Phylogenetics of rice (Oryza sativa L.) genotypes from Pakistan based on diversity of biochemical markers

Haseena Gulzar  
Department of Biotechnology, Shaheed Benazir Bhutto University, Sheringal Dir Upper, P.O Box 18000 Khyber Pakhtunkhwa, Pakistan

Sidra Pervez  
Department of Biochemistry, Shaheed Benazir Bhutto Women University, Peshawar, Khyber Pakhtunkhwa, Pakistan

Rehana Masood  
Department of Biochemistry, Shaheed Benazir Bhutto Women University, Peshawar

Asad Jan  
Institute of Biotechnology and Genetic Engineering, Agriculture University Peshawar, P.O. Box 706 Khyber Pakhtunkhwa Pakistan

Faheeda Soomro  
International Center for Chemical and Biological Sciences, University of Karachi, Karachi 75270, Pakistan

Muhammad Asif Nawaz  
Department of Biotechnology, Shaheed BB University, Sheringal Dir.

Research

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Abstract

In this study phylogenetics and diversity of 150 rice (Oryza sativa L.) genotypes originated from different regions of an agricultural country Pakistan, self-sufficient in its production were analyzed. Total seed protein profile of these genotypes was assessed using a technique of sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE). The seeds of accessions from different geographical regions of the country were provided by the Institute of Agri-Biotechnology and Genetic Resources, National Agricultural Research Centre (IABGRI-NARC) Islamabad, Pakistan. Total endosperm proteins from each genotype were extracted and used as biochemical markers. Three different protein extraction protocols were followed in comparison to optimize the procedure. An immense content of genetic variability was confirmed based on polymorphism data of polypeptide bands. The Jaccard's dissimilarity coefficient ranged from 0.09–0.94 for all 150 accessions. Based on the neighbour joining method of clustering, the whole germplasm was divided into two clusters, each further subdivided sorting centre of origin and depicting diversity. In the same manner, 126 accessions of province Khyber Pakhtunkhwa (KPK) were also assessed separately. This wealth of knowledge based on proteins profiling will be used for marker assisted breeding/selection, germplasm conservation strategies and easy identification of these genotypes from Pakistan.

Introduction

Rice, a member of family Poaceae is the staple food for more than half of the world’s population and South Asia being the biggest rice producer and consumer. Nevertheless, it is cultivated all over the world on about 154 million hectares every year occupying 11% of the world’s cultivated area, and 10% only in Pakistan. It adds 5.9% of value in agriculture and 1.3% in GDP in Pakistan, a value larger than other South Asian countries therefore, also called ‘Golden Grain of Pakistan (Simmonds 1986; Shah et al. 2011). Among the cereal crops rice has the smallest genome, well characterized and sequenced thus a model plant for genomic research having 24 chromosomes (Eckardt 2000; Swain et al. 2017). Pakistan is among one of the few South Asian countries, self-sufficient in rice production with increasing yield since 2001 and is the second largest exporter with a wealth of diverse traditional landraces still cultivating in villages (Shah et al. 2011). More than 2,500 accessions collected from different areas of Pakistan are preserved in the national gene bank, IABGRI-NARC, Islamabad. Moreover, 100,000 traditional rice varieties are stored in the gene banks throughout the world (Pervaiz et al. 2011; Bishwajit et al. 2013). In the last two decades many researchers have put their efforts to assess the genetic variability present in rice germplasm of Pakistan (Khan et al. 2013; Tahir 2014; Bibi et al. 2017).

Landraces are valuable genetic resources containing huge genetic variability also hold several unwanted agronomic traits like tall plant stature, long crop duration etc however, also exhibit strong characteristics such as resistance to biotic and abiotic stresses. In the last decade of 20th century, Pakistan has released a number of superior cultivars by crossing advanced breeding lines and traditional varieties, achieving high quality of grains and resistance to biotic and abiotic stresses (Pervaiz et al. 2011; Haque et al. 2015). SDS-PAGE is useful, easy, low cost and a quick way for studying genetic diversity in numerous
genotypes based on polymorphic endosperm storage proteins which are unlike morphological traits not under the influence of environment (Ali et al. 2007; Gulzar et al. 2015). Therefore, the information provided by protein profiling can truly be beneficial in the screening of desirable traits in these germplasm resources for efficient utilization in advanced crop improvement, genetic engineering and breeding strategies (Conrad et al. 2013; Gul et al. 2015).

This work is one of the efforts to attain these goals as to ensure food security, systematic evaluation of the valuable genetic resources is pre-requisite for using in crop improvement programs (Bibi et al. 2017).

**Materials And Methods**

This study was conducted at the Institute of Biotechnology and Genetic Engineering (IBGE), Agriculture University Peshawar, KPK, Pakistan. Rice germplasm was provided by IABGRI-NARC, Islamabad, Pakistan.

The work consisted of two experiments; optimization of protein extraction protocol and biochemical evaluation of 150 rice accessions originated from Pakistan. The accessions belonging to KPK province (126) were analyzed separately including check variety Fakhr-e-Malakand based on SDS-PAGE of endosperm storage proteins.

**Optimization of protein extraction protocol**

For optimization three different compositions of protein extraction buffer (PEB) were used to extract proteins from rice endosperms in order to obtain clear bands.

**PEB 1**: 0.5M Tris-HCl pH 8.0, 0.2% SDS, 5M urea, 1% 2-mercaptoethanol (2-ME) (Ranjan et al. 2012).

**PEB 2**: Phosphate buffered saline (PBS): 137mM NaCl, 2.7mM KCl, 10mM Na₂HPO₄ and 1.8 mM KH₂PO₄ with pH 7.4 (Walczyk et al. 2017).

**PEB 3**: Cold spring harbour protocol for 2X sample buffer; 1M Tris, pH 6.8, 50% glycerol, 10% SDS, 0.5% bromophenol blue (BPB) and 0.4 ml, 2-ME (Cold Spring Harb Protoc 2009)

Using PEB 1 and 3, the protocol followed include dehusking of rice seeds, ground into fine powder with mortar and pestle and centrifugation (0.01 g seed meal/400 µl PEB) at 10,000 rpm for 10 minutes to obtain supernatant and stored at -20°C (Pervaiz et al. 2011).

While using PEB 3 or PBS, the protocol followed include the steps of dehusking rice seeds, ground into fine meal, defatting rice meal with n-hexane twice, protein extraction by shaking 0.01 g defatted seed meal/500 µl PBS for 2 hours at 4°C and centrifugation at 12,000 rpm for 10 minutes to obtain supernatant and stored at -20°C until use. Before loading into gel, the PEB 1 was used as loading dye with proteins extracted in PBS in the same ratio (1:1) (Walczyk et al. 2017).

**Biochemical characterization using SDS-PAGE**
Total rice seed proteins were evaluated by one dimensional discontinuous system of SDS-PAGE with some modifications in Laemmli’s method (1970). Extracted proteins (12µl each) were loaded in 5% stacking and 12% separating gel, run at 25mA and100V for 60 to 90 min. Bio-Rad dual precision protein ladder was used as molecular weight marker (10 to 250 kDa). The gels were stained overnight in 0.02% Comassie Brilliant Blue (CBB) by continuous shaking and destained in distilled water until the background was clear (Khan et al. 2013).

**Data analysis for phylogenetics and diversity**

An electrophoregram was analyzed for the presence/absence (1/0) of polypeptide bands in each accession. The binary data made was used to generate similarity and dissimilarity matrix which was further processed to construct a phylogenetic tree by the unweighted pair group method with arithmetic means (UPGMA) introduced by Sneath and Sokal in (1973) using a Statistical Programme for Social Scientists (SPSS) and Ward’s method (Rahman et al. 2010; Gulzar et al. 2015).

**Results**

In this paper 150 different local rice (*Oryza sativa* L.) accessions originated from different regions of Pakistan—including north western regions (Province of KPK), including Parachinar and Kurrum agency, northern areas (Gilgit-Baltistan) and Punjab province—were analyzed for phylogenetic relationship and genetic diversity using SDS-PAGE of total seed proteins (Table 1). Most of these were from KPK Province (126) including a check variety Fakhr-e-Malakand and only 24 from the other regions of the country.

**Optimization of protein extraction method**

Three different protein extraction protocols were used and the procedure optimized. The electrophoregrams generated by first protocol and third (Ranjjan et al. 2012), resulted in clear bands however, we recommend the use of second protocol with PBS pH 7.4 (Walczyk et al (2017) for genetic variability studies in rice which gave us more clear and sharp bands (Figs. 1 & 2). Moreover, up to 12 samples can be run in a single gel when working with protocol 2, thus saving time and resources. While with protocol 1, only six samples can be run at a time because it produce more spread out and denser bands (Figs. 1 & 2).

We found that defatting seed meal with n-hexane improved protein extraction process with no floating fats on surface and gave clear bands.

**SDS-PAGE based Polymorphism**

In this study the polymorphism present in 150 local rice (*Oryza sativa* L.) accessions of Pakistan were biochemically analyzed for seed storage proteins using SDS-PAGE. A huge content of genetic variability was seen based on polypeptides banding pattern, size (10 to 250 kDa) and number (14 to 27). Both major and minor bands were kept under consideration, thus results showed comparatively significant variation in minor bands although some variation in major bands was there (Figs. 1 and 2) (Tahir et al. 2014). For
ease of scoring the entire banding pattern was divided into four zones, zone A (60-to-250 kDa), B (30-to-60 kDa), C (18-to-30 kDa) and D (10-to-18 kDa) as shown in Figs. 1 and 2. Maximum number of bands were found in zone A and B (each 11 bands) followed by C (08) and D (04) in different accessions. Therefore, during analysis 34 bands were observed with minimum number of 14 bands and maximum of 27 bands in different accessions (Table 2). Out of 34 bands 10 (29.4%) were monomorphic and 24 (70.5%) were polymorphic.

In zone A, nine bands showed polymorphism while eight, six and one band was polymorphic in zone B, C and D respectively. In present investigation, maximum number of bands were seen in the accessions belonging to province KPK, Chitral including accession no. 7284 with 27 bands followed by 25 bands in accessions no. 7626, 7805, 7794. While 24 bands were observed in accession no. 7598 (KPK, Malakand), 17940 (KPK, Swat), 7886 (Pakistan Agriculture Research Council (ARC) and 7801, 7791, 7792, 7808, 7811 accessions from Chitral District (Tables 1, 2, 3 & 5). Minimum number of bands i.e. 14 was observed in accessions number 7281 (KPK, Mansehra), 7291 & 7292 (KPK, Swat) and 7829 (KPK). It shows that the genetic resources of rice originated from district Chitral possess more diversity than that of other districts of the country (Bibi et al. 2017). In a check variety Fakhr-e-Malakand 17 (2, 6, 6, 3) bands were observed corresponding to zones A –D (Table 1–3, Figs. 1 & 2).

Rf values of the last bands of each zone (A, B, C, D) were recorded as 0.32, 0.58, 0.75 and 0.96 respectively, shown in Figs. 1 & 2 (Chen et al. 1987). The sum total of maximum bands observed in each zone is equal to 34 (A; 11, B; 11, C: 08, D; 04).

**Heirarchical Clustering:**

**Cluster analysis for the whole germplasm**

Cluster analysis (Ward's method) Ward, and Amer (1963) for the whole germplasm consisting of 150 rice landraces of Pakistan was carried out using binary data processed by SPSS software. A dendrogram was developed which divided the whole Germplasm into two major clusters; cluster A consisted of 58 accessions while the rest 92 genotypes grouped in cluster B, each further classified into two (A1, A2 &B1, B2) sub-clusters having 42, 16, 48 and 44 accessions respectively. A check variety Fakhr-e-Malakand (FM) was grouped in cluster B2 as shown in table 4. The Jaccard's dissimilarity coefficient ranged from 0.09 (lowest) to 0.94 (highest), while that of similarity coefficient from 0.009 to 0.635, revealing highest genetic diversity among the rice landraces of Pakistan.

**Cluster analysis for the accessions of Province KPK**

In addition to whole germplasm molecular polymorphism studies, phylogenetics studies for 126 accessions of KPK were also carried out and separately analyzed by cluster analysis.

A dendrogram was developed which divided the KPK Germplasm into two major clusters; the major cluster A consisted of 105 accessions and cluster B consisted of 21 accessions. Cluster A was further divided into two (A1, A2) sub-clusters with 59 (A1a; 26, A1b; 33) and 46 (A2a;16, A2b; 30) accessions
while cluster B into two (B1, B2) sub-clusters with 06 and 15 accessions respectively. A check variety Fakhr-e-Malakand (FM) was grouped in A2b (Fig. 5 and Table 5&6). Pair-wise genetic distance calculated to comprehend the genetic diversity ranged from 0.185 (lowest) to 0.965 (highest) and similarity coefficient from 0.018 to 0.632, thus revealing intraspecific genetic diversity within the accessions of KPK.

Discussion

In this study 150 diverse local rice (*Oryza sativa* L.) accessions originated from different regions of Pakistan were analyzed for genetic diversity using SDS-PAGE. Maximum number (126) of accessions was from province KPK, including a check variety Fakhr-e-Malakand. Fakhr-e-Malakand, a new cold tolerant and early maturing japonica rice variety was introduced in 2003 by the government of KPK, Pakistan, cultivated at more than 1000 meters altitude, and irrigated by snow fed rivers. It gives higher head rice (56.8%) on milling, higher quality index (1.33) and the best results in comparison to other rice varieties under the system of rice intensification (SRI) practices (Ahmad and Khan 2016). Fakhr-e-Malakand produced heavier grains and high yield as compared to other varieties when studied their sowing dates in relation to bacterial leaf blight (Raﬁ et al. 2013). The outstanding features of Fakhr-e-Malakand like higher yield, lodging and pests resistance, cold tolerance, along with more Benefit Cost Ratio of 3.24, made it most popular and economical commercial variety of the rice farming community (Ahmad et al. 2015).

In order to obtain of clear bands on the polyacrylamide gel, protein extraction method was first optimized. Three different protocols gave different results. One protocol gave denser bands and another sharp bands, on the basis of which, we proposed the use of second protocol with PBS pH 7.4 (Ranjan et al. 2012; Walczyk et al (2017). Defatting seed meal with n-hexane improved protein extraction process and gave clear bands. Hirano et al (2000) and Walkzyk et al (2017) reported the same results and found that defatting peanut meal with n-hexane resulted in significantly higher yields of crude proteins and allergens than other methods. Isotonic solutions like PBS is extensively used in the extraction of lectin proteins and food allergens which resulted in greater sensitivity of their molecular analysis than with Tris-Borate EDTA (TBS) or Tris. The yields of the compounds of interest were also affected by the composition of different buffers, their pH and other conditions.

One hundred and fty local rice (*Oryza sativa* L.) accessions of Pakistan, analyzed for seed storage proteins using SDS-PAGE showed significant polymorphism. A total of 14–27 bands were found in each accession with size range from 10 to 250 kDa. Pervaiz et al (2010-11) made one of the efforts to explore the genetic polymorphism of some Pakistan's landraces based on biochemical and molecular markers including Wx gene products (60 kDa), glutelin, albumin, globulin and prolamine polypeptide subunits as well as RAPD fragments.

Shah et al (2011) also reported 34 bands in different species of rice including *Oryza sativa* L. However, we have not seen 34 polypeptide bands in any accession as we have observed a maximum of 27 bands.
Rice accessions originated from district Chitral showed more genetic variation as compared to other districts of the country (Bibi et al. 2017). Pakistan's rice germplasm has proved to be such a diverse that there can possibly be more than what is explored to date. After Shah et al (2011), maximum number of polypeptide bands in various Pakistan's landraces has been reported by Habib et al (2000), equal to 32, Pervaiz et al in (2011), 25, followed by Asghar et al (2004) who observed 22 bands and Bibi et al (2017) reported overall 18 bands. Sharief et al (2005) reported a minimum of 12 bands. The primary reason for variation in number of polypeptide subunits reported by other authors is the diversity of the rice germplasm of Pakistan from different geographical locations. Moreover, it can also be due to difference in protein extraction buffer as we have used three different compositions, and other lab protocols and gel composition for the evaluation rice seed proteins (Asghar et al. 2004).

No major relationship of geographical location and variation in seed protein profile was examined. The rice genotypes of China have been found relatively low in genetic diversity as compared to the landraces of the South Asian countries (Swain et al. 2017; Warusawithana et al. 2017).

A phylogenetic tree was developed for 150 rice accessions which divided the whole Germplasm into two major groups with 58 and 92 genotypes respectively, including check variety Fakhr-e-Malakand in second group (Ward, and Amer, 1963). The Jaccard’s dissimilarity coefficient (0.09 to 0.94) and similarity coefficient (0.009 to 0.635), revealed highest genetic diversity among the rice genotypes of Pakistan. The beneficial characteristics of FM are expected to be present in the same group of accessions on the basis of hierarchical clustering, in which it has been grouped due to less genetic distance (Ahmad et al. 2016). Similar work has been done in these references on Oryza sativa L (Bibi et al. 2017). The accessions from different geographical zones were clustered together in same groups. This demonstrates that expression profile is least affected by environment, a statement given by a range of studies on other cereal crops and rice. Some work also linked protein markers to the agronomic traits clustering in the same group which need to be explored further (Pervaiz et al. 2011). Cluster analysis for 126 accessions of KPK were also carried out and a cladogram was developed which divided the KPK Germplasm into two major groups consisting of 105 accessions and 21 accessions respectively including check variety Fakhr-e-Malakand in the first group (A2b). Pair-wise genetic distance (0.185–0.965) and similarity coefficient (0.018 to 0.632), was also found to express intraspecific genetic diversity within.

As FM is an improved variety with desirable characteristics, the same traits are also expected to be present in group A2b, due to lowest genetic distance with them (Ahmad et al. 2015; Ahmad et al. 2016). Perwaiz et al (2011) reported a similarity coefficient of 0.81; this low genetic distance may be credited to the same genetic background, long-established farming practices and consumer preferences.

**Conclusion**

The findings in this work unveiled a considerable amount of genetic diversity in 150 landraces of the O. sativa L. from Pakistan and that of Province Khyber Pakhtunkhwa in particular. These facts offer an opportunity to be used for marker aided selection or breeding strategies and improvement of modern rice
cultivars very low in genetic variability. The information obtained from clustering behaviour of accessions can be helpful to design plans for their management in the gene-bank. The desirable and rich traits in these landraces can also be identified and exploited in order to ensure food security and to cope with changing environmental conditions thus providing an efficient utilization and emphasizing on the conservation of germplasm resources of Pakistan for the betterment of humanity.

**Abbreviations**

SDS-PAGE, sodium dodecyl sulphate polyacrylamide gel electrophoresis, (IABGRI-NARC) Agri-Biotechnology and Genetic Resources, National Agricultural Research Centre (IABGRI-NARC), Khyber Pakhtunkhwa (KPK), GDP, Gross Domestic Product, (IBGE), Institute of Biotechnology and Genetic Engineering, protein extraction buffer (PEB), Phosphate buffered saline, bromophenol blue (BPB), Comassie Brilliant Blue (CBB), unweighed pair group method with arithmetic means (UPGMA), Statistical Programme for Social Scientists (SPSS), Federally administered tribal areas (FATA), Fakhr-e-Malakand (FM), (Pakistan Agriculture Research Council (ARC), system of rice intensification (SRI)

**Declarations**

**Acknowledgements**

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**Author contributions**

MAN and SP conceived and designed research, HG and RM conducted experiments while AJ and FS analysed the data. HG and MAN wrote the manuscript. All authors read and approved the manuscript.

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**Availability of data and materials:**

The datasets used and/or analysed during the current study are available from the corresponding author on request.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**
Not applicable.

**Competing Interests**

This publication is approved by all authors and they do not have any conflict of interest regarding any financial, personal or other relationships with any other people or organizations.

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Tables

Due to technical limitations, table 1, 2, 3, 4, 5 , 6 is only available as a download in the Supplemental Files section.

Figures

Figure 1

Rice (Oryza Sativa L.) electrophoregram of endosperm proteins from Pakistan s accessions (7600-7599) extracted by protocol 1, divided into four zones (A, B, C, D) and Rf values of major bands given
Figure 1

Rice (Oryza Sativa L.) electrophoregram of endosperm proteins from Pakistan’s accessions (7600-7599) extracted by protocol 1, divided into four zones (A, B, C, D) and Rf values of major bands given.

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Figure 2

Rice (Oryza Sativa L.) electrophoregram of endosperm proteins from 12 Pakistan’s accessions (7815-7849) extracted by protocol 2, showing variation divided into four zones (A2B2C2D2) and Rf values of major bands given.

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Rice (Oryza Sativa L.) electrophoregram of endosperm proteins from 12 Pakistans accessions (7815-7849) extracted by protocol 2, showing variation divided into four zones (A2B2C2D2) and Rf values of major bands given.
Figure 3

A dendogram showing phylogenetic relationship and genetic diversity among 150 rice (Oryza Sativa L.) accessions of Pakistan based on SDS-PAGE
A dendogram showing phylogenetic relationship and genetic diversity among 150 rice (Oryza Sativa L.) accessions of Pakistan based on SDS-PAGE.
Figure 4

Graph showing relationship between Agglomeration schedule coefficients (values) and stages of clustering revealing diversity in 150 Pakistani rice (Oryza Sativa L.) accessions
Figure 4

Graph showing relationship between Agglomeration schedule coefficients (values) and stages of clustering revealing diversity in 150 Pakistani rice (Oryza Sativa L.) accessions
Figure 5

A dendrogram showing phylogenetic relationship and genetic diversity among 126 rice (Oryza Sativa L.) accessions of province Khyber Pakhtunkhwa (KPK) Pakistan based on SDS-PAGE
Figure 5

A dendogram showing phylogenetic relationship and genetic diversity among 126 rice (Oryza Sativa L.) accessions of province Khyber Pakhtunkhwa (KPK) Pakistan based on SDS-PAGE.
Figure 6

A graph showing relationship between agglomeration schedule coefficients (values) and stages of clustering revealing diversity in 126 rice (Oryza Sativa L.) accessions of KPK province from Pakistan.
Figure 6

A graph showing relationship between agglomeration schedule coefficients (values) and stages of clustering revealing diversity in 126 rice (Oryza Sativa L.) accessions of KPK province from Pakistan

Supplementary Files

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