Biochemical profile and bioactive potential of wild folk medicinal plants of Zygophyllaceae from Balochistan, Pakistan

Alia Ahmed¹, *Amjad Hameed² and Shazia Saeed¹

¹ Department of Botany University of Balochistan, Quetta Pakistan
² * Nuclear Institute for Agriculture and Biology (NIAB) Faisalabad, Pakistan

*Corresponding author: amjad46pk@yahoo.com

Abstract:

Recent focus is on analysis of biological activities of extracts from plant species. Zygophyllaceae is exceedingly important angiosperm family with many taxa being used in folk medicines widely dispersed in arid and semi-arid zones of Balochistan, Pakistan. Only a small proportion of them have been scientifically analyzed and many species are nearly facing extinction. Therefore present investigation explores the biochemical and bioactive potential of fourteen folk medicinal plants usually used for treatments of different ailments. Fresh aerial parts of nine taxa and two fruit samples were collected from plants growing in arid and semi-arid zones of Balochistan and analyzed for enzymatic, non-enzymatic and other biochemical activities. Higher phytochemical activities were detected in the aerial parts. Superoxide dismutase was detected maximum in Fagonia indica, (184.7±5.17 units/g), ascorbate peroxidase in Tribulus longipetalus subsp. longipetalus (947.5±12.5 Units/g), catalase and peroxidase was higher in Peganum harmala (555.0±5.0 and 2597.8±0.4 units/g respectively). Maximum esterase and alpha amylase activity was found in Zygophyllum fabago (14.3±0.44 and 140±18.8 mg/g respectively). Flavonoid content was high in T. longipetalus subsp. longipetalus (666.1±49 µg/ml). The highest total phenolic content and tannin was revealed in F. olivieri (72125±425 and 37050±1900 µM/g respectively). Highest value of ascorbic acid was depicted in F. bruguieri var. rechingeri (448±1.5 µg/g). Total soluble Proteins and reducing sugars were detected higher in P. harmala (372.3±54 and 5.9±0.1 mg/g respectively). Maximum total antioxidant capacity (TAC) was depicted in Z. simplex (16.9±0.01 μM/g). Pigment analysis exhibited the high value of lycopene and total carotenoids in T. terrestris (7.44±0.2 and 35.5±0.0 mg/g respectively). Chlorophyll a, b and total chlorophyll content was found maximum in T. longipetalus subsp. pterophorus (549.1±9.9, 154.3±10 and 703.4±20.2 ug/g respectively). All taxa exhibited anti-inflammatory activity as well as anti-diabetic inhibitory potential. Seed extracts of Zygophyllum eurypterum (96%) exhibited highest inhibitory potential, along with twelve other taxa of Zygophyllaceae indicated (96-76%) activity when compared with the standard drug diclofenac sodium (79%). Seeds of T. longipetalus subsp. longipetalus (85%) exhibited the highest anti-diabetic activity; other eleven taxa also exhibited inhibitory activity of α-amylase ranging from (85-69%) compared with Metformin (67%) standard drug. Phytochemical screening revealed that selected taxa proved to be the potential source of natural antioxidants and could further be explored for in-vivo studies and utilized in pharmaceutical industries as potent therapeutic agents validating their ethno-pharmacological uses.

Keywords: Zygophyllaceae, enzymatic antioxidants, Non-enzymatic antioxidants, Phytochemicals, anti-diabetic, anti-inflammatory, Balochistan Pakistan.
Introduction:
Most of the world’s population, particularly in the underdeveloped and developing countries, depends mainly on herbal medicines to cure various ailments [1]. Wild medicinal plants are being utilized as folk medicines globally. Numerous plant parts have various therapeutic properties and utilized as herbal medicines to treat various disorders and infections. Currently most prevailing and powerful drugs used were derived from medicinal plants [2,3] Thus, for biological and antioxidant capacity plant based derivatives must be explored. During the last few years herbal medicines and their derivatives seek much attention because of their traditional use. People of many developing countries depended on the medicinal plants for their healthcare. Focus on antioxidants specifically of herbal medicines originates from their ethno-pharmacological utilization [4]. Many medicinal plants extract constituted various types of chemicals, each have a property to control a variety of biological and pharmacological activities such as antimicrobial, anti-parasitic, anti-diabetic, antioxidant, anti-inflammatory and anticholinesterase. These chemicals may have a combined effect in controlling various diseases [5,6]. Diabetes mellitus is commonly related with metabolic disorders linked with numerous macro and micro vascular problems. These disorders accelerate morbidity and mortality [7-10]. It has also been revealed that oxidative stress, rise in free radicals and deterioration of antioxidant defense may mediate the prevalence of diabetes associated complications in diabetic patients [11,12]. Many degenerative disorders like rheumatoid arthritis, joints and shoulder inflammation, heart disease, muscular inflammation, asthma, cancer, and inflammation of gastrointestinal tract commonly related with inflammatory processes [13,14]. During various metabolic processes free radicals formed in the living systems. The reactive oxygen species (ROS) comprised mostly of hydroxyl radical (• OH), superoxide anion (O$_2^-$) and hydrogen peroxide (H$_2$O$_2$) formed in little quantity for diverse physiological processes [15]. Though, increased amount of these radicals resulted in various disorders like, Alzheimer disease, atherosclerosis, inflammation of various organs and cancer [16]. Oxidative stress, resulting from excessive ROS, causes detrimental effects on biomolecules, leading to many pathological and neurological disorders. The integrated antioxidant systems can balance the toxicity of oxidative reactive species which comprises of enzymatic and non-enzymatic antioxidants. Enzymatic antioxidants such as superoxide dismutase (SOD), peroxidase (POD), ascorbate peroxidase (APX) and catalase (CAT) remove hydrogen peroxide, scavenging free radicals and intermediates of oxygen [17]. Non-enzymatic antioxidants comprised of lipid soluble related with membrane like alpha tocopherol, beta carotene and water soluble reducers like ascorbate, glutathione and phenolic [18]. The antioxidants like polyphenols safeguard the biological systems and oxidative processes. Therefore, controlling diseases induced with oxidative stress by the consumption of herbal products as a rich source of phyto-antioxidants [19]. It has been proved that herbal extracts containing phyto-antioxidants particularly polyphenols, flavonoids, tannins and other associated compounds have progressive health effects and decrease disease risk. Recent researches revealed effects of bioactive and antioxidant potentials of medicinal plants. [20,21].

The Zygophyllaceae consists of diverse habits of wild flora including succulents, herbs, undershrub, shrubs and small trees. The habitat of these plants is predominantly desert or saline areas of temperate and tropical regions around the globe [22,23]. Tribulus, Fagonia, Zygophyllum and Peganum are being considered as the main genera. Many taxa of the family being used ethnomedicinally. Ref ?
*Fagonia* Linn., is a genus of wild, flowering plants of the family, Zygophyllaceae, having about 45 species all over the world. The distribution of the genus includes parts of Africa, the Mediterranean Basin, Asia and parts of the America. In Pakistan 10 species are reported earlier, in Balochistan it has 7 species present. While working in the field of southern Balochistan we collected 7 species including few sub-species of the genus.

*P. harmala* Linn., is a perennial herbaceous plant usually 25-60 cm tall yellowish white flowers appears in April to May and commonly dispersed in many regions of the province.

Genus *Tribulus* L. is considered as the most complex genus in Zygophyllaceae because of enormous number of invalid specific epithets and also of the variations present in various populations In Pakistan 4 species are reported.

*Zygophyllum* Linn., with about 90 species, grows mainly between northern Africa and central Asia mainly in arid and semi-arid areas.

Keeping in view the significance of the medicinal plants of Zygophyllaceae, the present study was conducted and most probably the first comprehensive report on phytochemical screening of leading genera of Zygophyllaceae from the arid and semi-arid regions of Balochistan, Pakistan. The specific objective was to evaluate the bioactive potential and to detect the anti-diabetic and anti-inflammatory activities of aerial parts and seed extracts of these medicinal plants to validate their folk-uses and subsequent utilization as a source of novel drugs.

**Materials and Methods:**

**Plant collection**

The selection of medicinal plants was based on ethno-botanical appraisal. Wild plants were collected from different areas of Balochistan, Pakistan. Voucher specimens were prepared, identified and submitted in Botanical garden Herbarium university of Balochistan, Quetta. Furthermore records also available in open herbarium for future reference (www.open herbarium.org). Fresh aerial parts of nine plants, fruits of two species of *Tribulus*, dry aerial parts of fourteen plants and ten seeds were collected from different ecological zones of Balochistan (Table 1). Fresh plant samples were transported and kept at -20°C. Samples were collected and shade dried and stored at room temperature. Experiments were conducted at Plant Breeding and Genetics Division (MAB Lab) Nuclear Institute for Agriculture and Biology (NIAB), Faisalabad, Pakistan.

Table 1: List of selected taxa of Zygophyllaceae with geographical coordinates of the collection sites and voucher specimen’s number

**Extraction of antioxidant enzymes**

Fresh aerial parts of nine plants, two fruits, dry aerial parts and seeds (0.15g) were subjected in 1ml phosphate buffer (50 mM, pH 7.8) to ground. Further the mixture was centrifuged at 14,000×g (20 min, 4°C). Now the supernatant of this extracted plant material used to perform further phytochemical activities. All the data were taken in replicates of three.

**Superoxide dismutase (SOD) assay:**

Method of [24] used to determine SOD activity by homogenizing the fresh aerial parts and fruits of selected taxa in phosphate buffer (50 mM, pH 7.8), EDTA (0.1 mM) and DTT (1 mM) following the procedure of [24]. Further analyzed by assessing its property to stop the photochemical reduction of nitro-blue tetrazolium as explicated by [25]. One unit of SOD activity was demarcated as the amount of enzyme causing 50% inhibition of photochemical reduction of nitro-blue tetrazolium.
Peroxidase (POD) assay:
The assessment of POD activity carried out using method [26] was followed with few changes. Homogenized mixture of the aerial parts and fruits prepared in 1 ml phosphate buffer (50 mM, pH 7.8), EDTA (0.1 mM) and DTT (1 mM). The assay solution contained 535µl distilled water, phosphate buffer 50 mM (pH 7.0), Guaiacol (20 mM), H₂O₂ (40 mM) and 15µl enzyme extract. The addition of enzyme extract initiated the reaction. At 470 nm absorbance rises was noted at interval of 20 sec. Absorbance change of 0.01 min⁻¹ was demarcated as one unit POD activity. Enzyme activity was expressed on the basis of fresh sample weight.

Catalase (CAT) assay:
Catalase activity was measured by homogenizing the aerial parts and fruit samples prepared in phosphate buffer (50 mM, pH 7.8), EDTA (0.1 mM) and DTT (1 mM). CAT was assessed by following the method of [26]. The activity was measured in a solution contained 59 mM H₂O₂ and 0.1 ml enzyme extract. At 240 nm decrease in absorbance recorded after interval of 20 seconds. Change in absorbance of 0.01 per min defined CAT activity of one unit. Enzyme activity was expressed on fresh weight basis.

Ascorbate peroxidase (APX) assay:
The assessment of APX activity carried out by following the [24] method. Samples were extracted in phosphate buffer (50 mM, pH 7.0). APX activity was analyzed by the method of [24]. For the measurement of APX the assay buffer contained potassium phosphate buffer (200 mM, pH 7.0), EDTA (0.5 M), ascorbate (10 mM), 1 ml of H₂O₂ and 50µl supernatant. At 290 nm absorbance decrease was noted after every thirty seconds to estimate the oxidation rate of ascorbic acid [27].

Hydrolytic Enzymes:
Esterase activity:
The α-esterase was determined by using the method [28]. α-naphthyl acetate used as substrates, the reaction mixture contained 30 mM α-naphthyl acetate (30 mM), acetone (1%), and phosphate buffer (0.04 M, pH=7) and enzyme extract. Incubate this mixture for 15 min at 27ºC in dark. After 15 min added staining solution 1 ml (Fast blue BB 1% and SDS 5% with ratio of 2:5) and again incubate in dark for (20 min, 27 ºC). Absorbance at 590nm was measured for α-naphthol produced amount. Using standard curve, enzyme activity was α-naphthol produced in µM min⁻¹ /g wt.

Alpha amylase activity:
A modified method for alpha amylase activity was followed for all plant samples as described by [29].

Other biochemical parameters:
Total oxidant status (TOS):
Method of [30] used to determined TOS. The assay established on ferrous ion oxidation into ferric ion. The presence of oxidants in the sample in acidic medium and ferric ion measurement produced by xylenol orange [31]. The assay based on two mixtures R1 (stock xylenol orange solution (0.38g in 500µL of 25Mm H₂SO₄) 0.4g NaCl, 500 µL glycerol and volume up to 50Ml with 25mM H₂SO₄ sample extract and R2 (0.0317 g o-dianisidine, 0.0196 g ferrous ammonium sulphate (II). Absorption measured at 560nm after 5 min by using spectrophotometer.

Pigment analysis:
The concentration of Lycopene, chlorophyll (a and b), Total chlorophyll and carotenoids were examined by method of [32]. Samples (0.2 g) were grind in acetone (80%) and centrifuged at
10,000 g for 5 minutes. Absorbance measured at 645, 663 and 480 nm by using a spectrophotometer.

**Total phenolic contents (TPC) and Tannin:**

Micro colorimetric assay was used to measure TPC, by using Folin-Ciocalteu (F-C) reagent [33] with some modifications. 0.05 g sample was kept in 95% methanol in dark for 48 hours. After 48 hours take the supernatant add 150 µl FC reagent (10%) and 1.2 ml sodium carbonate (700 mM). Place this mixture at room temperature for 1 hour and take reading at 765 nm. Linear regression equation was calculated by using standard curve of Gallic acid at different concentrations. To measure Tannin add (0.1g) PVPP and in above prepared sample vortex vigorously and centrifuge again at 14000 rpm, measured reading at 765nm.

**Determination of Total Flavonoid Content**

Assay was determined by colorimetric method using Quercetin as standard. Take 200 µl sample prepared in 95% methanol extract and phosphate buffer (40 mM, pH 6.8). Add 50 µl AlCl₂ (10%) 50 µl Potassium Acetate (1M) Incubate the mixture at room temperature for 40 min and take reading at 415nm absorbance.

**Total Antioxidant Capacity:**

A modified method of TAC was followed as described by [34]. Due to the presence of antioxidants in sample, ABTS assay represents decrease of 2, 2-azino-bis (3-ethylbenzothiazoline-6- sulfonate) radical cation ABTS⁺⁺ (blue-green in color) into original ABTS (colorless compound). The antioxidants of the sample extract according to their content decolorize the ABTS⁺⁺ radical cation. The reaction mixture contained reagent R1, sample extract and reagent R2. After 5 min at wavelength of 660nm, the absorption of each reaction mixture was measured. This analysis used AsA (ascorbic acid) to develop a calibration curve. The results for antioxidant contents found in plant extracts were measured as µM AsA equivalent to1g.

**Reducing Sugars (Sugar Content)**

Assessment of reducing sugars level in plant samples was determined by dinitrosalicylic acid method proposed by [35].

**Total soluble protein content**

Protein estimation of plant samples was based on quantitative protein analysis described by [36]. Aerial parts and fruits samples were homogenized in potassium phosphate (50 mM, pH 7.0). Supernatant 5µl and NaCl (0.1N) mixed with1.0 ml of Bradford dye. Incubate the mixture for 30 min to get a protein dye complex. Measure the quantity at 595 nm absorbance by spectrophotometer.

**In Vitro Anti-Diabetic Activity (enzyme α-amylase inhibition method)**

The in Vitro anti-diabetic activity was determined by assaying the inhibitory activity of the enzyme α-amylase which involves in the breakdown of starch to produce glucose [37]. In this method, 1 ml of methanolic extracts of all species were tested separately and thus added to 1 ml of the enzyme α-amylase in a test-tube and incubated for 10 min at 37°C. Then 1 ml of 1% starch solution was added into it and again incubated for 15 min at 37°C. Then 2 ml 3, 5-dinitrosalicylic acid reagent was added into it, in order to terminate the reaction. The reaction mixture was then incubated in boiling water bath for 5 min and then allows it to cool at room temperature. The absorbance of the reaction mixture was then measured at 546 nm in a spectrophotometer. The standard (control) of the reaction without the extract represents the 100% enzyme activity. The % age inhibition of enzyme activity of α-amylase was determined by:

\[
\%age \text{ inhibition of } \alpha-\text{amylase} = \frac{\text{Enzyme activity of control} - \text{Enzyme activity of extract}}{\text{Enzyme activity of control}} \times 100 \]

5
**In Vitro Anti-Inflammatory Activity (Protein Denaturation Method)**

The protein denaturation assay was determined using a modified method as described by [38]. Briefly, the reaction mixture (0.5 mL; pH 6.3) consisted of 0.45 mL of bovine serum albumin (5% aqueous solution) and 0.05 mL of distilled water. The pH was adjusted to 6.3 using a small amount of 1 N HCL. 1 mL of acetone or aqueous extract with final concentrations of (0.1 to 0.5 mg/mL) was added to the reaction mixture. These were incubated at 37°C for 30 min and then heated at 57°C for 5 min. After cooling the samples, 2.5 mL of phosphate buffer solution (pH 6.4) was added. Turbidity was measured spectrophotometrically at 660 nm. For the negative control, 0.05 mL of distilled water and 0.45 mL of bovine serum albumin were used. Diclofenac sodium with the final concentration of 100, 200, 300, 400, and 500 μg/mL was used as reference drug. The percentage inhibition of protein denaturation was calculated by using the following formula:

\[
\text{% age inhibition} = \frac{\text{Abs control} - \text{Abs sample}}{\text{Abs control}} \times 100
\]

Statistical Analysis:

Data was recorded in mean±SEM. Resulting data were analyzed by applying descriptive statistics. Two-way ANOVA with three replications were used in analyses. Significance of data was tested by analysis of variance and Tukey (HSD) Test at p<0.001 using XL-STAT software.

Results and Discussion

Present study is based on analysis of biological activities of extracts from fifteen plant species of Zygophyllaceae, remarkably important angiosperm family with many taxa being used in folk medicines. There are not much data available in literature about these plants so making a comparison of the results we obtained is not easy. Nonetheless, a few papers reported few biological activities of *T. terrestris*, *P. harmala* and few species of *Fagonia*. Present investigation explores the presence of enzymatic constituents such as SOD, POD, APX, CAT, esterase, alpha amylase, non-enzymatic antioxidants and other phytochemical like AsA, TOS, TAC, TSP, TPC, TF, tannin and pigments. Selected plants also provided evidence for anti-diabetic and anti-inflammatory potential of varying extent in seeds and aerial parts. The difference in pro-oxidants and antioxidants causes oxidative stress and chronic diseases in the body [39]. Cellular damage results in causing cancer. One of the mechanism behind the anti-oxidation is free radical scavenging action [40]. POD helps in scavenging the reactive oxygen species (ROS), causing cell oxidative injury [41]. Highest value of peroxidase and catalase were depicted in the aerial parts of *P. harmala* (2597.8±0.4 units/g f. wt. and 555.0±5.0 Units/g f. wt. respectively) show in S. Fig 1a and b respectively. Plant species having high antioxidant activities can be utilized for different therapeutic applications for treatment of oxidative stress induced diseases.

Antioxidant enzyme catalase present in all animal tissues with its highest activities in liver and red blood cells which defends the tissues from highly reactive hydroxyl radicals by decomposing hydrogen peroxide. Decrease amount of catalase causes numerous damages due to hydrogen peroxide and superoxide radical assimilation [42]. Superoxide dismutase enzyme referred as the significantly involved in cellular defense, therefore it is considered as an indicator of antioxidant
The highest value of superoxide dismutase (184.7±5.17 units/g f. wt.) was observed in *F. indica* shown in S. Fig. 1c. Traditionally this plant being used for anticancer treatment and possibly detected high concentrations of SOD and TAC may be responsible for this therapeutic effect.

Highest APX value was recorded in *T. longipetalus* subsp. *longipetalus* (947.5±12.5 Units/g f. wt.) S. Fig. 1d. Ascorbate peroxidase (APX) enzyme is crucial for the protection from damage by H$_2$O$_2$ and hydroxyl radicals (•OH) [44]. Antioxidant enzymes activities namely, CAT, POD and SOD were reported earlier in *Rumex obtusifolius* a wild medicinal plants and found to have good antioxidant capacity [45]. Similarly, *Calamintha officinalis* also has potent antioxidants [46]. Alpha amylase and esterase activity was higher in *Z. fabago* (140±18.8 mg/g. and 14.3±0.44 µM/min/g f. wt. respectively) S. Fig. 1e and f respectively. Esterase plays an important role in the disintegration of natural constituents and industrial pollutants and other toxic chemicals. It is also beneficial for the production of optically pure compounds, perfumes, and antioxidants [47].

Maximum TAC was depicted in fresh samples of *Z. simplex* (16.9±0.01 µM/g. f. wt.) followed by *F. indica* (15.6±0.04µM/g. f. wt.) shown S. Fig. 2a. No significant variation was detected among all dry aerial parts of selected plants. In general maximum TAC was detected in seed of *Z. eurypterum* (15.8±2.2 µM/g. dry wt.) followed by aerial parts of *Z. simplex* (15.7±2.33 µM/g.dry wt.). In seeds of selected taxa no significant variation was detected the maximum TAC was in *F. olivieri* (15.6±2.4 µM/g. s. wt.) second highest was in *F. bruguieri* var. *rechingeri* (15.5±2.5 µM/g. s. wt.). Previously aerial parts of *F. longispina* reported to be a good source for natural antioxidants [48]. Previously *F. cretica* was also found to have high antioxidant and radical scavenging potential due to high TPC and TFC [49]. *F. olivieri* can serve as a natural source to develop the free radical scavengers beneficial in the prevention of oxidative stress development [50].

Total Flavonoid content (TFC) in methanolic extract was showed maximum quantity in fresh samples of *T. longipetalus* subsp. *longipetalus* (666.1±49µg/ml sample) shown S. Fig 2b. Fruit samples (fresh) highest value of flavonoid and ascorbic acid was detected in *T. terrestris* (566.1 ±5.1 µg/ml sample and 456.5±9.5 µg/g f. wt. respectively). In dry aerial parts *T. longipetalus* subsp. *longipetalus* flavonoid content gives maximum amount of TFC (495.4±16.4 µg /mL sample) followed by *F. bruguieri* var. *laxa* (395.1±37 µg/mL sample). While in seeds *P. harmala* showed maximum TFC (418.8±16.7µg/mL sample) followed by *F. bruguieri* var. *bruguieri* (391.8±5.12 µg/mL sample) Flavonoids are considered to be effective free radical scavengers in fruits, vegetables, and medicinal plants. Highest ascorbic acid reported in *P. harmala* shown S. Fig 2c. Ascorbic acid involved in a number of physiological processes. such as POD and SOD [51].The highest flavonoids and ascorbic acid content in fruits of *T. terrestris* and aerial parts of *T. longipetalus* subsp. *longipetalus* in this study validated its traditional medicine use and may be responsible to cure various ailments. *F. olivieri* shown S. Fig 2c gives the highest amount of tannins.
Total soluble protein and reducing sugar were high in fresh aerial parts of *P. harmala* shown in S Fig. 3a and b respectively. Earlier [52] isolated antioxidant protein from *P. harmala*. Seeds possessed antioxidant activity, and this activity was due to the presence of hydrophobic amino acids. *P. harmala* is one of the most frequently used medicinal plants to treat hypertension and cardiac disease worldwide [53]. The maximum total soluble proteins (168.3±6.3 mg/g dry wt.) were depicted in dry aerial parts of *Z. simplex*. No significant variation was observed in seed samples, the highest value of total soluble proteins (248.6±30 mg/g s. wt.) was found in *F. bruguieri* var. *rechingeri*. Total oxidant status (TOS) was lower in *F. indica* (1275±475 µM/g. f. wt.) shown in S Fig. 3d.

S. Fig. 3: Comparison of a) Total soluble proteins b) Reducing Sugar c) Total Oxidant Status d) Total Antioxidant Capacity

Total flavonoid content was calculated in and expressed as µg/mL in methanolic extract using quercetin as standard (Table 2). Significant difference was observed among all selected species of Zygophyllaceae. In aerial parts *T. longipetalus* subsp. *longipetalus* flavonoid content gives maximum amount of TFC (495.4±16.4 µg/mL sample) followed by *F. bruguieri* var. *laxa* (395.1±37 µg/mL sample). While in seeds *P. hermala* showed maximum TFC (418.8±16.7 µg/mL sample) followed by *F. bruguieri* var. *bruguieri* (391.8±5.12 µg/mL sample) (Table 3). Total Phenolic Content (TPC) was estimated, in aerial parts of selected taxa no significant TPC variation was found among (Table 4). In general, highest TPC was depicted in *T. longipetalus* subsp. *pterophorus* (63025±1725 µM/g. dry wt.) followed by *F. bruguieri* var. *rechingeri* (54600±1350 µM/g. dry wt.). Seeds of selected plant samples showed significant variation. Highest TPC was detected in *Z. propinquum* (69225±775 µM/g. s. wt.) followed by *F. bruguieri* var. *rechingeri* (66850±3900 µM/g. s. wt.) shown in Table 3. No significant Tannin variation was detected among aerial parts as well as in seeds of all tested taxa (Table 2). However, highest amount of tannin was estimated in *T. longipetalus* subsp. *pterophorus* (40375±4125 µM/g dry wt.) followed by *Z. fabago* (39175±4825 µM/g. dry wt.). In seeds highest amount of tannin was estimated in *P. hermala* (47525±2575 µM/g s. wt.) followed by *Z. propinquum* (42625±175 µM/g. s. wt.) shown in Table 3. Ascorbic Acid was observed, in aerial parts of all selected taxa and significant difference found among studied taxa (Table 4). Highest value of AsA was found in *F. olivieri* (744.2±2.7 µg/g dry wt.) followed by *F. bruguieri* var. *bruguieri*. AsA content in seeds showed no significant variation. In general the highest AsA content was found in *F. bruguieri* var. *laxa* (740.8±2.19 µg/g s. wt.) given in Table 3. Alpha amylase activity in dry aerial parts of selected plants was assessed (Table 2) no significant variation was found among the taxa of Zygophyllaceae. However, highest α-amylase activity was found in *Z. simplex* (164.9±3.39 mg/g dry wt.) followed by *Z. fabago* (153±6.6 mg/g dry wt.). In seeds of selected taxa significant variation was found among various taxa is shown in (Fig 19 f). The highest value was observed in *F. bruguieri* var. *laxa* (159.5±11.8 mg/g s. wt.) followed by *F. ovalifolia* subsp. *pakistanica* (133.9±0.37 mg/g. s. wt.) given in Table 3. Significant variation of total soluble protein was observed among all tested taxa dry samples (Table 2). The maximum total soluble proteins (168.3±6.3 mg/g dry wt.) were depicted in *Z. simplex*. No significant variation was observed in seed samples, the highest value of total soluble proteins (248.6±30 mg/g s. wt.) was found in *F. bruguieri* var. *rechingeri* (Table 3). TAC was measured in aerial parts of selected taxa of Zygophyllaceae (Table 2), No significant variation
was detected among all selected plants. In general maximum TAC was detected in *Z. eurypterum* (15.8±2.2 µM/g. dry wt.) followed by *Z. simplex* (15.7±2.33 µM/g. dry wt.). In seeds of selected taxa no significant variation was detected the maximum TAC was in *F. olivieri* (15.6±2.4 µM/g. s. wt.) second highest was in *F. bruguieri* var. *rechingeri* (15.5±2.5 µM/g. s. wt.) given in Table 3. Reducing Sugar was measured in aerial parts of all selected plants (Table 2) there was a significant difference found among all taxa. Highest value of reducing sugar was recorded in *Z. fabago* (7.47±0.2 mg/g. s. wt.) followed by *Z. propinquum* and with minimum difference (7.19±0.55 mg/g. s. wt.). In seeds highest value of reducing sugar was found in *T. terrestris* (7.9±0.1 mg/g. s. wt.) given in Table 3.

**Table 2: Phytochemical analysis in dry aerial parts of selected taxa of Zygophyllaceae**

**Table 3: Phytochemical analysis in seeds of selected taxa of Zygophyllaceae**

Lycopene content in dry aerial parts of various taxa was measured (Table 4). The higher value was found in *T. longipetalus* subsp. *pterophorus* (9.25± 1.8 mg/g dry wt.) followed by *T. terrestris* (8.12±0.01 mg/g dry wt.). Lycopene content in seeds of various taxa of Zygophyllaceae was investigated (Table 5). The higher value of lycopene was found in *F. ovalifolia* subsp. *pakistanica* (5.87±0.75 mg/g s. wt.) followed by *F. bruguieri* var. *rechingeri* (4.92±0.19 mg/g s. wt.). Chlorophyll a content was estimated in dry aerial parts of dry samples of various taxa of Zygophyllaceae (Table 4). The Chlorophyll a content ranged from maximum in *T. longipetalus* subsp. *pterophorus* (572.1±0.05 ug/g dry wt.) followed by *T. terrestris* (548.8±0.2 ug/g dry wt.) to minimum with significant difference in *Z. eurypterum* (92.5±8.0 ug/g dry wt.). Chlorophyll a content in seeds of various taxa (Table 5) ranged from maximum in *F. bruguieri* var. *rechingeri* (197±0.76 ug/g s. wt.) followed by *F. ovalifolia* subsp. *pakistanica* (171.2±5.3 ug/g s. wt.). Chlorophyll b content was assessed in dry aerial parts of different taxa of Zygophyllaceae (Table 4) Chlorophyll b content varied among various taxa. The Chlorophyll b content ranged from maximum in *T. longipetalus* subsp. *pterophorus* (228 ±63 ug/g dry wt.) followed by *T. terrestris* (170.2±1.6 ug/g dry wt.). In seeds of different taxa Chlorophyll b content varied ranged from maximum in *F. ovalifolia* subsp. *pakistanica* (70.7±11.2 ug/g s. wt.) next highest was *F. bruguieri* var. *rechingeri* (38±1.9 ug/g s. wt.) to minimum in *Z. propinquum* (2.9±0.01ug/g s. wt.) with a significant difference (Table 5). Total carotenoids were investigated in dry aerial parts of dry samples of different species of Zygophyllaceae (Table 4). Significantly highest value of total carotenoids was found in *T. longipetalus* subsp. *pterophorus* (38.6±0.6 mg/g dry wt.) followed by *T. terrestris* (38.2±0.02 mg/g dry wt.). In seeds samples significantly highest value of total carotenoids was found in *F. bruguieri* var. *rechingeri* (19.3±0.00 mg/g s. wt.) (Table 5) followed by *F. ovalifolia* subsp *pakistanica* (16.6±1.5 mg/g s. wt.). Total chlorophyll content of dried aerial parts was measured among different taxa of zygophyllaceae is shown in (Table 4). Maximum Content of total chlorophyll was depicted in *T. longipetalus* subsp. *pterophorus* (800±62.9 ug/g dry wt.) with significant difference. It was followed by *T. terrestris* (719±1.8 ug/g dry wt.). In seeds measured among different taxa of Zygophyllaceae is given in (Table 5). Significant variation was observed among various taxa. Maximum Content of total chlorophyll was depicted in *F. ovalifolia* subsp. *pakistanica* (242±16.6 ug/g s. wt.) Second highest in *F. bruguieri* var. *rechingeri* (235.3±3.9 ug/g s. wt.). The liquid chlorophyll supplements are important in enhancing energy, detoxification of liver and stomach and colon, eliminating the body and mouth odor, helping in anemia and aiding in the elimination of mould from the body. The carotenoids are vital because of their strong colours, antioxidant activity as
well as their role as precursors of vitamin A. These can be used as safe chemicals for
neutraceutical purposes and food supplementation [54].

**Table 4: Pigment analysis in dry aerial parts of selected taxa of Zygophyllaceae**

**Table 5: Pigment analysis in seeds of selected taxa of Zygophyllaceae**

Pigment analysis in fresh samples shown in S. Fig 4 a, b, c, d, and e gives the significant
variation among the selected taxa. The chlorophyll contents (a, b and total), were higher in aerial
parts of *T. longipetalus* subsp. *pterophorus* followed by *T. terrestris*. While carotenoids and
lycopene were depicted maximum in *T. terrestris* followed by *T. longipetalus* subsp.
*pterophorus*. The main carotenoids such as zeaxanthin, b-carotene, canthaxanthin, astaxanthin as
well as lycopene are prepared synthetically in neutraceutical industry [55]. Furthermore,
Lycopene is suggested as one of the efficient carotenoids group for quenching ability. The plants
used in current study detected pigments in Zygophyllaceae taxa can be characterized for various
above mentioned applications can be utilized as supplements or medicines.

S. Fig.4: Comparison of Pigment a) Lycopene content b) Chlorophyll a content c) Chlorophyll b
content d) Total carotenoids e) Total chlorophyll content

This study was carried out to assess the anti-inflammatory and anti-diabetic potential of naturally
occurring medicinal plants of Zygophyllaceae from desert and semi-desert areas of Balochistan,
Pakistan. The protein denaturation assay was followed for in vitro anti-inflammatory activity
while anti-diabetic activity was determined by α-amylase inhibitory assay for 13 extracts of
aerial parts and 12 seeds extracts. Selected plants also used in traditional medicines by the folks
in Balochistan. Selection of plants was based on Ethnobotanical appraisal of local communities,
their uses in folk medicines to cure various ailments like fever, cough, inflammation of organs,
gonorrhea, urinary tract infection (UTI), diabetes and cancer etc. Previously secondary
metabolites such as flavonoids and alkaloids derivatives were evaluated in Zygophyllaceae
[56,57].

Inflammation in medical terms is demarcated as a pathophysiological procedure described by
soreness, fever, swelling of body parts, loss of function and discomfort [58]. The inhibitory
effect among different plants of Zygophyllaceae on albumin denaturation is shown in (Fig. 5).
Significant inhibition of albumin was observed among different taxa. *Z. eurypterum* seeds
depicted maximum inhibition with the highest value (96.85±1.85 % Inh.). Seeds of *T.
longipetalus* subsp. *pterophorus* revealed the next highest value (95.85±2.85 %), Seeds of *T.
terrestris* exhibited the (95.35±3.35% Inh.) activity. *T. longipetalus* subsp. *longipetalus*
depicted (91.1±1.1% of Inh.). Earlier in the fruit of *T. terrestris* significant anti-inflammatory activity was
evaluated [59]. All species of *Fagonia* also exhibited anti-inflammatory activity significant and
comparable with the standard drug diclofenac sodium. In seeds extract highest value were found
in *F. bruguieri* var. *laxa* (88.9 ± 4.9% Inh). *F. ovalifolia* subsp. *pakistanica* showed (84.8±0.8%)
of inhibition. *F. olivieri* and *F. bruguieri* var. *rechingeri* showed (80.9 ± 0.9 and 79.7±0.7% Inh.
respectively). Seeds of *P. harmala* constituted (76.4±1.4 % Inh.) activity for inhibition. Seeds of
*F. bruguieri* var. *bruguieri* (66.2±1.2% Inh.) revealed less inhibition when compared with the
standard drug. Previously anti-inflammatory activity of *F. cretica* was studied by [60]. Minimum
inhibition of albumin was observed in seeds of *Z. propinquum* (15.79±0.20% Inh.) in comparison
with all studied taxa. Previously, *Nitraria schoberi* from Zygophyllaceae fruit extract has been
found to have anti-inflammatory effects [61] that are quite in line with presently observed
activities for seeds and aerial parts. Aerial parts of all the studied taxa of Zygophyllaceae also exhibited significant inhibition of albumin. *T. longipetalus* subsp. *pterophorus* revealed maximum inhibition (90.1±2.1 % Inh.) followed by *T. terrestris* and *Fagonia bruguieri* var. *laxa* (89.9±0.9 and 89.4±0.9 % Inh. respectively) anti-inflammatory inhibition. *Z. fabago* and *F. olivieri* indicated (88.3 ± 1.3 % Inh.) inhibition in both plants. *F. ovalifolia* subsp. *pakistanica* ( endemic to the region) revealed (87.1±1.1% Inh.) albumin inhibition activity. *T. longipetalus* subsp. *longipetalus*, *Z. simplex* and *F. bruguieri* var. *bruguieri* exhibited inhibition activity as (86.5 ± 0.5, 83.2 ±1.2 and 80.4 ± 1.2 % Inh. respectively). *P. harmala* and *Z. propinquum* aerial parts also showed significant inhibition as compared with standard drug (79.6±0.6 and 75.9±1.9 % Inh.). *F. indica* (49.6±0.3 % Inh.) depicted less inhibition when compared with all other taxa as well as standard drug. Previously chloroform and methanol extract of aerial parts of *T. terrestris* exhibited significant anti-inflammatory activities at a dose of 200 mg/kg [59]. All examined *Fagonia* spp. in present study exhibited therapeutic potential especially *F. bruguieri* var. *laxa* suggested to be used as anti-inflammatory as well as anti-diabetic agent authenticating its folk use. Earlier, *F. cretica* was identified for its high therapeutic effects against various types of hematological, liver disorders, neurological and inflammatory conditions. Furthermore, aqueous extract is one of the seventeen ingredients in Norm acid syrup used in the treatment of high acidity and gastritis [62]. Furthermore, the endemic taxa of the region i.e. *F. ovalifolia* subsp. *pakistanica* also exhibited high anti-inflammatory potential. Previously *F. longipina* traditionally used as a preventive for cancer, also used for the treatment of inflammation of the urinary tract [48]. *F. schweinfurthii* plant extract gel could be developed as a therapeutic agent for wound healing and anti-inflammatory properties [63].

Fig 5: Comparison of Anti-inflammation activity among different taxa of Zygophyllaceae. Data are presented as mean values ± SEM (n = 3). Statistical analysis: ANOVA test and Tukey (HSD). The different letters above the values in the same column indicate significant differences with Tolerance: 0.0001.

Anti-diabetic activity was analyzed by using amylase inhibition assay. Seeds and aerial parts of selected species revealed significant differences in anti-diabetic activity when compared with standard drug Metformin (Fig.6). The seeds of *T. longipetalus* subsp. *longipetalus* (85.65±0.34 % Inh.) exhibited the highest anti-diabetic activity among all selected species. It was followed by seeds of *Z. eurypterum* (83.63±0.63% Inh.), next highest value was found in *T. terrestris* (82.8±0.1%). Seeds of *Z. propinquum* indicated (81.5±0.4%) activity. *F. bruguieri* var. *rechingeri* depicted (80.9±1.9 %) anti-diabetic activity. *F. bruguieri* var *bruguieri* revealed (80.3±0.6 %) enzyme inhibition in the seeds of *F. bruguieri* var. *laxa* (79.5±1.0%) inhibition was found. Seeds of *T. longipetalus* subsp. *pterophorus* reported (77.7±0.2%) enzyme inhibition activity. The endemic plant of the region *F. ovalifolia* subsp. *pakistanica* depicted (76.5±0.4%) anti-diabetic activity. *P. harmala* showed (71.4±1.4%) anti-diabetic activity. Seeds of *F. olivieri* exhibited (69.8±0.8%) enzymatic inhibitory activity Seeds of *P. harmala* used as anti-inflammatory, antidiabetic [64]. Same therapeutic potential of seeds of *P. harmala* was attained in the current study.

Aerial parts of selected taxa also exhibited significant enzyme inhibition activity when compared with standard drug. Highest value was found in *F. bruguieri* var. *rechingeri* (83.59±0.40 % Inh.). Second highest value was in *T. longipetalus* subsp. *pterophorus* (83.37±0.62 % Inh.). Next highest value was found in *Z. simplex* (81.3±1.3% Inh.) followed by *Z. propinquum* (80.2±1.2 % Inh.) succulent plants. *F. bruguieri* var *laxa* depicted (78.3±1.1 % Inh.) anti-diabetic activity. *F. indica* showed (77.7±0.2 % Inh.) activities followed by *T. longipetalus* subsp. *longipetalus* (77
±1.06% Inh.) inhibition. Aerial parts of *Z. fabago* shows (76.2±1.2% Inh.) inhibitory effect. *F. bruguieri* var. *bruguieri* showed (75.5±0.5% Inh.) enzymatic inhibitory activity. Aerial parts of endemic plant *F. ovalifolia* subsp. *pakistanica* also showed significant inhibition of amylase enzyme (64.9±0.05% Inh.) Earlier reported *F. indica* alone or combined with *Aloe vera* can be used as a natural blood glucose lowering agent [65]. In present research the other examined *Fagonia* species are potentially active therapeutic agents in comparison with *F. indica*. *P. harmala* and *Z. gaetulum* frequently used to treat hypertension and diabetes mellitus [53].

**Fig 6**: Comparison of Anti-diabetic activity of different taxa of Zygophyllaceae Data are presented as mean values ± standard deviation (n = 3). Statistical analysis: ANOVA test and Tukey (HSD). The different letters above the values in the same column indicate significant differences with Tolerance: 0.0001

**Conclusion**

Phytochemical screening of all selected of Zygophyllaceae revealed that these plants have significant potential of enzymatic, non-enzymatic activities and other phytochemicals like flavonoids, total phenolic compounds, tannins, and pigments. All the taxa proved to have natural antioxidants and could not only be used to treat various ailments but also contribute in the prevention of degenerative diseases and manufacturing of new drugs. Research findings could be utilized for isolation of potential phytophobicological active compounds from these wild medicinal plants for future research. Species such as *Z. eurypterum*, *T. longipetalus* subsp. *longipetalus*, *T. terresetris*, *F. bruguieri* and *F. ovalifolia* subsp. *pakistanica* had greater potential for identification, isolation and purification of novel therapeutic agents.

**References**

1. Joshi LS, Pawar HA (2015) Herbal cosmetics and cosmeceuticals: An overview. Nat Prod Chem Res 3: 170.
2. Uniyal SK, Singh K, Jamwal P, Lal B (2006) Traditional use of medicinal plants among the tribal communities of Chhota Bhangal, Western Himalaya. Journal of ethnobiology and ethnomedicine 2: 14.
3. Srivastava JP, Lambert J, Vietmeyer N (1996) Medicinal plants: An expanding role in development: The World Bank.
4. Ullah M, Khan MU, Mahmood A, Malik RN, Hussain M, et al. (2013) An ethnobotanical survey of indigenous medicinal plants in Wana district south Waziristan agency, Pakistan. Journal of ethnopharmacology 150: 918-924.
5. Rates SMK (2001) Plants as source of drugs. Toxicon 39: 603-613.
6. Houghton P, Howes M-J, Lee C, Steventon G (2007) Uses and abuses of in vitro tests in ethnopharmacology: visualizing an elephant. Journal of Ethnopharmacology 110: 391-400.
7. Fowler MJ (2008) Microvascular and macrovascular complications of diabetes. Clinical diabetes 26: 77-82.
8. Halder N, Joshi S, Gupta S (2003) Lens aldose reductase inhibiting potential of some indigenous plants. Journal of Ethnopharmacology 86: 113-116.
9. Altan VM (2003) The pharmacology of diabetic complications. Current medicinal chemistry 10: 1317-1327.

10. Thomas M, Tsalamandris C, Maclsaac R, Jerums G (2005) Anaemia in diabetes: an emerging complication of microvascular disease. Current diabetes reviews 1: 107-126.

11. Jin L, Xue H-Y, Jin L-J, Li S-Y, Xu Y-P (2008) Antioxidant and pancreas-protective effect of aucubin on rats with streptozotocin-induced diabetes. European journal of pharmacology 582: 162-167.

12. Vos T, Flaxman AD, Naghavi M, Lozano R, Michaud C, et al. (2012) Years lived with disability (YLDs) for 1160 sequelae of 289 diseases and injuries 1990–2010: a systematic analysis for the Global Burden of Disease Study 2010. The lancet 380: 2163-2196.

13. Iwalewa E, McGaw L, Naidoo V, Eloff J (2007) Inflammation: the foundation of diseases and disorders. A review of phytomedicines of South African origin used to treat pain and inflammatory conditions. African Journal of Biotechnology 6.

14. Polya G (2003) Biochemical targets of plant bioactive compounds: a pharmacological reference guide to sites of action and biological effects: CRC press.

15. Scartezzini P, Speroni E (2000) Review on some plants of Indian traditional medicine with antioxidant activity. Journal of ethnopharmacology 71: 23-43.

16. Shehab NG, Abu-Gharbieh E, Bayoumi FA (2015) Impact of phenolic composition on hepatoprotective and antioxidant effects of four desert medicinal plants. BMC complementary and alternative medicine 15: 401.

17. Lee Y-P, Kim S-H, Bang J-W, Lee H-S, Kwak S-S, et al. (2007) Enhanced tolerance to oxidative stress in transgenic tobacco plants expressing three antioxidant enzymes in chloroplasts. Plant cell reports 26: 591-598.

18. Jaleel CA, Riadh K, Gopi R, Manivannan P, Ines J, et al. (2009) Antioxidant defense responses: physiological plasticity in higher plants under abiotic constraints. Acta Physiologiae Plantarum 31: 427-436.

19. Afsar T, Razak S, Khan MR, Mawash S, Almajwal A, et al. (2016) Evaluation of antioxidant, anti-hemolytic and anticancer activity of various solvent extracts of Acacia hydaspica R. Parker aerial parts. BMC complementary and alternative medicine 16: 258.

20. Agbor GA, Moumbeigna P, Oluwasola EO, Nwosu LU, Njoku R-C, et al. (2011) Antioxidant capacity of some plants foods and beverages consumed in the
eastern region of Nigeria. African Journal of Traditional, Complementary and Alternative Medicines 8.

21. Vayalil PK (2002) Antioxidant and antimutagenic properties of aqueous extract of date fruit (Phoenix dactylifera L. Arecaceae). Journal of Agricultural and Food Chemistry 50: 610-617.

22. Bellstedt D, Van Zyl L, Marais E, Bytebier B, De Villiers C, et al. (2008) Phylogenetic relationships, character evolution and biogeography of southern African members of Zygophyllum (Zygophyllaceae) based on three plastid regions. Molecular Phylogenetics and Evolution 47: 932-949.

23. Beier B-A, Chase M, Thulin M (2003) Phylogenetic relationships and taxonomy of subfamily Zygylloideae (Zygophyllaceae) based on molecular and morphological data. Plant Systematics and Evolution 240: 11-39.

24. Dixit V, Pandey V, Shyam R (2001) Differential antioxidative responses to cadmium in roots and leaves of pea (Pisum sativum L. cv. Azad). Journal of Experimental Botany 52: 1101-1109.

25. Giannopolitis CN, Ries SK (1977) Superoxide dismutases: I. Occurrence in higher plants. Plant physiology 59: 309-314.

26. Chance B, Maehly A (1955) [136] Assay of catalases and peroxidases. Methods in enzymology 2: 764-775.

27. Chen G-X, Asada K (1989) Ascorbate peroxidase in tea leaves: occurrence of two isozymes and the differences in their enzymatic and molecular properties. Plant and Cell Physiology 30: 987-998.

28. Van Asperen K (1962) A study of housefly esterases by means of a sensitive colorimetric method. Journal of insect physiology 8: 401-416.

29. Varavinit S, Chaokasem N, Shobsngob S (2002) Immobilization of a thermostable alpha-amylase. Science Asia 28: 247-251.

30. Erel O (2005) A new automated colorimetric method for measuring total oxidant status. Clinical biochemistry 38: 1103-1111.

31. Harma M, Harma M, Erel O (2005) Oxidative stress in women with preeclampsia. American Journal of Obstetrics & Gynecology 192: 656-657.

32. LICHTENTHALER HK, Wellburn AR (1983) Determinations of total carotenoids and chlorophylls a and b of leaf extracts in different solvents. Portland Press Limited.

33. Ainsworth EA, Gillespie KM (2007) Estimation of total phenolic content and other oxidation substrates in plant tissues using Folin–Ciocalteu reagent. Nature protocols 2: 875.
34. Nenadis N, Lazaridou O, Tsimidou MZ (2007) Use of reference compounds in antioxidant activity assessment. Journal of agricultural and food chemistry 55: 5452-5460.
35. Miller GL (1959) Use of dinitrosalicylic acid reagent for determination of reducing sugar. Analytical chemistry 31: 426-428.
36. Bradford MM (1976) A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Analytical biochemistry 72: 248-254.
37. Prabhakar P, Kumar A, Doble M (2014) Combination therapy: a new strategy to manage diabetes and its complications. Phytomedicine 21: 123-130.
38. Murugan R, Parimelazhagan T (2014) Comparative evaluation of different extraction methods for antioxidant and anti-inflammatory properties from Osbeckia parvifolia Arn.–An in vitro approach. Journal of King Saud University-Science 26: 267-275.
39. Wu B, Ootani A, Iwakiri R, Sakata Y, Fujise T, et al. (2006) T cell deficiency leads to liver carcinogenesis in Azoxymethane-treated rats. Experimental Biology and Medicine 231: 91-98.
40. Oshaghi EA, Tavilani H, Khodadadi I, Goodarzi MT (2015) Dill tablet: a potential antioxidant and anti-diabetic medicine. Asian Pacific Journal of Tropical Biomedicine 5: 720-727.
41. Vicuna D (2005) The role of peroxidases in the development of plants and their responses to abiotic stresses.
42. Rashid U, Khan MR, Sajid M (2016) Hepatoprotective potential of Fagonia Olivieri DC. Against acetaminophen induced toxicity in rat. BMC complementary and alternative medicine 16: 449.
43. Sudipta K, Kumara Swamy M, Balasubramanya S, Anuradha M (2014) Assessment of genetic fidelity, antioxidant enzyme activity and proline content of micropropagated and field grown plants of Leptadenia reticulata(wight & arn.)-an endangered medicinal plant. Plant Cell Biotechnol Mol Biol 15: 127-135.
44. Gangwar S, Singh VP, Tripathi DK, Chauhan DK, Prasad SM, et al. (2014) Plant responses to metal stress: the emerging role of plant growth hormones in toxicity alleviation. Emerging technologies and management of crop stress tolerance: Elsevier. pp. 215-248.
45. Alici EH, Arabaci G (2016) Determination of SOD, POD, PPO and cat enzyme activities in Rumex obtusifolius L. Annual Research & Review in Biology 11: 1-7.
46. Moattar FS, Sariri R, Giahi M, Yaghmaee P, Ghafoori H, et al. (2015) Antioxidant and anti-proliferative activity of Calamintha officinalis extract on breast cancer cell line MCF-7. Journal of Biological Sciences 15: 194-198.
47. Panda T, Gowrishankar B (2005) Production and applications of esterases. Applied microbiology and biotechnology 67: 160-169.
48. Hamidi N, Lazouni H, Moussaoui A, Ziane L, Djellouli M, et al. (2014) Ethnopharmacology, antibacterial and antioxidant activities, phytochemical screening of bioactive extracts from the aerial parts of Fagonia longispina. Asian J Nat Appl Sci 3: 53-63.
49. Iqbal P, Ahmed D, Asghar MN (2014) A comparative in vitro antioxidant potential profile of extracts from different parts of Fagonia cretica. Asian Pacific journal of tropical medicine 7: S473-S480.
50. Rashid U, Khan MR, Jan S, Bokhari J, Shah NA (2013) Assessment of phytochemicals, antimicrobial and cytotoxic activities of extract and fractions from Fagonia olivieri (Zygophyllaceae). BMC complementary and alternative medicine 13: 167.
51. Cavusoglu K, Bilir G (2015) Effects of ascorbic acid on the seed germination, seedling growth and leaf anatomy of barley under salt stress. J Agric Biol Sci 10: 124-129.
52. Ahmed H, ELZAHAB HA, Alswiai G (2013) Purification of antioxidant protein isolated from Peganum harmala and its protective effect against CCl4 toxicity in rats. Turkish Journal of Biology 37: 39-48.
53. Tahraoui A, El-Hilaly J, Israili Z, Lyoussi B (2007) Ethnopharmacological survey of plants used in the traditional treatment of hypertension and diabetes in south-eastern Morocco (Errachidia province). Journal of ethnopharmacology 110: 105-117.
54. Liu D, Shi J, Ibarra AC, Kakuda Y, Xue SJ (2008) The scavenging capacity and synergistic effects of lycopene, vitamin E, vitamin C, and β-carotene mixtures on the DPPH free radical. LWT-Food Science and Technology 41: 1344-1349.
55. Del Campo JA, García-González M, Guerrero MG (2007) Outdoor cultivation of microalgae for carotenoid production: current state and perspectives. Applied microbiology and biotechnology 74: 1163-1174.
56. Tulyaganov T, Allaberdiev FK (2003) Alkaloids from plants of the Nitraria genus. Structure of sibiridine. Chemistry of natural compounds 39: 292-293.
57. Salem JH, Chevalot I, Harscoat-Schiavo C, Paris C, Fick M, et al. (2011) Biological activities of flavonoids from Nitraria retusa (Forssk.) Asch. and their acylated derivatives. Food Chemistry 124: 486-494.

58. Hyun E, Bolla M, Steinhoff M, Wallace JL, Del Soldato P, et al. (2004) Anti-inflammatory effects of nitric oxide-releasing hydrocortisone NCX 1022, in a murine model of contact dermatitis. British journal of pharmacology 143: 618-625.

59. Mohammed MS, Alajmi MF, Alam P, Khalid HS, Mahmoud AM, et al. (2014) Chromatographic finger print analysis of anti-inflammatory active extract fractions of aerial parts of Tribulus terrestris by HPTLC technique. Asian Pacific journal of tropical biomedicine 4: 203-208.

60. Rawal A, Nath D, Yadav N, Pande S, Meshram S, et al. (2009) Rubia cordifolia, Fagonia cretica linn and Tinospora cordifolia exert anti-inflammatory properties by modulating platelet aggregation and VEGF, COX-2 and VCAM gene expressions in rat hippocampal slices subjected to ischemic reperfusion injury. Int J Appl Res Nat Prod 2: 19-26.

61. Sharifi-Rad J, Hoseini-Alfatemi SM, Sharifi-Rad M, Da Silva JAT (2015) Antibacterial, antioxidant, antifungal and anti-inflammatory activities of crude extract from Nitraria schoberi fruits. 3 Biotech 5: 677-684.

62. Shah K, Kumar RG, Verma S, Dubey R (2001) Effect of cadmium on lipid peroxidation, superoxide anion generation and activities of antioxidant enzymes in growing rice seedlings. Plant Science 161: 1135-1144.

63. Alqasoumi SI, Yusufoglu HS, Alam A (2011) Anti-inflammatory and wound healing activity of Fagonia schweinfurthii alcoholic extract herbal gel on albino rats. African Journal of Pharmacy and Pharmacology 5: 1996-2001.

64. Khan AM, Qureshi RA, Gilani SA, Ullah F (2011) Antimicrobial activity of selected medicinal plants of Margalla hills, Islamabad, Pakistan. Journal of Medicinal Plants Research 5: 4665-4670.

65. Mahdy A, Shehab NG (2015) Hypoglycemic Activity of Fagonia indica and Aloe vera in Alloxan-Induced Hypergly-cemia in Mice. EC Pharmaceutical Science 2: 239-244.
Fig 5: Comparison of Anti-inflammatory activity among different taxa of Zygophyllaceae. Data are presented as mean values ± SEM (n = 3). Statistical analysis: ANOVA test and Tukey (HSD). The different letters above the values in the same column indicate significant differences with Tolerance: 0.0001.
Fig 6: Comparison of Anti-diabetic activity of different taxa of Zygophyllaceae Data are presented as mean values ± standard deviation (n = 3). Statistical analysis: ANOVA test and Tukey (HSD). The different letters above the values in the same column indicate significant differences with Tolerance: 0.0001