Compositional studies and biological activities of some mash bean (Vigna mungo (L.) Hepper) cultivars commonly consumed in Pakistan

Muhammad Zia-Ul-Haq1*, Shakeel Ahmad2, Shazia Anwer Bukhari3, Ryszard Amarowicz4, Sezai Ercisli5* and Hawa ZE Jaafar6*

Abstract

Background: In recent years, the desire to adopt a healthy diet has drawn attention to legume seeds and food products derived from them. Mash bean is an important legume crop used in Pakistan however a systematic mapping of the chemical composition of mash bean seeds is lacking. Therefore seeds of four mash bean (Vigna mungo (L.) Hepper, family Leguminosae) cultivars (NARC-Mash-1, NARC-Mash-2, NARC-Mash-3, NARC-Mash-97) commonly consumed in Pakistan have been analyzed for their chemical composition, antioxidant potential and biological activities like inhibition of formation of advanced glycation end products (AGE) activity and tyrosinase inhibition activity.

Results: The investigated cultivars varied in terms of biochemical composition to various extents. Mineral composition indicated potassium and zinc in highest and lowest amounts respectively, in all cultivars. The amino acid profile in protein of these cultivars suggested cysteine is present in lowest quantity in all cultivars while fatty acid distribution pattern indicated unsaturated fatty acids as major fatty acids in all cultivars. All cultivars were found to be rich source of tocopherols and sterols. Fourier transform infrared spectroscopy (FTIR) fingerprints of seed flour and extracts indicated major functional groups such as polysaccharides, lipids, amides, amines and amino acids. Results indicated that all investigated cultivars possessed appreciable antioxidant potential.

Conclusions: All cultivars are rich source of protein and possess sufficient content of dietary fiber, a balanced amino acid profile, low saturated fatty acids and antioxidant capacity that rationalizes many traditional uses of seeds of this crop besides its nutritional importance. The collected data will be useful for academic and corporate researchers, nutritionists and clinical dieticians as well as consumers. If proper attention is paid, it may become an important export commodity and may fetch considerable foreign exchange for Pakistan.

Keywords: Nutrients, Antioxidant potential, Mash bean cultivar, Pakistan
Seeds are believed as spermatopoetic, and used for treating erectile dysfunction and premature ejaculation. Seeds are used for lengthening the hair, keeps them black and curing dandruff. Hot aqueous extracts of the leaves are used in the treatment of brain disorders, stomach, jaundice, rheumatic pain and inflammatory disorders. Seeds are considered fattening and flour made from seeds is excellent substitute for soap, leaving the skin soft and smooth and used in cosmetics in preparation of facial mask [2-6].

The mash bean occupies an important position in agriculture system of Pakistan and is grown annually on area of 27.6 thousand hectares with annual production of 13,600 tonnes with 493 kg/ha as average yield [7]. It is grown all over the country, but its cultivation is concentrated mainly in Punjab, the major mash production province. It is the least researched crop among pulses in Pakistan as is apparent from scarcity of literature on it and as a result its area of cultivation and production are decreasing gradually [8].

The food industry globally is searching functional foods, nutraceuticals and botanicals to meet demand of consumers for natural, immunity-boosting and health-promoting plant based food products. To our knowledge, there is no study indicating chemical composition and antioxidant potential of seeds of mash bean cultivar indigenous to Pakistan. As part of our research studies to investigate the biochemical composition and antioxidant capacity of indigenous flora of Pakistan [9-13] this study has been conducted to determine the chemical composition, antioxidant activity and biological activities of seeds of mash bean cultivars.

**Results and discussion**

Composition and contents of various constituents and components like various bioactive constituents and secondary metabolites, fixed and essential oil, fatty acids, tocopherol and sterol profile, mineral, amino acid, vitamin, protein and carbohydrate contents present in a food commodity like seed, fruit, vegetable, spice, grain or any other product derived from them varies depending upon many factors like plant variety, agronomic practices utilized in cultivation, stage of collection and geological and climatic conditions of area from where that food commodity or plant part (seed or fruit) is collected, and the method employed for its determination. So there is need to establish food composition database on regional and country level for various food commodities for various regions and countries respectively. Previously our research group has compiled compositional and nutritional information on various other legumes like chickpea, pea, cowpea, lentil and mung bean. In current study we have determined biochemical composition, their impact on health as well biological activities of a less-researched legume crop i.e. mash bean.

The data on the proximate composition is summarized in Table 1. The observed range for protein was 24.62% for NARC-Mash-97 to 25.48% for NARC-Mash-2 Mash 2. The crude fiber content ranged from 4.25% to 5.09%. The range observed for fat content was between 1.80 and 2.25% while carbohydrates ranged from 53.43% to 55.55%. The high carbohydrate contents present in mash bean seeds indicate its potential use as a prime source of energy to prevent marasmus in infants especially. Like other legumes, its seeds are also rich in protein, contain sufficient amount of dietary fibre and lesser amount of oil. The results are in partial agreement to those reported earlier for mash bean [14,15] and other legumes [9-13]. Regular intake of dietary fibre is associated with low chances of cardiovascular disease, obesity, certain cancers and diabetes. High dietary fiber contents may be responsible for its traditional use as anti-cancer food. Since dietary fibre containing foods are used in bakery products, it also indicates its potential use in bakery and pastry products.

The data of vitamin contents is summarized in Figure 1. Niacin content was highest in NARC-Mash-1 (1.80 ± 0.07 mg/g) while NARC-Mash-3 had lowest content of riboflavin (0.19 ± 0.19 mg/g). Regarding vitamin contents of seeds of mash bean, niacin was present in higher concentration among all cultivars. As there is no report available on vitamin contents of mash bean, so vitamin contents cannot be compared to previous results. However the vitamin contents are in close proximity to that of *Pisum sativum* as per our previous studies [16]. High contents of niacin are good from medical point of view as this water-soluble vitamin is excreted by urine from human body and its continuous supply by eating mash bean seeds will complete its deficiency. Various agro-geo-climatological conditions affect vitamin contents in legume seeds.

Mineral contents (Table 2) indicated potassium as major mineral from 1599.82 ± 1.74 mg/100 g in NARC-Mash-97 to 1646.01 ± 0.92 mg/100 g in NARC-Mash-3. Phosphorus ranked second in quantity from 439.79 ± 0.42 and 500.17 ± 1.85 in same varieties. Zinc was present in lowest content (1.94 ± 0.76 mg/100 g) in NARC-Mash-97. All cultivars contained sufficient contents of potassium, phosphorus and copper. It is perhaps this high potassium content that makes it an aphrodisiac. The high content of potassium is useful for patients who use diuretics to manage hypertension and there is unnecessary seepage of potassium from their body fluids. The low content of sodium compared to potassium led to a low sodium: potassium ratio, which is favorable from nutritional point of view, as foods with low Na:K ratio are linked with lower frequency of blood hypertension. Na:K ratio is from 0.14 to 0.17 in NARC-Mash-1 and NARC-Mash-3 respectively. For prevention of high blood pressure, Na/K ratio of less than one is suggested. This may explain the
rationale behind the traditional use of its seeds in managing hypertension. Low Ca:P ratio leads to loss of Ca in the urine more than normal amount, so Ca concentration in bones is reduced. Food is considered “poor” if Ca:P ratio is less than 0.5 and “good” if it is above one. In present study, Ca:P ratio ranged from 0.78 to 1.00 in NARC-Mash-2 and NARC-Mash-1 respectively indicating regular consumption of mash bean seeds will serve as fine source of calcium for formation of bones. High levels of calcium are required during growth, gravidity and lactation of animals [13]. The results are in par to those already reported for mash bean elsewhere [17,18]. It is well-known that mineral contents of plant and crops parts like fruit and seeds depend on cultivars, collection time and maturity stage, climatological conditions, agronomic practices like type of fertilizer and water requirements of the amino acids, a human body needs. A protein-rich diet is not a guarantee to fulfill the requirements of the amino acids, a human body needs. A balanced protein diet should comprise all amino acids in sufficient amount and essential and non-essential amino acid ratio denotes the nutritional quality of protein. Glutamic acid (19.19 ± 0.62 to 21.49 ± 0.07 g/100 g) and aspartic acids (11.53 ± 0.11 to 13.20 ± 0.27 g/100 g) were present in highest amount in all cultivars. Except tryptophan and S-containing amino acids, all essential amino acids are present in sufficient amounts in all analyzed cultivars as is evident by data (Table 3). Most amino acids derived from plant sources are believed to possess antimicrobial, anti-inflammatory, immune-stimulating and antioxidant properties besides their role in nutrition. Results are comparable to those of previous studies on mash bean amino acids [17]. The deficient amino acids can be acquired by including large quantity of mash bean in diet, or by taking mash bean as well as other legumes.

Besides amino acid composition, protein digestibility is crucial for determining the protein quality. In-vitro protein digestibility data (Figure 2) suggested that values are lowest in NARC-Mash-1 (29.30 ± 0.82%) and highest in NARC-Mash-97 (38.53 ± 0.21%) while starch digestibility was 59.93 ± 0.17 to 67.09 ± 0.02 for same cultivars. Protein digestibility was below 50 percent while starch digestibility was above 50 percent in all analyzed cultivars. A significant variation has been observed for protein digestibility of legume seeds previously for mash bean and other legumes [16,17]. The sensory, textural and nutritional characteristics of products made from legumes are due to various functional properties of proteins. Anti-nutritional components like tannins, phytates and tryspin inhibitors, and structural distinctiveness of storage proteins slow down the digestibility of legume proteins. Treatments like roasting; autoclaving and cooking may be utilized to increase the legume proteins digestibility. In vitro starch digestibility values are close to those reported earlier [20]. Since legume starches generally contain more amylase, therefore these are less digestible. This low digestibility is useful as it decreases release of glucose in blood and so is helpful for patients suffering from diabetes. It may be reason of prescribed use of mash bean for diabetic patients by traditional healers. The low-digestibility however may be managed by utilization of legume seeds along with husk since dietary fibre present in husk will decrease the transit time in intestines and will help in bowel motility.

Fatty acids profile determines the oil quality of seeds or fruits or any other part of plant and products derived from them. Fatty acids profile of oil of seeds of investigated cultivars is summarized in Table 4. All cultivars were found to be rich source of δ-linolenic acid (49.52 ± 0.09 to 51.80 ± 0.03%) and oleic acid (26.62 ± 0.07 to 27.34 ± 0.25%). Bulk of the oil consisted of unsaturated fatty acids for all cultivars. The results are comparable to previously published works for low-oil bearing legumes in general [21] and for mash bean in particular [22]. Saturated fatty acids were a small percentage of total fatty acids present. There is reduced risk of cholesterol-related heart diseases by consuming oils containing more unsaturated fatty acids. However since oil content is very low in seeds therefore it cannot be considered as commercial source of vegetable oil.

Data about tocopherol composition is summarized in Table 5. Despite differences, γ-tocopherol contents were present in highest quantity in all cultivars while considerable contents of δ-tocopherol followed α-tocopherol were also noted. Oil of seeds of all mash bean cultivars studied contained all major tocopherols. Like many other traits,

| Component       | NARC-Mash-1          | NARC-Mash-2          | NARC-Mash-3          | NARC-Mash-97         |
|-----------------|----------------------|----------------------|----------------------|----------------------|
| Crude protein   | 27.91 ± 1.71         | 26.48 ± 1.66         | 25.07 ± 1.60         | 28.60 ± 1.72         |
| Total lipids    | 5.13 ± 0.05          | 6.00 ± 0.05          | 5.80 ± 0.09          | 6.22 ± 0.09          |
| Carbohydrates   | 56.55 ± 1.82         | 54.81 ± 1.73         | 58.13 ± 1.10         | 54.81 ± 1.75         |
| Crude fiber     | 5.44 ± 1.20          | 6.84 ± 1.60          | 4.25 ± 1.20          | 5.11 ± 1.60          |
| Ash             | 4.97 ± 0.19          | 5.87 ± 0.18          | 6.72 ± 0.19          | 5.26 ± 0.18          |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).
no previous report is present regarding tocopherol and sterol contents of mash bean seeds. However all values are close to those reported for Indian mash bean seeds [22]. Since naturally occurring tocopherols are used for oils and fats stabilization against oxidative degradation, it suggests their usage in pharmaceutical, biomedical, and nutritional products.

Sterol profile is summarized in Table 6. Substantial amounts of campesterol, avenasterol and stigmasterol were found in oils of seeds of all four cultivars. The main sterol in oil of seeds of all investigated mash bean cultivars was β-sitosterol which is in agreement with previous studies for low-oil bearing legumes like chickpea, mungbean, cowpea, *Albizia lebbeck* and *Acacia leucophloea* in general [10,12,13,21] and for mash bean in particular [23]. Various agro-geo-climatological factors as well as solvent used for extraction of oil are believed to be responsible for the distribution of tocopherols and sterols in oils extracted from plant parts. Sitosterol, campestrol and stigmasteprosterol have been observed to be major sterols in oils from most of plants belonging to family *Leguminosae* [10,12,13,21].

FTIR-fingerprints, give a quick check of identification, classification and discrimination of food samples by providing a general outline of pattern and trends indicating presence of various chemical compounds in samples. FTIR spectrum of mash bean seed powder indicated the presence of various types of aliphatic and aromatic compounds, especially carboxylic acids, esters, alkyl halides and nitro compounds. The presence of carboxylic acids is indicated by peaks at 2929.40 (O-H stretching), 1249.94 cm\(^{-1}\) (C-O stretching). The peak at 1728.04 corresponds to C = O stretching frequency of aldehydic group. Unsaturated compounds presence is indicated by the peak at 1658.57 cm\(^{-1}\) (C = C stretching, alkene) and 1556.41 cm\(^{-1}\) (C-C stretching, aromatic compounds). Saturated compounds presences is shown by the peak at 1450.17 cm\(^{-1}\) (C-H bending, alkane). Nitro compounds and aromatic amine presence is indicated by the peaks at 1343.93 and 1319.41 cm\(^{-1}\). The peaks at 1160.05 and 1074.23 cm\(^{-1}\) showed the presence of aliphatic amines. Alkyl halides presence is pointed out by the peak at 849.49 cm\(^{-1}\).

For mash bean extract, the highly intensified OH region with intensified shoulder peak of amine group was present. A new peak in the region of 1700 – 1800 was observed which may be attributed to presence of ester. Saturated compounds presence is indicated by the peaks at 2925.31, 2859.93 (C-H stretching, alkanes) and 1384.79 cm\(^{-1}\) (C-H rocking, alkane). Carboxylic acid presences is confirmed by the peaks at 3023.38 (O-H stretching), 1736.21 and 1695.35 cm\(^{-1}\) (C = O stretching). Nitro compounds presences are indicated by the peak at 1515.55 cm\(^{-1}\) (N-O asymmetric stretching). Primary aliphatic amines presences is indicated by peaks at 1466.52 (N-H bending) and 1221.34 cm\(^{-1}\) (C-N stretching).
Aromatic compounds presence is shown by peak at 1466.52 cm\(^{-1}\) (C-C stretching in ring). The observed bands for amines, amides, amino acids confirmed the presence of proteins, whereas presence of other bio-molecules like carboxylic acids, carbohydrates and oil was indicated by other absorption bands. Bhat et al. [24] and Zia-Ul-Haq et al. [25] have reported previously similar functional groups in *Gnetum gnemon* L. and *Pisum sativum* L. respectively.

Especial attention is being given to the identification of phenolic acids, flavonoids and tannins from extracts of legume seeds. Total phenolic content (TPC, mg GAE/g) of seed extracts from selected mash bean cultivars are presented in Table 7. The TPC was observed in highest amount in NARC-Mash-97 (86 mg GAE/g), whereas the lowest TPC was noted for NARC-Mash-1 (75 mg GAE/g). The total flavonoids contents (TFCs) and condensed tannins (CTCs) were expressed in catechin equivalents (CAE/g). The cultivars differed significantly (\(P<0.05\)) in TFCs and CTCs. The chromatograms (RP-HPLC) of extracts of seeds of mash bean were recorded at 330 nm and two dominant peaks (1–2) with a retention times of 28 and 28.8 min respectively (Figure 3), were observed.

| Minerals      | NARC-Mash-1       | NARC-Mash-2       | NARC-Mash-3       | NARC-Mash-97       |
|---------------|-------------------|-------------------|-------------------|-------------------|
| Phosphorus    | 461.24 ± 0.22\(^a\) | 480.47 ± 3.02\(^b\) | 500.15 ± 2.91\(^a\) | 440.90 ± 0.80\(^b\) |
| Potassium     | 1603.39 ± 1.66\(^b\) | 1638.88 ± 2.86\(^a\) | 1646.11 ± 3.17\(^a\) | 1600.03 ± 2.61\(^b\) |
| Sodium        | 227.01 ± 4.55\(^c\) | 244.90 ± 1.41\(^b\) | 284.08 ± 2.01\(^a\) | 261.33 ± 1.79\(^a\) |
| Calcium       | 462.90 ± 2.07\(^a\) | 375.01 ± 3.66\(^b\) | 485.38 ± 1.14\(^a\) | 394.19 ± 2.04\(^b\) |
| Magnesium     | 263.85 ± 3.56\(^a\) | 239.70 ± 1.36\(^b\) | 20845 ± 1.21\(^c\) | 221.77 ± 1.18\(^a\) |
| Iron          | 5.89 ± 0.25\(^b\) | 6.14 ± 0.21\(^b\) | 6.38 ± 0.18\(^c\) | 6.55 ± 0.33\(^a\) |
| Manganese     | 2.39 ± 2.07\(^b\) | 3.27 ± 0.05\(^c\) | 3.32 ± 0.11\(^a\) | 3.22 ± 0.18\(^a\) |
| Zinc          | 2.40 ± 0.14\(^a\) | 2.28 ± 0.12\(^a\) | 2.50 ± 0.22\(^a\) | 1.94 ± 0.76\(^b\) |
| Copper        | 3.92 ± 0.47\(^b\) | 4.03 ± 0.08\(^c\) | 4.26 ± 0.66\(^a\) | 4.51 ± 0.34\(^a\) |
| Na/K          | 0.14              | 0.15              | 0.17              | 0.16              |
| Ca/P          | 1.00              | 0.78              | 0.97              | 0.89              |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (\(p < 0.05\)).

### Table 3 Percentage composition of amino acids in seeds

| Amino acid   | NARC-Mash-1       | NARC-Mash-2       | NARC-Mash-3       | NARC-Mash-97       |
|--------------|-------------------|-------------------|-------------------|-------------------|
| Alanine      | 4.63 ± 0.17\(^b\) | 5.20 ± 0.07\(^a\) | 4.35 ± 0.05\(^a\) | 4.17 ± 0.21\(^c\) |
| Arginine     | 6.03 ± 0.27\(^c\) | 6.30 ± 0.04\(^b\) | 6.53 ± 0.03\(^a\) | 6.64 ± 0.15\(^a\) |
| Aspartic acid| 13.20 ± 0.27\(^a\) | 12.40 ± 0.08\(^b\) | 11.98 ± 0.07\(^c\) | 11.53 ± 0.11\(^c\) |
| Cystine      | 0.75 ± 0.29\(^b\) | 0.90 ± 0.04\(^a\) | 0.45 ± 0.03\(^c\) | 0.72 ± 0.23\(^b\) |
| Glutamic acid| 21.07 ± 0.65\(^a\) | 21.49 ± 0.07\(^a\) | 20.44 ± 0.09\(^b\) | 19.19 ± 0.62\(^c\) |
| Glycine      | 4.39 ± 0.12\(^a\) | 4.61 ± 0.05\(^a\) | 3.73 ± 0.03\(^b\) | 4.34 ± 0.24\(^a\) |
| Histidine    | 2.36 ± 0.31\(^b\) | 2.13 ± 0.02\(^b\) | 3.21 ± 0.01\(^d\) | 3.26 ± 0.26\(^a\) |
| Isoleucine   | 4.48 ± 0.17\(^a\) | 4.37 ± 0.07\(^a\) | 3.79 ± 0.05\(^b\) | 4.25 ± 0.09\(^a\) |
| Leucine      | 8.89 ± 0.12\(^a\) | 7.31 ± 0.03\(^b\) | 7.79 ± 0.04\(^b\) | 7.54 ± 0.45\(^b\) |
| Lysine       | 4.19 ± 0.88\(^d\) | 7.69 ± 0.01\(^a\) | 6.90 ± 0.08\(^b\) | 5.07 ± 0.74\(^c\) |
| Methionine   | 1.92 ± 0.74\(^a\) | 1.12 ± 0.05\(^b\) | 1.42 ± 0.09\(^b\) | 1.29 ± 0.20\(^b\) |
| Phenylalanine| 5.59 ± 0.18\(^d\) | 4.88 ± 0.06\(^b\) | 5.80 ± 0.07\(^a\) | 5.67 ± 0.12\(^a\) |
| Proline      | 4.30 ± 0.21\(^b\) | 3.69 ± 0.03\(^c\) | 5.01 ± 0.01\(^a\) | 4.20 ± 0.08\(^b\) |
| Serine       | 5.18 ± 0.30\(^a\) | 5.31 ± 0.05\(^a\) | 4.14 ± 0.08\(^c\) | 4.78 ± 0.07\(^b\) |
| Threonine    | 3.95 ± 0.35\(^b\) | 3.80 ± 0.04\(^b\) | 4.50 ± 0.03\(^a\) | 3.99 ± 0.28\(^b\) |
| Tryptophan   | 1.01 ± 0.14\(^d\) | 1.70 ± 0.09\(^b\) | 2.80 ± 0.02\(^a\) | 3.15 ± 0.28\(^a\) |
| Valine       | 5.09 ± 0.11\(^a\) | 4.80 ± 0.08\(^b\) | 4.94 ± 0.04\(^d\) | 5.08 ± 0.04\(^a\) |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (\(p < 0.05\)).
The spectra (UV) of both compounds (peaks 1–2) displayed maxima at 269 and 334 nm. Compounds 1 and 2 were identified as chlorogenic acid and caffeic acids when compared with standards run simultaneously. The mash bean extracts investigated in this study were characterized by several times higher content of flavonoids and condensed tannins when compared to desi chickpea, kabuli chickpea, lentil, cowpea, *Albizia lebbeck* and *Acacia leucophloea* varieties [11-13,25,26]. Presence of higher contents of various phenolic compounds was noted in extracts (Table 8). Various phenolic acids have been identified earlier in extracts from other legumes like chickpea, cowpea and pea [11,25,26]. Consumption of phenolic-rich foods is associated with low risk of several chronic diseases such as cardiovascular disease, ageing, cancer, neurodegenerative disease and Alzheimer disease as is evident from various epidemiological studies which highlights importance of presence of ample contents of phenolic acids noted in mash bean extract.

The human body has several mechanisms to shield bio-molecules against damage caused by reactive oxygen and nitrogen species. However, the instinctive protection may not be adequate to counter the rigorous or continuous oxidative stress. Hence, certain amounts of exogenous antioxidants are frequently required to maintain sufficient antioxidants level to balance the reactive nitrogen and oxygen species-pressure in the human body. Scientists are exploring antioxidants from natural sources like legume seeds as these are natural, cost effective and without side effects. The scavenging activity of mash bean extracts was expressed by antiradical assays against DPPH$^-$ and ABTS$^{•+}$ assay as well as by FRAP and reducing power assays as shown in Table 9. DPPH values of mash bean varieties ranged from 34.72 in NARC-Mash-2 to 39.49 μmol Trolox/g in NARC-Mash-3. Sufficient scavenging of DPPH radical was observed by extracts. It indicates that antioxidants present in extracts quench free radicals by donating them hydrogen atoms thereby converting them to non-toxic species. Although assessment of antiradical activity of an extract by DPPH protocol is fast and trouble-free, it usually has a relatively small linear reaction range therefore antiradical activity against ABTS$^{•+}$ was measured. The ABTS$^{•+}$ scavenging data indicated that the extracts may scavenge free radicals by hydrogen/electron donation mechanism and may protect biomatrices from oxidative degradation resulting from free radicals. Substantial antiradical activity for DPPH and ABTS$^{•+}$ was observed with same order of scavenging in both protocols. It was noted

![Figure 2](image-url)

**Figure 2** In-vitro protein and starch digestibility (%) of mash bean seeds.

The spectra (UV) of both compounds (peaks 1–2) displayed maxima at 269 and 334 nm. Compounds 1 and 2 were identified as chlorogenic acid and caffeic acids when compared with standards run simultaneously. The mash bean extracts investigated in this study were characterized by several times higher content of flavonoids and condensed tannins when compared to desi chickpea, kabuli chickpea, lentil, cowpea, *Albizia lebbeck* and *Acacia leucophloea* varieties [11-13,25,26]. Presence of higher contents of various phenolic compounds was noted in extracts (Table 8). Various phenolic acids have been identified earlier in extracts from other legumes like chickpea, cowpea and pea [11,25,26]. Consumption of phenolic-rich foods is associated with low risk of several chronic diseases such as cardiovascular disease, ageing, cancer, neurodegenerative disease and Alzheimer disease as is evident from various epidemiological studies which highlights importance of presence of ample contents of phenolic acids noted in mash bean extract.

The human body has several mechanisms to shield bio-molecules against damage caused by reactive oxygen and nitrogen species. However, the instinctive protection may not be adequate to counter the rigorous or continuous oxidative stress. Hence, certain amounts of exogenous antioxidants are frequently required to maintain sufficient antioxidants level to balance the reactive nitrogen and oxygen species-pressure in the human body. Scientists are exploring antioxidants from natural sources like legume seeds as these are natural, cost effective and without side effects. The scavenging activity of mash bean extracts was expressed by antiradical assays against DPPH$^-$ and ABTS$^{•+}$ assay as well as by FRAP and reducing power assays as shown in Table 9. DPPH values of mash bean varieties ranged from 34.72 in NARC-Mash-2 to 39.49 μmol Trolox/g in NARC-Mash-3. Sufficient scavenging of DPPH radical was observed by extracts. It indicates that antioxidants present in extracts quench free radicals by donating them hydrogen atoms thereby converting them to non-toxic species. Although assessment of antiradical activity of an extract by DPPH protocol is fast and trouble-free, it usually has a relatively small linear reaction range therefore antiradical activity against ABTS$^{•+}$ was measured. The ABTS$^{•+}$ scavenging data indicated that the extracts may scavenge free radicals by hydrogen/electron donation mechanism and may protect biomatrices from oxidative degradation resulting from free radicals. Substantial antiradical activity for DPPH and ABTS$^{•+}$ was observed with same order of scavenging in both protocols. It was noted

### Table 4 Fatty acid composition (%) of oil of mash bean seeds

| Fatty acid         | NARC-Mash-1         | NARC-Mash-2         | NARC-Mash-3         | NARC-Mash-97         |
|--------------------|---------------------|---------------------|---------------------|---------------------|
| Palmitic acid      | 11.31 ± 2.20³       | 10.99 ± 1.99³       | 11.23 ± 1.87³       | 12.09 ± 1.58³       |
| Stearic acid       | 2.09 ± 0.63³        | 2.70 ± 0.24³        | 2.89 ± 0.43³        | 2.17 ± 0.77³        |
| Behenic acid       | 0.99 ± 0.14³        | 1.00 ± 0.29³        | 0.87 ± 0.22³        | 0.93 ± 0.30³        |
| Oleic acid         | 26.62 ± 0.07³       | 26.74 ± 0.15³       | 27.34 ± 0.25³       | 26.65 ± 0.35³       |
| Linoleic acid      | 07.19 ± 4.47³       | 08.93 ± 5.11³       | 07.08 ± 3.74³       | 08.64 ± 3.87³       |
| α-Linolenic acid   | 51.80 ± 0.03³       | 49.64 ± 0.06³       | 50.59 ± 0.05³       | 49.52 ± 0.09³       |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).
that reducing potential of extracts increased with increasing amount of extracts. Butylated hydroxanisole was used as standard to compare the reducing power of extracts. Mechanistic studies indicate that antioxidant potential of extracts is closely linked with their reducing power. The results were close to reported earlier [27-29].

The FRAP assay determines antioxidant activity of extracts as their potential to reduce ferric ions to ferrous ions. The FRAP values of the extracts of seeds of selected cultivars are presented in Table 9. Similar to that in DPPH analyses, high variations of FRAP values were observed and FRAP values of cultivars ranged from 9.65 mmol Fe²⁺/g in NARC-Mash-3 to 13.76 mmol Fe²⁺/g in NARC-Mash-97. Our results for FRAP are different from those reported earlier [27-29]. Antioxidant activity of the extracts of seeds of other plant like pea, cowpea, lentil, garden cress, capper and chickpea has been reported in several studies [25,26,30-32] by our research group.

It is generally believed that diabetes can be cured with more consumption of legumes however the mechanism behind this remained unexplored till now. The recent studies indicated that legumes cure diabetes by reducing AGE-formation. Advanced glycation end products (AGE) formation is increased in diabetes mellitus, so search for (AGEs)-inhibitor is a new approach in diabetes treatment. Two models used mostly for quantification of AGE-inhibition of plant extracts are BSA-MGO and BSA-glucose models. In advanced glycation end (AGE) products inhibition activity, NARC-Mash-97 exhibited the highest inhibition (86.67%), followed by NARC-Mash-3 (74.84%) in BSA-glucose method. BSA-MGO inhibition model showed the same trend like that of BSA-glucose model (Figure 4). It is believed that phenolic compounds present in legume seeds inhibit the AGE-formation by inhibiting production of free radical during glycation process and subsequently inhibiting protein modification. The results (Figure 4) obtained in our study are in agreement with those reported previously for other legume seeds [33-38]. Same trend was observed in tyrosinase inhibition activity as was for AGE inhibition. Tyrosinase inhibition potential of extracts of seeds of mash bean may be ascribed to the presence of phenolic contents since hydroxyl groups present in various phenolic acids make a hydrogen bond at active site of the tyrosinase and as a result tyrosinase activity is decreased or stopped. Tyrosinase inhibitors have potential applications in food and cosmetic industry because they are used to stop or slow-down browning of various food commodities like fruits, vegetable and fisheries products and impart whitening effects to skin by stopping human skin hyper-pigmentation. The browning of food commodities leads to decrease in attractive appearance and loss of nutritional quality. It rationalized traditional use of mashbean in facial massages by indigenous communities and proves its anti-freckles, anti-wrinkling, anti-ageing and skin-whitening activity. For the first time tyrosinase inhibition activities of extracts of seeds of mash beans are being reported.

Statistically non-significant and very low correlations were found between different parameters viz. FRAP with BSA-G, BSA-MGO and TI; FRAP with BSA-MGO and TI at P < 0.05 (Table 10). TPC was having statistically no correlation with FRAP. Similarly, there was very low correlation between ABST and BSA-G. Medium correlations were found between different parameters viz. FRAP with BSA-G, BSA-MGO and TPC, which is statistically significant. It is generally believed that diabetes can be cured with more consumption of legumes however the mechanism behind this remained unexplored till now. The recent studies indicated that legumes cure diabetes by reducing AGE-formation. Advanced glycation end products (AGE) formation is increased in diabetes mellitus, so search for (AGEs)-inhibitor is a new approach in diabetes treatment. Two models used mostly for quantification of AGE-inhibition of plant extracts are BSA-MGO and BSA-glucose models. In advanced glycation end (AGE) products inhibition activity, NARC-Mash-97 exhibited the highest inhibition (86.67%), followed by NARC-Mash-3 (74.84%) in BSA-glucose method. BSA-MGO inhibition model showed the same trend like that of BSA-glucose model (Figure 4). It is believed that phenolic compounds present in legume seeds inhibit the AGE-formation by inhibiting production of free radical during glycation process and subsequently inhibiting protein modification. The results (Figure 4) obtained in our study are in agreement with those reported previously for other legume seeds [33-38]. Same trend was observed in tyrosinase inhibition activity as was for AGE inhibition. Tyrosinase inhibition potential of extracts of seeds of mash bean may be ascribed to the presence of phenolic contents since hydroxyl groups present in various phenolic acids make a hydrogen bond at active site of the tyrosinase and as a result tyrosinase activity is decreased or stopped. Tyrosinase inhibitors have potential applications in food and cosmetic industry because they are used to stop or slow-down browning of various food commodities like fruits, vegetable and fisheries products and impart whitening effects to skin by stopping human skin hyper-pigmentation. The browning of food commodities leads to decrease in attractive appearance and loss of nutritional quality. It rationalized traditional use of mashbean in facial massages by indigenous communities and proves its anti-freckles, anti-wrinkling, anti-ageing and skin-whitening activity. For the first time tyrosinase inhibition activities of extracts of seeds of mash beans are being reported.

| Table 5 Tocopherol content (mg/100 g) in oil of seeds of mash bean cultivars |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| α-Tocopherol               | α-Tocopherol               | α-Tocopherol               | α-Tocopherol               |
| NARC-Mash-1                | NARC-Mash-2                | NARC-Mash-3                | NARC-Mash-97               |
| 3.04 ± 0.89b               | 2.97 ± 0.55b               | 3.49 ± 0.17b               | 3.17 ± 0.34b               |
| γ-Tocopherol               | γ-Tocopherol               | γ-Tocopherol               | γ-Tocopherol               |
| 722.09 ± 2.17b             | 724.34 ± 4.13b             | 722.21 ± 1.16a             | 720.33 ± 2.01a             |
| δ-Tocopherol               | δ-Tocopherol               | δ-Tocopherol               | δ-Tocopherol               |
| 16.69 ± 3.3b               | 16.18 ± 4.2c               | 17.14 ± 2.66a              | 17.12 ± 4.0a               |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).

| Table 6 Sterol content (mg/100 g) in oil of seeds of mash bean cultivars |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| Sterols                     | Sterols                     | Sterols                     | Sterols                     |
| β-Sitosterol                | β-Sitosterol                | β-Sitosterol                | β-Sitosterol                |
| NARC-Mash-1                 | NARC-Mash-2                 | NARC-Mash-3                 | NARC-Mash-97                |
| 56.5 ± 0.2a                 | 55.1 ± 0.6a                 | 55.1 ± 0.61a                | 56.1 ± 0.51a                |
| Stigmastanol                | Stigmastanol                | Stigmastanol                | Stigmastanol                |
| 34.0 ± 0.4a                 | 34.4 ± 0.8a                 | 34.4 ± 0.8a                 | 33.4 ± 0.1b                 |
| Δ⁵-Vanasterol               | Δ⁵-Vanasterol               | Δ⁵-Vanasterol               | Δ⁵-Vanasterol               |
| 4.00 ± 0.22b                | 3.64 ± 0.38b                | 3.64 ± 0.38b                | 4.51 ± 0.12b                |
| Stigmastanol                | Stigmastanol                | Stigmastanol                | Stigmastanol                |
| 2.66 ± 0.05b                | 3.79 ± 0.12a                | 3.79 ± 0.12a                | 3.21 ± 0.38b                |
| Δ⁵-avenasterol              | Δ⁵-avenasterol              | Δ⁵-avenasterol              | Δ⁵-avenasterol              |
| 1.01 ± 0.17a                | 1.05 ± 0.69a                | 1.05 ± 0.69a                | 1.09 ± 0.43a                |
| Campesterol                 | Campesterol                 | Campesterol                 | Campesterol                 |
| 0.87 ± 0.40b                | 0.98 ± 0.80a                | 0.98 ± 0.80a                | 0.66 ± 0.18c                |
| Unidentified                | Unidentified                | Unidentified                | Unidentified                |
| 1.00 ± 0.03a                | 1.00 ± 0.27a                | 1.00 ± 0.27a                | 1.00 ± 0.13a                |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).
Table 7 Total phenolic contents, total flavonoid contents and condensed tannin contents in extracts of seeds of mash bean cultivars

| Cultivar       | Total phenolic contents | Total flavonoid contents | Condensed tannin contents |
|----------------|-------------------------|--------------------------|---------------------------|
| NARC-Mash-1    | 75.91 ± 2.72<sup>c</sup> | 51.78 ± 1.85<sup>b</sup> | 86.79 ± 1.56<sup>a</sup>  |
| NARC-Mash-2    | 79.33 ± 1.52<sup>b</sup> | 47.11 ± 2.47<sup>c</sup> | 89.14 ± 1.11<sup>b</sup>  |
| NARC-Mash-3    | 82.22 ± 1.36<sup>a</sup> | 42.66 ± 1.81<sup>d</sup> | 93.68 ± 1.65<sup>a</sup>  |
| NARC-Mash-97   | 86.99 ± 1.19<sup>a</sup> | 55.73 ± 1.92<sup>a</sup> | 79.20 ± 1.77<sup>c</sup>  |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).

Figure 3 HPLC spectra of mash bean (NARC-Mash-97) seed extract.
Very high and positive correlations were found among certain variables like BSA-G and BSA-MGO, BSA-G and TI, BSA-MGO and TI, TPC and BSA-G, TPC and BSA-MGO and TPC and TI. The correlation between DPPH and FRAP was also statistically significant at p < 0.05, DPPH and ABST at p < 0.01, FRAP and ABST were also highly correlated (p < 0.01).

Conclusion
The results suggested mash bean seeds as a rich source of nutrients and extracts of seeds exhibited good anti-oxidant and biological activities. Seeds are rich source of protein and carbohydrate and good source of dietry fibre. These also contain ample amount of essential minerals like Ca, K, Na, Mg, Cu and Zn and various essential and non-essential amino acids. Seeds also have acceptable fatty acids, tocopherol and sterol profile. Various functional groups were detected in FTIR of seeds and extracts. Antioxidant results suggested them as rich source of phenolic acids, flavonoids and condensed tannin contents. The extracts indicated good tyrosinase and AGE-inhibition activity. These results suggest that mash bean seed may be used in food industry as functional food and nutraceutical as well as in cosmetic and pharmaceutical industry as ingredient of skin-whitening creams and as cure for diabetes respectively. The data obtained will be helpful for labeling of nutrients as well as for monitoring the quality and authenticity of foods containing mash bean in indigenous markets. Further investigations are necessary to evaluate the toxic effects (if any), to determine the ant-nutrients factors present and to understand mechanism of action of tyrosinase-inhibitory and AGE-inhibitory potential of extracts.

Methods
Material
Analytical grade solvents were used. All chemicals were purchased from Sigma except where indicated. The seeds of four mash bean cultivars namely, NARC-Mash-1, NARC-Mash-2, NARC-Mash-3 and NARC-Mash-97 were procured from National Agricultural Research Centre, Islamabad (Pakistan). Seeds of four cultivars were stored in stainless-steel containers at 4°C prior to analysis.

Proximate analysis
Proximate chemical analysis of seeds was carried out according to AOAC International methods as per our previous studies [39]. Results are shown in Table 1.

Vitamin contents
Powdered sample (5 g) was steamed with concentrated H₂SO₄ (30 ml) for half an hour. After cooling, distilled H₂O was added to this suspension to make its volume up to 50 ml and filtered. Basic lead acetate (60%, 5 ml) was added to this filtrate (25 ml). The pH was adjusted (9.5) and supernatant was collected after centrifugation. To this supernatant, concentrated H₂SO₄ (2 ml) was added. After 1 hr, this mixture was centrifuged and then ZnSO₄ (5 ml, 40%) was added. The pH was adjusted (8.4) and supernat was collected after centrifugation. The pH of resulting supernatant was adjusted (7) and this was utilized as niacin extract. One ml of this extract was made 6 ml by distilled H₂O; after addition of cyano-gren bromide (3 ml) and shaking, aniline (4%, 1 ml) was added. After 5 min, yellow color formed was spectrophotometrically measured at 420 nm against blank and niacin contents were calculated by a standard graph [16]. Thiochrome method and fluorescence method were used to determine thiamine, riboflavin, nicotinic acid and pyridoxine respectively.

Table 8 Content of two main phenolic compounds in the extracts and seeds of mash bean cultivars

| Cultivar     | Compound 1 (mg/g extract) | Compound 2 (mg/g extract) | Compound 1 (mg/g fresh seeds) | Compound 2 (mg/g fresh seeds) |
|--------------|---------------------------|---------------------------|-------------------------------|-------------------------------|
| NARC-Mash-1  | 4.09 ± 0.20                  | 5.22 ± 0.26                | 0.39 ± 0.02                  | 0.50 ± 0.02                  |
| NARC-Mash-2  | 5.94 ± 0.30                  | 8.09 ± 0.40                | 0.55 ± 0.03                  | 0.74 ± 0.04                  |
| NARC-Mash-3  | 5.48 ± 0.27                  | 7.35 ± 0.37                | 0.48 ± 0.02                  | 0.65 ± 0.03                  |
| NARC-Mash-97 | 5.07 ± 0.25                  | 6.94 ± 0.35                | 0.44 ± 0.02                  | 0.44 ± 0.02                  |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).

Table 9 Antioxidant capacity of extracts of seeds of mash bean cultivars

| Cultivar     | Reducing power (mg/g) | DPPH· scavenging capacity (µmol Trolox/g) | FRAP (mmol Fe²⁺/g) | ABTS scavenging capacity (µmol trolox/g) |
|--------------|-----------------------|------------------------------------------|--------------------|------------------------------------------|
| NARC-Mash-1  | 1.09 ± 0.18           | 41.64 ± 0.18                            | 12.81 ± 0.03       | 33.81 ± 0.45                            |
| NARC-Mash-2  | 0.87 ± 0.02           | 34.72 ± 0.29                            | 11.70 ± 0.19       | 27.09 ± 0.58                            |
| NARC-Mash-3  | 1.02 ± 0.09           | 39.49 ± 0.11                            | 9.65 ± 0.37        | 29.74 ± 0.83                            |
| NARC-Mash-97 | 0.95 ± 0.06           | 46.56 ± 0.05                            | 13.76 ± 0.57       | 35.93 ± 0.22                            |

Values in the same row having different letters differ significantly with least significant difference (LSD) at probability (p < 0.05).
used for determination of thiamine and riboflavin contents respectively [40,41] Figure 1.

Minerals contents
A muffle furnace was used to incinerate seeds (450°C; 12 h) and the resulting samples were digested by acid mixture (nitric/perchloric; 2:1). Na and K were estimated by taking aliquots from this digested material by flame photometer. Other minerals like Mn, Mg, Ca, Fe, Cu and Zn were estimated spectrophotometrically (AAS; Perkin-Elmer 5000) while phosphovanado-molybdate method was used to measure phosphorus contents. Standard solutions of known concentration were run concurrently to quantify the samples [31,32] (Table 2).

Amino acid analysis
HCl (6 M) was used to hydrolyze samples (300 mg) in an evacuated test tube (105°C; 24 h). Citrate buffer (pH 2.2) was used to dissolve the dried residue resulting from flash evaporation. Hitachi Perkin-Elmer (KLA 3B) amino acid analyzer was utilized to quantify amino acids by taking aliquots from above solution. After treatment with performic acid followed by hydrolysis (HCl), cystine and methionine were analyzed separately from same solution. Alkali hydrolysis (NaOH) method was used to measure tryptophan [22,23] (Table 3).

Protein and starch digestibility (In-vitro)
In-vitro digestibility of protein was evaluated enzymetically while starch digestibility was evaluated as starch hydrolyzed (%) out of total starch present in sample [42-44] (Figure 2).

Fatty acid (FA) composition
Petroleum ether as solvent was used to extract oil from seeds by Soxhlet apparatus (6 hr) as per official AOCS method [39]. The fatty acid profile of oils obtained was evaluated by a method reported earlier [45]. Briefly, n-heptane (1 mL) was used to dissolve oil (1 drop), sodium methanolate (50 μL; 2 M) was added, and shaken in a closed tube (1 min). Water (100 μL) was added and the tube was centrifuged (4500 g; 10 min) and resulting aqueous phase was separated. To remaining heptane phase, HCl (50 μL; 1 M) was added, both phases were mixed for short period of time and resulting aqueous phase was
discarded. After addition of sodium hydrogen sulphate (20 mg) and centrifugation (4500 g; 10 min), n-heptane phase was stored in a vial and inserted in a gas chromato-

**Tocopherol contents**

Twenty five ml of n-heptane was mixed with oil (250 mg) and tocopherol contents were was analyzed by HPLC system (Merck-Hitachi), containing a pump (L-6000), a fluorescence spectrophotometer (Merck-Hitachi F-1000), excitation wavelength (295); emission wavelength (330 nm) and a D-2500 integration system; 20 μL of samples were inserted by a Merck 655-A40 autosampler in a dual phase HP (Merck) having column column (25 cm × 4.6 mm) while flow rate was adjusted at 1.3 mL/min. Mobile phase used was n-heptane: tert-butyl methyl ether (99:1) [46] (Table 5).

**Sterol composition**

The sterols were quantified by a gas chromatograph (Perkin Elmer model 8700), having flame-ionization detector (FID) and OV-17 capillary column (methyl phenyl polysiloxane coated; ID: 30 m × 2.25 mm, film: 20 μm). The column was operated isothermally (255°C) while temperature for injector and detector were 275 and 290°C, respectively. Carrier gas selected was extra pure nitrogen with 3 mL/min as flow rate. Sterols were recognized and quantified by comparing with a sterol standard mixture [21,31] (Table 6).

**Extraction**

The mash bean seeds were ground to flour by a mill (IKA Works Inc.) and were sieved (60-mesh). After maceration with 5 L solvent mixture of aqueous: methanol (80:20) for 15 days at room temperature and extracts were collected. The process was carried out three times. The resulting extracts were collected and filtered by filter paper. The extra solvent present was evaporated under reduced pressure by using a rotary evaporator. A thick gummy mass was obtained which was then dried in a desicator and utilized for assessment of biological activities.

**FTIR of Mash bean powder and crude extract**

Functional groups present in flour and extracts of seeds of mash bean cultivars were identified by FTIR spectroscopy (Perkin Elmer; UK) [24,25] Figure 5 and 6.

**Total phenolic, flavonoid and condensed tannin contents (TPC, TFC, CTC)**

Total phenolics were were estimated using the Folin and Ciocalteau’s phenol reagent [47] and results were reported as gallic acid equivalents [48,49]. A previously reported method was used for estimation of flavonoids contents [50]. Condensed tannins were quantified by acidified vanillin reagent [51] and results were expressed as mg of CAE/g (Table 7).

**DPPH radical scavenging assay**

Scavenging potential of extracts of mash bean seeds against DPPH• was estimated by a previously reported method [53]. The absorbance of extracts (A\text{sample}) was measured spectrophotometrically (Shimadzu, Kyoto, Japan) at 517 nm and ethanol was used as blank. The extraction solvent (0.2 mL) after addition of DPPH• was used as negative control (A\text{control}). Following equation was used to assess antiradical activity:

\[
\text{Antiradical activity} \% = \left(1 - \frac{\text{Absorbance}_{\text{sample}}}{\text{Absorbance}_{\text{control}}} \right) \times 100
\]

Calibration curve of Trolox was used to calculate results and indicated as micromoles of Trolox equivalent (μmol Trolox/g) Table 9.

**Ferric reducing antioxidant power (FRAP) activity**

FRAP assay was carried out to assess antioxidant activity [54]. Deionized water was used to dilute properly the sample solution to fit within the linearity range of Fe²⁺. The calibration curve of Fe²⁺ was used to calculate FRAP value as mmoles of Fe²⁺ equivalent (mmol Fe²⁺/g) Table 9.

**Reducing power**

Reducing potential of investigated extracts was determined by a reported method [55]. Aliquots (2.5 ml) of extracts dissolved in phosphate buffer (pH 6.6, 0.2 M) were mixed with C₆H₉FeK₃ (10 mg/ml; 2.5 ml) and resulting solution was incubated (20 min; 50°C). To this reaction mixture, trichloroacetic acid (100 mg/ml solution;
(2.5 ml) was added and centrifuged (1000 rpm; 10 min). The resulting supernatant (2.5 ml) was mixed with an equal volume of H2O (distilled) and FeCl3 (1 mg/ml solution; 0.5 ml) was added. Spectrophotometer was used to measure absorbance at 700 nm against ascorbic acid Table 9.

ABTS** scavenging assay
Scavenging activity of extracts of seeds was also evaluated against ABTS** [56]. ABTS aqueous solution (5 mM) was passed from the oxidizing reagent (MnO2), on filter paper (Fisher Brand P8) to prepare ABTS**. The solution was filtered from fisher membrane (0.2 mm) to remove extra MnO2. Phosphate buffered saline (5 mM; pH 7.4) was used to dilute extracts to an absorbance of approximately 0.700 (±0.020) at 734 nm. The extracts (1.0 mL) were added to ABTS** solution (5 mL), and the absorbance was measured after 10 min. The blank used was PBS Table 9.

Evaluation of AGE inhibition activity
Inhibitory potential of mash bean extracts on the formation of advanced glycation end (AGE) products was determined by BSA-MGO and BSA-glucose models (Table 10). Briefly, BSA (5 g) and D-glucose (14.4 g) were dissolved in phosphate buffer (1.5 M; pH 7.4) to get a control solution
containing D-glucose (0.8 M) and BSA (50 mg/mL). Two mL of this solution was incubated at 37°C (1 week) in the absence or presence of bean extracts (1 mL) in phosphate buffer. After one week, fluorescent intensity (excitation: 330 nm; emission: 410 nm) was measured. The BSA-MGO assay was performed as reported elsewhere. Briefly, MGO (31 µL) was mixed with BSA (40 mg) in phosphate buffer (pH 7.4; 0.1 M) to make a control solution of MGO (5 mM) and BSA (1 mg/mL). Two mL of control solution was incubated (6 days) with or without bean extracts (1 mL) in phosphate buffer [48-52]. Inhibition (%) of formation of AGE by AGE for both models was calculated using the following equation:

\[
\text{Percent Inhibition} = 1 - \frac{\text{fluorescence with extract}}{\text{fluorescence without extract}} \times 100
\]

Measurement of tyrosinase inhibition activity
Microtiter plates (96-well) were used to perform assays and absorbance was measured (475 nm) by a plate reader. Each well contained sample (40 µL) and phosphate buffer (pH 6.8, 80 µL 0.1 M), tyrosinase (31 units/mL, 40 µL) and L-DOPA (2.5 mM; 40 µL), the samples were incubated (37°C) for half an hour and results are shown in Table 10. A control was prepared having all ingredients except tyrosinase [48-52]. The tyrosinase inhibition percentage was calculated as follows:

\[
\text{Percent Inhibition} = \frac{A_{\text{control}} - A_{\text{sample}}}{A_{\text{control}}} \times 100
\]

Statistical analysis
All experiments were performed in triplicate and values marked by same letter in same column are not significantly different (P < 0.05). Data are expressed as the mean ± standard deviation. Data were analyzed by using the “MSTATC” statistical computer package [57].

Competing interests
The authors state that there are no competing interests.

Authors’ contributions
MZUH and RA contributed to the experiment. SA and SAB contributed in statistical interpretation of the data while SE and HZJ helped in preparation of the manuscript. All authors approved the final form of the manuscript.

Author details
1The Patent Office, Karachi, Pakistan. 2Department of Agronomy, Bahauddin Zakariya University, Multan 60800, Pakistan. 3Department of Applied Chemistry and Biochemistry, Government College University, Faisalabad, Pakistan. 4Institute of Animal Reproduction and Food Research of the Polish Academy of Sciences, Tuwima Str. 10, 70-473 Częstochowa, Poland. 5Agricultural Faculty, Department of Horticulture, Ataturk University, Erzurum, Turkey. 6Department of Crop Science, Faculty of Agriculture, 43400 UPM Serdang, Selangor, Malaysia.

Received: 31 March 2014 Accepted: 22 May 2014
Published: 30 May 2014

References

1. Bhattacharya S, Lahra RB, Bhat KR: Controlled stress rheological measurement of black gram flour dispersions. J Food Eng 2006, 63:135-139.
2. Anitha K, Ranjith K, Vakula K, Thirupathi G, Balaji B: Protective effect of Vigna mungo (L) against carbon tetrachloride induced hepatotoxicity. Int J Pharm Res 2012, 22:9-24.
3. Battu G, Anjana CR, SK, Priya TH, Malleswari VN, Reeshma S: A phytopharmacological review on Vigna species. Pharmacnet 2011, 2,67-69.
4. Nitin M, Ifthehar S, Mumtaz M: Hepatoprotective activity of methanolic extract of Vigna mungo (Linn). Hepper in ethanol-induced hepatotoxicity in rats. Ayush J Pharm Sci 2012, 2:62-67.
5. Zia-Ul-Haq M, Ahmad S, Rezwan GH, Qayum M, Ahmed S, Harif M: Platelet aggregation inhibition activity of selected flora of Pakistan. Pak J Pharm Sci 2012, 25:863-865.
6. Zia-Ul-Haq M, Landa P, Kutl Z, Qayum M, Ahmad S: Evaluation of anti-inflammatory activity of selected legumes from Pakistan: in vitro inhibition of cyclooxygenase-2. Pak J Pharm Sci 2013, 26:185-187.
7. Ahmed Z, Ansar M, Saleem A, Arif ZU, Zaved H, Saleem R: Improvement of mash bean production under rainfed conditions by rhizobium inoculation and low rates of starter nitrogen. Pak J Agric Res 2012, 25:154-160.
8. Achalas MM, Taran SA: Effect of seed rate on growth, yield components and yield of mash bean grown under irrigated conditions of arid uplands of Balochistan, Pakistan. Pak J Bot 2011, 43:936-939.
9. Zia-Ul-Haq M, Iqbal S, Ahmad M: Characteristics of oil from seeds of 4 mungbean (Vigna radiata (L) wilczek) cultivars grown in Pakistan. J Am Oil Chem Soc 2008, 85:851-856.
10. Zia-Ul-Haq M, Ahmad S, Chavara E, Ahmad S: Studies of oil from cowpea (Vigna unguiculata (L) Walp.) cultivars commonly consumed in Pakistan. Pak J Bot 2010, 43:1333-1341.
11. Zia-Ul-Haq M, Ahmad S, Ahmad M, Iqbal S, Khawar KM: Phytochemical, cooking and nutritional characteristics of some cowpea (Vigna unguiculata(L) Walp.) cultivars commonly consumed in Pakistan. Molecules 2013, 18:2005-2017.
12. Zia-Ul-Haq M, Cavar S, Qayum M, Khan I, Ahmad S: Compositional studies and antioxidant potential of Acacia leucophloea Roxb. Acta Bot Croat 2013, 72:27-31.
13. Zia-Ul-Haq M, Ahmad S, Qayum M, Erciçli S: Compositional studies and antioxidant potential of Albizia lebbeck (L) Betn. Turk J Bio J 2013, 37:25-32.
14. Khalil H, Intikhab J: Characterization and biochemical studies of the oils extracted from four cultivars of Vigna mungo grown in Pakistan. J Rashid Latif Med Coll 2013, 1:19-24.
15. Singh N, Kaur M, Sandhu KS, Sodhi NS: Physicochemical, cooking and textural characteristics of some Indian black gram (Phaseolus mungo L) varieties. J Sci Food Agric 2006, 86:977-982.
16. Zia-Ul-Haq M, Ahmad S, Amarnovicz R, Erciçli S: Compositional studies of some pea (Pisum sativum L) seed cultivars commonly consumed in Pakistan. Ital J Food Sci 2013, 25:305-302.
17. Soro TP, Kala KB, Mohan VR, Vedvial V: The biochemical composition and nutritional potential of three varieties of Vigna mungo (L) Hepper. Adv Biotech 2010, 15:6-16.
18. Miyamoto Y, Kajikawa A, Zaidi JH, Nakashita K, Sakamoto K: Minor and trace element determination of food spices and pulses of different origins by NAA and PAA. J Radioanal Nucl Chem 2000, 243:747-765.
19. NRC/NAS B: Recommended Dietary Allowances. 10th edition. Washington DC, USA: National Academy Press; 1989.
20. Jarrad A, Lubra B, Hamid Y: Studies on Vigna mungo L, effect of processing on carbohydrate fractionation and influence of grain starch on protein utilization in albino rats. Pak J Biol Sci 1999, 2:1298-1262.
21. Zia-Ul-Haq M, Ahmad S, Ahmad M, Iqbal S, Khawar KH: Effects of cultiver and row spacing on tocopherol and sterol composition of chickpea (Cicer arrietium L) seed oil. Tarim Bilimleri Dergisi 2009, 15:25-30.
22. Gopala RAG, Prabhakar JN, Aitzetmuller K: Tocopherol and fatty acid composition of some Indian pulses. J Am Oil Chem Soc 1997, 74:1603-1606.
23. Alisha T, Nishimura Y, Nakamura N, Roy K, Gosh P, Thakur S, Tamura T: Sterols of Cajanus cajan and three other Leguminosae seeds. Phytochem 1992, 31:1765-1768.
24. Bhat R, Yahya N: Evaluating belinjau (Gnetum gnemon L.) seed flour quality as a base for development of novel food products and food formulations. Food Chem 2014, 156:42-49.
25. Zia-Ul-Haq M, Amarowicz R, Ahmad S, Riaz M: Antioxidant potential of some pea (Pisum sativum L.) cultivars commonly consumed in Pakistan. Oix Commun 2013, 36:1046–1057.

26. Imran I, Zia-Ul-Haq M, Calani L, Mazzeo T, Pellegrini N: Phenolic profile and antioxidant potential of selected plants of Pakistan. J Appl Biol Food Qual 2014, 87:30–35.

27. Girish TK, Pratape VM, Rao UJSP: Protection of DNA and erythrocytes from free radical induced oxidative damage by black gram (Vigna mungo L.) husk extract. Food Chem Toxic 2012, 50:1690–1696.

28. Girish TK, Pratape VM, Rao UJSP: Nutrient distribution, phenolic acid composition, antioxidant and alpha-glucosidase inhibitory potentials of black gram (Vigna mungo L.) and its milled by-products. Food Res Int 2012, 46:570–377.

29. Marathe SA, Rajalakshmi V, Jamdar SN, Sharma A: Comparative study on antioxidant activity of different varieties of commonly consumed legumes in India. Food Chem Toxic 2011, 49:2005–2012.

30. Zia-Ul-Haq M, Amarowicz R, Ahmad S, Qayum M, Ercişli S: Antioxidant potential of mungbean cultivars commonly consumed in Pakistan. Oix Commun 2013, 36:15–25.

31. Zia-Ul-Haq M, Ahmad S, Calani L, Mazzeo T, Rio DD, Pellegrini N, DeFeo V: Compositional study and antioxidant potential of (Ipomoea hederacea Jacq. and Lepidium sativum L.) seeds. Molecules 2012, 17:10306–10321.

32. Zia-Ul-Haq M, Čavara S, Qayum M, Imran I, DeFeo V: Compositional studies, antioxidant and anti-diabetic activities of Capparis decidua (Forsk.) Edgew. Int J Mol Sci 2011, 12:8896–8861.

33. Tiwari AK, Swapna M, Ayesha SB, Zehra A, Agarwale SR, Madhusudana K: Identification of proglycemic and anti-hyperglycemic activity in antioxidant rich fraction of some common food grains. Int Food Res J 2011, 18:915–923.

34. Peng XF, Zheng ZP, Cheng KW, Shan F, Ren GX, Chen F, Wang MF: Inhibitory effect of mung bean extract and its constituents vexitin and isovitexin on the formation of advanced glycation endproducts. Food Chem 2008, 106:475–481.

35. Yao Y, Cheng X, Wang S, Wang L, Ren G: Influence of altitudinal variation on the antioxidant and anti-diabetic potential of azuki bean (Vigna angularis). Int J Food Sci Nutr 2011, 63:117–124.

36. Yao Y, Cheng X, Wang L, Wang S, Ren G: Biological potential of sixteen legumes in China. Int J Mol Sci 2011, 12:7048–7058.

37. Yao Y, Cheng X, Wang L, Wang S, Ren G: Major phenolic compounds, antioxidant capacity and anti-diabetic potential of rice bean (Vigna umbellata L.) in China. Int J Mol Sci 2012, 13:2707–2716.

38. Lim TY, Lim YY, Yule CM: Evaluation of antioxidant, antibacterial and anti-tyrosinase activities of four Macaranga species. Food Chem 2009, 114:594–599.

39. Association of Official Analytical Chemists (AOAC): Official Methods of Analysis of the Association of Official Analytical Chemists. 14th edition. Washington, DC, USA: AOAC; 1990.

40. Gotmer F: Chemisch-Phänotische Vitamin Ermittlungs-Methoden. Stuttgart, Germany: Ferdinand Enke Verlag; 1965.

41. Arinathan V, Mohan VR, Britto D, John A: Chemical composition of certain tribal pulses in South India. Int J Food Sci Nutr 2003, 54:209–217.

42. Ekpenyong TE, Barchers RL: Digestibility of proteins of winged bean seed. J Food Sci Tech 1979, 16:92–95.

43. Hu H, Vayak DL, Satterlee LD: A multienzyme technique for estimating protein digestibility. J Food Sci 1977, 42:1269–1271.

44. Goni I, Garcia-Alonso A, Saura-Calixto FA: A starch hydrolysis procedure to estimate glycemic index. Nutr Res 1997, 17:427–437.

45. ISO/IDoS 5509: International Standards. 1st edition. Genève, Switzerland: International Organization for Standardization; 1997.

46. Bätz M, Shuttle E, Their HP: Trennung von Tocopherol und Tocotrienolendurch HPLC. Fat Sci Tech 1992, 94:209–213.

47. Singleton VL, Rossi JA: Colorimetry of total phenolic with phophomolybdicyphosphotungstic acid reagents. Am J Environ Sci 1965, 16:144–158.

48. Heimler D, Vignolini P, Dini MG, Romani A: Rapid tests to assess the antioxidant activity of Phaseolus vulgaris L. dry bean. J Agric Food Chem 2005, 53:3033–3036.

49. Xu BJ, Chang SKC: A comparative study on phenolic profiles and antioxidant activities of legumes as affected by extraction solvents. J Food Sci 2007, 72:5159–5166.