

According to the results of the SDS-PAGE, the overall pattern of seed storage-proteins showed the diversity of pistachio cultivars. The diversity in seed storage proteins has also been reported by Khan et al. for wheat varieties [18]. Moreover, identification of three wheat genotypes including ILC-195, CM-2000 and CM-98/99 has also been reported by protein markers [19].

Since in mature seeds, type and amount of proteins are more constant than other plant tissues [20] therefore, the SDS-PAGE pattern of seed storage proteins of pistachio showed polymorphism on the basis of difference in protein intensity among genotypes. The presence or absence of protein bands has also been applied for detection of polymorphism of Brassica cultivars [21].

The present investigation revealed variation in different cultivars of pistachio seeds with regard to their total seed protein profiles. Regarding interspecific variation among cultivars this investigation revealed some variations. The genetic affinities within cultivars of the same species generally corroborated the morphological analysis. Similar to our finding the result of differentiation of yellow sarson and brown seeded types of Brassica clearly separated the yellow seeded and brown seeded varieties by SDS PAGE [22]. However, we can conclude that, SDS-PAGE can reveal the differences among seed storage proteins of pistacia cultivars.

5. ACKNOWLEDGEMENTS

Authors wish to thank University of Isfahan for their support.

REFERENCES


