

Invasive success of star weed (*Parthenium hysterophorus* L.) through alteration in structural and functional peculiarities

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A study was conducted on fifteen distinct populations of the star weed (*Parthenium hysterophorus* L.) to investigate the factors contributing to its widespread distribution in diverse environmental conditions. The results revealed significant variations in growth performance, physiological traits, and internal structures among populations from different habitats. The populations from wastelands exhibited superior growth, with higher accumulation of soluble proteins (TSP) and chlorophyll content (Chl a&b, TChl, Car, and Chl a/b). These populations displayed increased root and stem area, storage parenchyma, vascular bundle area, metaxylem area, and phloem area. Significant leaf modifications included thicker leaves, sclerification around vascular bundles, and widened metaxylem vessels. Roadside populations possessed larger leaf area, enhanced antioxidant activity, increased thickness of leaves in terms of midrib and lamina, and a higher cortical proportion. Populations found in agricultural fields depicted enhanced shoot biomass production, higher levels of chlorophyll b, and an increased total chlorophyll/carotenoid ratio. Additionally, they exhibited increased phloem area in their roots, stems, and leaves, with a thick epidermis only in the stem. All these outcomes of the study revealed explicit structural and functional modifications among *P. hysterophorus* populations collected from different habitats. These variations were attributed to the environmental variability and could contribute to the widespread distribution of this species.

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Abstract

10 A study was conducted on fifteen distinct populations of the star weed (*Parthenium hysterophorus*
11 L.) to investigate the factors contributing to its widespread distribution in diverse environmental
12 conditions. The results revealed significant variations in growth performance, physiological traits,
13 and internal structures among populations from different habitats. The populations from
14 wastelands exhibited superior growth, with higher accumulation of soluble proteins (TSP) and
15 chlorophyll content (Chl *a&b*, TChl, Car, and Chl *a/b*). These populations displayed increased root
16 and stem area, storage parenchyma, vascular bundle area, metaxylem area, and phloem area.
17 Significant leaf modifications included thicker leaves, sclerification around vascular bundles, and
18 widened metaxylem vessels. Roadside populations possessed larger leaf area, enhanced
19 antioxidant activity, increased thickness of leaves in terms of midrib and lamina, and a higher
20 cortical proportion. Populations found in agricultural fields depicted enhanced shoot biomass
21 production, higher levels of chlorophyll b, and an increased total chlorophyll/carotenoid ratio.
22 Additionally, they exhibited increased phloem area in their roots, stems, and leaves, with a thick
23 epidermis only in the stem. All these outcomes of the study revealed explicit structural and
24 functional modifications among *P. hysterophorus* populations collected from different habitats.
25 These variations were attributed to the environmental variability and could contribute to the
26 widespread distribution of this species.

27 **Keyword:** *P. hysterophorus*, invasiveness, osmoregulation, surface hairiness, storage parenchyma,
28 ubiquitous

29

30 **1. Introduction**

31 Climate change poses an existential threat to global food security, ecosystems, and public health,
32 with atmospheric carbon dioxide (CO₂) levels projected to exceed 700 ppm by the century's end
33 and average temperatures increasing by 4 °C. These changes are expected to enhance the growth
34 and reproductive capabilities of many weeds and invasive plants, enabling them to compete more
35 effectively with crops and pastures (IPCC, 2014; Mao et al., 2021). Moreover, climate change may
36 reduce the efficacy of chemical herbicides and biological control agents. In addition to long-term
37 climate shifts, the increased frequency and intensity of extreme weather events, such as floods and
38 droughts, can disturb ground cover, create colonization opportunities, and facilitate weed dispersal
39 (Sun et al., 2020). The aggressive nature and potential impacts of parthenium weed raise concerns
40 about the effects of climate change, particularly rising atmospheric CO₂ levels and temperatures,
41 on its demography and competitive ability, as well as its management strategies. Increased
42 temperature and reduced humidity can negatively impact biological control agents, leading to
43 fluctuations in field population density (Hasan & Ansari, 2016). Furthermore, recent research
44 focusing on elevated CO₂ levels suggests the need to adjust current management approaches.
45 Therefore, a comprehensive review of parthenium weed's biology, ecology, and management
46 options under various climate change scenarios is crucial for informing future management
47 decisions and adapting strategies to address these emerging climate-induced challenges (Shabbir
48 et al., 2020; Mao et al., 2021).

49

50 In response to water scarcity and other stresses, plants adopt various survival strategies. They
51 increase root biomass and reduce shoot growth, along with making changes in leaf orientation,
52 size reduction, and shedding (Leukovic et al., 2009; Oliveira et al., 2018). At the anatomical level,
53 these plants exhibit reduced cell size, enlargement in vascular tissues, alterations in the
54 xylem/phloem ratio, and reductions in xylem and phloem vessel size (Makbul et al., 2011;
55 Boughalleb et al., 2014). Additionally, under drought or salinity stress, plants significantly reduce
56 xylem vessel diameter and increase the thickness of epidermis, phloem, and mesophyll tissues in
57 aerial parts (El Afry et al., 2012; Iqbal et al., 2023). They also accumulate substantial amounts of
58 protective compounds like glycine betaine, proline, and total soluble proteins to combat the
59 adverse effects of these abiotic stresses. Ionic homeostasis is a crucial physiological mechanism
60 in plants that contributes to their vitality and vigor even under harsh conditions (Siringam et al.,

61 2011). This mechanism involves processes such as noxious ion accumulation, selective ion uptake,
62 and excretion of toxic ions through specialized structures like leaf hairs, trichomes, leaf sheaths,
63 and excretory organs (Iqbal et al., 2022).

64 Parthenium weed (*Parthenium hysterophorus* L.) is a highly invasive plant species that has spread
65 across five continents, posing significant environmental, agricultural, and health threats.
66 Originating from the neotropical region, it has rapidly expanded its range due to accidental
67 introductions and unchecked trade. This invasive weed has invaded diverse ecosystems, including
68 grasslands, pastures, urban areas, and croplands, impacting biodiversity, and reducing livestock
69 and crop production (Adkins & Shabbir, 2014; Maharjan et al., 2020). Its competitive and
70 allelopathic effects have challenged farming systems' sustainability. Additionally, it directly
71 endangers human and livestock health. Efforts to control and manage parthenium weed are crucial
72 for safeguarding the environment, agriculture, and public well-being (Shabbir et al., 2013; Bajwa
73 et al., 2016). Parthenium weed exerts deleterious impacts on various cropping systems,
74 socioeconomic structures, and native ecosystems. The severity of these impacts is largely
75 attributed to its swift and effective invasion mechanism, as highlighted by research (Tanveer et al.
76 2015; Bajwa et al., 2016). However, a comprehensive understanding of this invasion mechanism
77 and its associated characteristics is currently lacking. To develop effective management strategies,
78 it is imperative to gain insights into how parthenium weed invades, as well as its interactions and
79 responses to biological and physical factors within invaded regions, which are essential for a more
80 robust ecological comprehension.

81 *Parthenium hysterophorus* L., is an annual herb known for its aggressive invasion of disturbed
82 lands and roadsides. While native to North America and Mexico, it has become an invasive species
83 in Pakistan. This plant exhibits notable characteristics, including strong competitiveness, high
84 drought tolerance, insensitivity to temperature fluctuations, and a remarkable capacity for seed
85 production. Its adaptability to diverse habitats makes it an invaluable tool for studying their
86 structural and functional responses to heterogenic environmental conditions. It was hypothesized
87 that the invasive success of *P. hysterophorus* in diverse habitats is influenced by its phenotypic
88 plasticity, allowing it to adapt to a wide range of environmental conditions. In this scenario, a
89 comprehensive study was aimed to answer the following research questions: a) how does *P.*
90 *hysterophorus* respond to heterogeneous environmental conditions at the levels of growth,

91 anatomy, and physiology? b) What types of micro-structural, physiological, and morphological
92 adaptations enable *P. hysterophorus* to mitigate the detrimental effects of prevailing stresses? By
93 examining the responses and modifications at different levels, the researchers sought to gain a
94 comprehensive understanding of the weed's ability to thrive in diverse environmental conditions.

95 **2. Materials and Methods**

96 **2.1 Study surveys, sampling and collection sites**

97 Sampling was done from distinct habitats of Punjab province to determine the growth,
98 physiological and anatomical response of *Parthenium hysterophorus* towards heterogenic
99 environmental conditions (Fig. 1 & 2, Table 1). Samples were collected during the peak of
100 flowering season (March to October) in 2022. Each study site was thoroughly searched in radius
101 of 1km and total 50 plants were ear marked. Ten plants (n=10) per population were finalized based
102 on growth habit, plant height, shoot length, leaf number and size, flowers and inflorescence for the
103 measurement of morpho-anatomical and physiological parameters. The populations were collected
104 from four prominent regions such as i) near wasteland (RYK-Rahim Yar Khan, SDK-Sadiqabad,
105 KHP-khanpur), ii) along water channels (BWP-Bahawalpur, LAP-Liaquatpur, AHP-Ahmadpur,
106 MUL-Multan), iii) along roadside (VEH-Vehari, DGK-DG Khan, RJP-Rajanpur, JHG-Jhang), iv)
107 near agriculture fields (MUZ-Muzaffargarh, SAR-Sargodha, FSD-Faisalabad, LYH-Layyah).
108 Coordinates were measured with the help of google positioning system (GPS, model: Garmin E-
109 Trex 20, GPS accuracy ± 1 m) (Table 2). Climatic data was taken from the meteorological
110 department situated in each district.

111 **2.2 Soil physiological parameters**

112 The soil texture was assessed using the USDA textural triangle, which categorizes soils into
113 distinct textural classes according to the relative proportions of sand, silt, and clay present in the
114 soil sample. The Walkley method (1947) was employed to measure the organic matter content
115 (OM) in the soil. This method involves oxidation of organic matter by dichromate in the presence
116 of sulfuric acid. A combined pH and ECe meter (WTW series InoLab pH/Cond 720, USA) was
117 used to measure the soil pH and electrical conductivity. Saturation paste prepared by saturating the
118 soil with water and extracting the solution, was used for these measurements. The saturation paste
119 was analyzed to determine the concentrations of different ions, including Na^+ , K^+ , and Ca^{2+} ,
120 utilizing a flame photometer (Jenway, PFP-7, UK). The nitrate content (NO_3^-) in the soil was

121 assessed using the micro-Kjeldahl method, which involves digesting the soil sample with sulfuric
122 acid. The resulting ammonia was then distilled and titrated using a semi-automatic ammonia
123 distillation unit (UDK-132, NIB-B (3)-DSU-003 Italy). The soil phosphate content (PO_4^{3-}) was
124 measured following the protocol described by Wolf (1982). This method typically involves
125 extracting the available phosphorus from the soil using a suitable extractant, followed by
126 colorimetric analysis. The chloride content in the soil was assessed using the Mohrs' titration
127 method (Mohrs, 1856). To determine the soil saturation percentage (SP), the soil samples were
128 dried in an oven at 70°C , and 200 g of the dried soil was used to prepare a composite saturation
129 paste, which was then analyzed. Saturation percentage assayed by following formula:

$$130 \quad SP (\%) = \frac{\text{Amount of water added}(g)}{\text{Oven dried soil } (g)} \times 100$$

131 Where SP % is saturation percentage.

132

133 **2.3 Morphological parameters**

134 To collect the necessary measurements, a meter rod was utilized to measure the plant, as well as
135 the length of shoot and root directly. A digital loading balance was employed to determine the fresh
136 weights of the shoot and root. Immediately after harvesting, the plant parts were weighed to obtain
137 their fresh weights. For dry weight analysis, the plant samples were subjected to oven-drying at a
138 temperature of 65°C until a constant weight was achieved. This ensured the complete removal of
139 moisture from the samples. The dry weights of the shoot and root were then measured using a
140 digital loading balance. To assess the leaf characteristics, the number of leaves on each plant was
141 counted. The leaf area was determined using cm-graph paper, providing a quantitative
142 measurement of the area occupied by the leaves. The leaf area was calculated using a formula
143 provided by Lopes et al. (2016).

144 **2.4 Physiological parameters**

145 2.4.1 Osmolytes and soluble proteins

146 Fresh samples were taken in falcon tubes and stored (at -80°C) for chlorophyll pigments,
147 osmoprotectants, and antioxidants activity. For the analysis of proline, fresh leaf samples were
148 thoroughly homogenized in sulfo-salicylic acid. Then was transferred into cuvette containing
149 ninhydrin solution. After subjected to water bath (100°C) toluene was added for extraction of

150 proline. Lastly, readings were taken on a spectrophotometer (Model 220, Hitachi, Japan) at 520
 151 nm wavelength (Bates et al., 1973).

$$152 \quad \text{Proline } (\mu\text{mol g}^{-1} \text{ fresh weight}) = \frac{\mu\text{g proline ml}^{-1} \times \text{ml of toluene}/115.5}{\text{sample weight (g)}}$$

153

154 To measure the glycine betaine content in the leaf samples, fresh leaf samples weighing 0.5 g were
 155 soaked in 20 ml of deionized water (H₂O) at a temperature of 25°C for a duration of 24 hours.
 156 Following the soaking period, an extract was prepared from the soaked samples and assayed using
 157 the established protocols outlined by Grattan & Grieve (1998). For the analysis of total soluble
 158 proteins, fresh leaf samples weighing 0.2 g were sliced and thoroughly crushed in 5 ml of
 159 phosphate buffer at a pH of 7.0. The buffer facilitated the extraction of proteins from the crushed
 160 leaf samples. The mixture of crushed leaf samples and buffer was then subjected to centrifugation
 161 at 5000 rpm for 5 minutes. This centrifugation step effectively separated the solid components of
 162 the mixture from the liquid supernatant. The supernatant, containing the soluble proteins, was
 163 collected for further analysis. To quantify the protein content in the supernatant, the method
 164 developed by Lowry et al. (1951) was employed. This method relies on a colorimetric assay to
 165 measure the protein concentration present in the sample

166 2.4.2 Photosynthetic parameters

167 To estimate the photosynthetic pigments, including chlorophylls (Chl a, Chl b, and TChl.) and
 168 carotenoids, the methods described by Arnon in 1949 and Davis in 1979 were followed. A
 169 spectrophotometer (Hitachi-220, Japan) was used for the measurements. The formulas used for
 170 calculations were:

171

$$172 \quad \text{Chl. a } (\text{mg g}^{-1} \text{ f.wt.}) = [12.7(\text{OD663}) - 2.69(\text{OD645})] \times \frac{V}{1000} \times W$$

$$173 \quad \text{Chl. b } (\text{mg g}^{-1} \text{ f.wt.}) = [22.9(\text{OD645}) - 4.68(\text{OD663})] \times \frac{V}{1000} \times W$$

$$174 \quad \text{Total chl. } (\text{mg g}^{-1} \text{ f.wt.}) = [20.2(\text{OD645}) - 8.02 (\text{OD663})] \times \frac{V}{1000} \times W$$

$$175 \quad \text{Carotenoids } (\text{mg g}^{-1} \text{ f.wt.}) = [12.7(\text{OD480}) - 0.114 (\text{OD663})] - 0.638 (\text{OD645})/2500$$

176 2.4.3 Total antioxidant activity

177 For the measurement of total antioxidant activity, a dried leaf sample weighing 0.5 g was placed
178 in a test tube. To facilitate the extraction of antioxidants from the leaf tissue, 20 mL of a 0.45%
179 salt solution was added to the test tube. The sample was then subjected to heating in a water bath
180 at 40°C for a duration of 20 minutes. After the heating process, the test tube was centrifuged at
181 3000 rpm for 30 minutes, enabling the separation of the supernatant from the solid residue. The
182 supernatant, which contained the extracted antioxidants, was carefully separated and stored at -
183 20°C until further analysis. To measure the total antioxidant activity, the FTC (Ferric Thiocyanate)
184 method described by Rahmat et al. (2003) was employed. This method involves assessing the
185 ability of the antioxidants to inhibit lipid peroxidation by reacting with ferric ions.

186 **2.5 Anatomical parameters**

187 To examine the anatomy of the root, stem, and leaf, the largest plant from each replicate was
188 selected. For leaf anatomy, a 2 cm section was obtained from the leaf base of fully mature and sun-
189 exposed leaves. For stem anatomy, a section was taken from the base of the internode of the main
190 stem. Similarly, for root anatomy, a section was obtained from tap root near the junction of the
191 root and shoot. The collected plant material was fixed using a formaldehyde acetic alcohol solution
192 consisting of 10% formaldehyde, 5% acetic acid, 50% ethanol, and 35% distilled water. The plant
193 material was immersed in the fixative solution for 48 hours, followed by transfer to an acetic
194 alcohol solution containing 25% acetic acid and 75% ethanol for long-term storage. To prepare the
195 sections for microscopic analysis, free-hand sections were made from the fixed plant material.
196 These sections underwent a series of dehydration steps using ethanol. For staining, the sections
197 were subjected to the standard safranin and fast green double-staining technique, as outlined by
198 Ruzin (1999). Measurements of the sections were taken using a light microscope (Nikon SE Anti-
199 Mould, Japan) equipped with an ocular micrometer that was calibrated using a stage micrometer.
200 Micrographs of the stained sections were captured using a digital camera (Nikon FDX-35)
201 mounted on a stereomicroscope (Nikon 104, Japan).

202 **2.6 Statistical analysis**

203 The morphological, physiological, and anatomical trait data were subjected to statistical analysis
204 using a One-way analysis of variance (ANOVA) in a complete randomized design with ten
205 replicates. Mean values were compared using the least significant difference (LSD) test at a
206 significance level of 5%. The statistical analysis was conducted using the Minitab software
207 package (version 17.1.0, Pennsylvania State University, USA). To examine the relationships

208 between the different morphological, physiological, and anatomical traits and the soil
209 physicochemical parameters of the collection sites, Principal Component Analysis (PCA) was
210 conducted. The analysis was carried out using the R-studio software, and the data were plotted to
211 visualize the patterns and associations. Furthermore, heatmaps were constructed using the
212 pheatmap package in R-studio. These heatmaps were used to cluster the selected groups based on
213 (i) soil physicochemical attributes and morphophysiological parameters, (ii) soil physicochemical
214 attributes and root anatomy, (iii) soil physicochemical attributes and stem anatomy, and (iv) soil
215 physicochemical attributes and leaf anatomy. The heatmaps provide a visual representation of the
216 relationships and similarities among the different variables.

217 **3. Results**

218 *3.1 Soil physicochemical characteristics*

219 The soil in most of the habitats was sandy (Table 2). The loamy soil was observed in five habitats
220 RYK (near the wasteland), KHP (near waste deposit), VEH (near the roadside), JHG (along rice
221 field) and LYH (wheat field) whereas loamy sand was observed in two habitats such as MUL
222 (along river Chenab) and RJP (near M5 motorway). Clayey loam was seen in DGK habitat (along
223 railway track). The soil electrical conductivity ranged from 0.76 to 6.73 dSm⁻¹, the maximum value
224 of soil ECe was recorded at RYK (near the wasteland) and KHP (near waste deposit) sites and the
225 minimum was observed at SDK (along barren land) and RJP (near M5 motorway). Habitats like
226 water channel (LAP), along Chenab river (MUL) and near GT road (JHG) showed exceptionally
227 highly level of soil ECe than rest of the populations. Most of the habitat comprised of alkaline pH,
228 ranging from 6.2 to 8.9. The acidic pH was observed only in one habitat RYK (near the wasteland).
229 The soil organic matter (OM) varied from 0.21 to 0.56%. The maximum organic matter was noted
230 in soil of Chenab river (MUL) and the minimum was measured in soil of roadside population
231 (VEH). The soil saturation percentage (SP) ranged from 15 to 42%. The maximum saturation
232 percentage was observed in soil of wheat filed (LYH) population. It was the minimum in soil of
233 water canal (LAP) and GT road (JHG) populations. The soil Phosphate concentration varied from
234 1.6 mg Kg⁻¹ in the SDK habitat to 3.6 mg Kg⁻¹ in the LYH habitat. The nitrate content in the LYH
235 habitat exhibited the highest value, while the MUZ habitat recorded the lowest value. The soil
236 chloride ion (Cl⁻) reached its maximum (567.8 mg Kg⁻¹) in the RYK habitat, while the minimum
237 (72.1 mg Kg⁻¹) was observed in both the DGK and MUL habitats. The soil calcium ion (Ca²⁺)

238 concentration ranged from 54.2 to 156.1 mg Kg⁻¹. The RYK habitat showed the highest soil
239 calcium concentration, while the SDK habitat exhibited the lowest concentration. The soil sodium
240 ion (Na⁺) ranged between 54.2 and 398.9 mg Kg⁻¹, with the RYK population having the highest
241 value and the SDK habitat recording the lowest. The maximum soil potassium ion (K⁺)
242 concentration was observed in the MUL and LYH habitats, while the minimum was found in the
243 SDK habitat.

244 **3.2 Growth characteristics**

245 Plant height was the maximum (56.5cm) in BWP population and the minimum (16.3 cm) in FSD
246 population (Fig. 2, Table 3). The maximum shoot length (44.7 cm) was recorded in BWP
247 population while the minimum (11.3 cm) of this parameter was noted in FSD population. Three
248 populations, KHP, VEH and SAR showed maximum shoot fresh (11.5 g plant⁻¹) and dry weight
249 (5.8 g plant⁻¹), while population FSD had least shoot fresh (3.0 g plant⁻¹) and dry weight (1.2 g
250 plant⁻¹). Root length was the maximum (11.5 cm) in BWP and the minimum (4.5 cm) in VEH
251 population. Four populations namely RYK, KHP, LAP and SAR showed maximum root fresh
252 weight (1.5 g plant⁻¹), while the population RJP exhibited low value of dry weight (0.4 g plant⁻¹).
253 Population RYK showed the maximum dry weight (1.2 g plant⁻¹) and populations BWP, AHP,
254 RJP and LYH possessed the minimum dry weight (0.2 g plant⁻¹). The maximum number of leaves
255 (29.5) were recorded in RYK population, while their minimum value (9.0) was observed in FSD
256 population. Two populations, BWP (65.3 cm²) and VEH (65.4 cm²) showed the maximum value
257 of leaf area, while the minimum (14.9 cm²) of that parameter was measured in RYK population.

258 **3.3 Physiological characteristics**

259 The population from RYK exhibited the highest total soluble protein content (47.9 µg g⁻¹ d.wt.),
260 while the population from VEH had the lowest (9.4 µg g⁻¹ d.wt.) (Table 3). Population BWP
261 showed the maximum proline content (19.8 µmol g⁻¹ d.wt.), whereas populations AHP and LYH
262 possessed the minimum (1.6 µmol g⁻¹ d.wt.). Glycine betaine content was highest in the BWP
263 population (10.2 µmol g⁻¹ d.wt.) and lowest in the FSD population (1.9 µmol g⁻¹ d.wt.). For
264 chlorophyll a content, the SDK population had the highest value (2.4 mg g⁻¹ f. wt.), while
265 populations MUL, DGK, RJP, JHG, and FSD had the lowest value (1.3 mg g⁻¹ f. wt.). Four
266 populations, SDK, JHG, MUZ, and FSD showed the highest chlorophyll b content (2.0 mg g⁻¹ f.
267 wt.), whereas the RYK population showed the lowest value (0.3 mg g⁻¹ f. wt.). The SDK
268 population had the maximum total chlorophyll content (4.4 mg g⁻¹ f. wt.), while the RYK and

269 DGK populations had the minimum ($2.1 \text{ mg g}^{-1} \text{ f. wt.}$). The LAP population had the highest
270 carotenoid content ($2.8 \text{ mg g}^{-1} \text{ f. wt.}$), and the MUZ population had the lowest ($1.0 \text{ mg g}^{-1} \text{ f. wt.}$).
271 The chlorophyll a/b ratio was highest in the RYK population (6.3) and lowest in the SAR
272 population (0.3). The MUZ population had the maximum total chlorophyll/carotenoid ratio (3.7),
273 whereas the VEH population had the minimum (0.3). Antioxidant activity was the maximum (9.9
274 %) in three populations, MUL, VEH and DGK, whereas it was the minimum (3.5%) in LAP
275 population.

276 ***3.4 Anatomical characteristics***

277 *3.4.1 Root anatomy*

278 The maximum root area ($400.4 \mu\text{m}^2$) was recorded in two populations, SAR and RYK, whereas
279 the minimum ($259.1 \mu\text{m}^2$) was in three populations such as KHP, BWP and FSD (Fig. 3, Table 4).
280 The population from MUL had the maximum epidermal thickness ($31.4 \mu\text{m}$), while the population
281 from LYH had the minimum epidermal thickness ($9.4 \mu\text{m}$). Population RYK showed the
282 maximum cortical thickness ($94.2 \mu\text{m}$), and population FSD had the smallest ($31.4 \mu\text{m}$). The
283 largest cortical cells ($41.1 \mu\text{m}^2$) were recorded in RYK and KHP populations, whereas the smallest
284 cells ($7.4 \mu\text{m}^2$) were seen in two populations, MUZ and SAR. Population BWP possessed the
285 largest vascular bundles ($121.3 \mu\text{m}^2$) than rest of the populations. On the other hand, population
286 MUL had smallest vascular bundles ($55.0 \mu\text{m}^2$). Three populations namely KHP, BWP and MUZ
287 exhibited widened metaxylem vessels ($15.7 \mu\text{m}^2$), whereas the populations of VEH and SAR had
288 the narrowest vessels ($9.4 \mu\text{m}^2$). Phloem area was the maximum ($2.5 \mu\text{m}^2$) in four populations,
289 KHP, LAP, MUZ and FSD, but the minimum ($0.5 \mu\text{m}^2$) was recorded in BWP and JHG.

290 *3.4.2 Stem anatomy*

291 The maximum value of stem area ($440.4 \mu\text{m}^2$) was observed in populations KHP and MUZ, while
292 their minimum value ($182.6 \mu\text{m}^2$) was noted in JHG (Fig. 4, Table 4). Epidermal thickness was
293 the maximum ($23.6 \mu\text{m}$) in SAR and KHP, and the minimum ($9.4 \mu\text{m}$) in RYK, MUL, RJP
294 and MUZ. Population KHP showed the highest cortical proportion ($70.7 \mu\text{m}$), whereas the
295 populations of RYK and MUL had lowest region ($18.8 \mu\text{m}$) of that character. Cortical cells area
296 was the maximum ($14.1 \mu\text{m}^2$) in population AHP, FSD and LYH, and the minimum ($6.3 \mu\text{m}^2$) was
297 in BWP. Population AHP and KHP showed largest vascular bundles ($164.9 \mu\text{m}^2$) as compared to
298 other populations, while populations of RYK, SDK and FSD represented smallest vascular regions
299 ($94.2 \mu\text{m}^2$). The largest metaxylem vessels ($18.8 \mu\text{m}^2$) were recorded in KHP and MUL, and the

300 smallest vessels ($9.4 \mu\text{m}^2$) were noted in BWP, VEH, MUZ and LYH populations. Phloem area
301 was the maximum ($69.1 \mu\text{m}^2$) in population LYH, and the minimum ($14.1 \mu\text{m}^2$) in SDK.

302 3.4.3 Leaf anatomy

303 Leaf thickness greatly varied in all populations of *P. hysterophorus* (Fig. 5, Table 4). Midrib
304 thickness was the maximum ($420.8 \mu\text{m}$) in SDK, and the minimum ($235.5 \mu\text{m}$) in FSD population.
305 The maximum value of lamina thickness ($38.1 \mu\text{m}$) was observed in population VEH, while the
306 minimum value ($11.0 \mu\text{m}$) was observed in RJP. Thicker epidermis ($23.6 \mu\text{m}$) was measured in
307 three populations, KHP, LAP and AHP, whereas the thinner ($10.6 \mu\text{m}$) of this parameter was noted
308 in MUL. Enhanced cortical region ($185.3 \mu\text{m}$) was observed in VEH, and their reduced (100.1
309 μm) was in FSD. The population from roadside habitats (VEH) exhibited the largest cortical cells
310 ($18.8 \mu\text{m}^2$), while the populations from FSD and LYH had the smallest cortical cells ($9.4 \mu\text{m}^2$).
311 The vascular bundle area was highest ($117.8 \mu\text{m}^2$) in the SDK population, whereas the RYK
312 population had the lowest vascular bundle area ($47.1 \mu\text{m}^2$). Among the populations, SDK had the
313 largest metaxylem vessels ($37.7 \mu\text{m}^2$), while BWP had the smallest ($10.9 \mu\text{m}^2$). The phloem area
314 was greatest (28.3) in the SAR population but was minimal ($11.8 \mu\text{m}^2$) in the DGK and AHP
315 populations.

316 **4. Multivariate analysis**

317 **4.1 Principal component analysis (PCA)**

318 Principal component analysis (PCA1) exhibited 27.4% and 21.2% (48.6%) variability among
319 morpho-physiological and soil physicochemical characteristics of *P. hysterophorus*. The Chl a,
320 TChl/Car, TChl, RDW, RFW, SFW and GB showed strong influence of soil NO_3 , SP, PO_4 and
321 pH, whereas Chl b, Chl a/b, TSP, SDW, SL, RL and LA represented least influence of soil OM
322 (Fig. 6a). Principal component analysis revealed significant influence of soil physiochemical
323 characters on anatomical traits of species. PCA2 represented the variability of 33.2% and 18.4%
324 (51.6%) among root anatomy and soil physicochemical attributes, as the CCA showed close
325 influence of soil Ca^{2+} , ECe, Cl^- and Na^+ , while MA, RA, CT, EpT, PhA and VBA had least
326 influence of soil OM (Fig. 6b). PCA3 indicated 36.5% and 18.9% (55.4%) variations between stem
327 anatomy and soil parameters, for example the MA represented very close influence of soil K and
328 NO_3 , whereas the CCA showed with soil ECe and VBA with soil pH (Fig.6c). PCA4 exhibited
329 28.8% and 19.9% (58.8%) variability amid leaf anatomy and soil attributes, as the LMT, MrT and

330 CCA showed strong influence of soil NO_3 , K^+ , SP and PO_4 , while the EpT with soil Ca^{2+} and Cl^- ,
331 and the VBA and PhA with soil pH (Fig. 6d).

332 **4.2 Clustered heatmaps**

333 Heatmap between soil physicochemical characters and morpho-physiological attributes exhibited
334 six major clusters (Fig. 7A). In the first cluster, soil attribute, OM form cluster with Ca and Car
335 content. The second cluster indicated the clustering of soil ECe, Cl and Na with LN, RFW and
336 RDW. In the third cluster, RL form cluster with chlb, Tchl, TSP and GB. The fourth group showed
337 clustering of soil attributes K, NO_3 , SP and PO_4 . The fifth cluster exhibited the clustering of LA
338 and soil pH, and the sixth cluster showed the clustering of Chl a/b, Chla and TChl/Car. The seventh
339 cluster indicated the clustering of SDW, SFW, PH and SL. Heatmap between root anatomical
340 characteristics and soil attributes indicated four major clusters (Fig. 7B). The first cluster indicates
341 the clustering of OM and MA. In the second cluster, soil pH form cluster with RA, CT, PhA, EpT
342 and VBA. In the third cluster, soil attributes like NO_3 , K, SP and PO_4 form clustering. In the
343 fourth cluster, soil Ca, Na, ECe and Cl showed clustering with CCA.

344 The heatmap between soil physicochemical attributes and stem anatomical features exhibited three
345 clusters (Fig. 7C). In the first cluster, soil pH and OM form clustering with VBA, EpT and PhA.
346 In the second cluster, soil ECe, Cl, Na and Ca show clustering with CCA and CT. The third cluster
347 indicated the clustering of K, PO_4 , SP and NO_3 with MA. Heatmap between soil physicochemical
348 attributes and leaf anatomical features exhibited four clusters (Fig. 7D). In the first cluster, soil
349 OM and pH form cluster with PhA and VBA, whereas in the second cluster, soil NO_3 , SP and PO_4
350 form cluster with LMT and CCA. The third cluster indicates the clustering of EPT and CT, while
351 the fourth cluster showing clustering of soil ECe, NA, Cl and Ca with MrT and MA.

352 **5. Discussion**

353 A summary of specific adaptive strategies of differently adapted populations of *Parthenium*
354 *hysterophorus* collected from different regions of Punjab province are highlighted in Figure 8. The
355 evaluation of morpho-anatomical and physio-biochemical adaptive markers is crucial for
356 understanding the underlying mechanisms of adaptation in differently adapted populations to
357 multiple stresses (Hameed et al., 2011; Nawaz et al., 2023). In the face of severe drought conditions
358 or physiological drought induced by other environmental stresses, water conservation becomes a
359 primary strategy (Sun et al., 2018). In water-scarce conditions, water conservation in plants is
360 achieved through mechanisms such as water storage in parenchymatous tissues like pith and cortex

361 (Alvarez et al., 2008; Iqbal et al., 2021), efficient water translocation facilitated by widening of
362 vessels, and reduction of water loss through the presence of mechanical tissues and a thick cuticle
363 on the surface of plant organs (Micco & Aronne, 2012). Herein, we tested the strength of
364 adaptation and the extent of these adaptations in plant survival, different populations of *P.*
365 *hysterophorus* were sampled from a wide range of habitats. It was hypothesized that the invasive
366 success of *P. hysterophorus* in diverse habitats is influenced by its phenotypic plasticity, allowing
367 it to adapt to a wide range of environmental conditions.

368 The investigation revealed significant variations in morphological characteristics among the
369 populations of *P. hysterophorus*, which can be attributed to the diverse environmental conditions
370 in which these populations were originally adapted. Under diverse conditions where the *P.*
371 *hysterophorus*, populations were collected, the genetically fixed characteristics of each population
372 were expressed, reflecting their adaptation to their respective habitats (Mojica et al., 2012; Paccard
373 et al., 2013). The population from the BWP site, which is located along a water channel with
374 relatively soft soil texture, exhibited the maximum growth (Table 3). This type of habitat seems to
375 be more favorable for the growth and development of *P. hysterophorus*, as reported for other
376 hydrophytes (Qadir et al., 2008; Hasanuzzaman et al., 2014). The compactness of the soil directly
377 influenced the growth and propagation of the species, with habitats consisting of compact soil
378 showing shorter plants, such as the FSD and VEH populations. Similar findings were reported by
379 Hamza & Anderson (2005), who observed shorter stature plants in compact soil. Biomass
380 production, both in roots and shoots, is a reliable criterion for assessing tolerance potential of a
381 species (Khosroshahi et al., 2014). The RYK and KHP populations demonstrated good overall
382 growth response, indicating their potential for stress tolerance. The SAR population also exhibited
383 vigorous growth, suggesting its complete adaptation to its specific habitat. Root and shoot
384 parameters, such as length, number, fresh and dry weights, have been previously associated with
385 abiotic stresses like drought or physiological drought in other plant species (Talukdar, 2013; Ye et
386 al., 2015). The RYK population displayed a high number of leaves per plant, although they were
387 smaller in size. Having a large number of leaves can enhance a plant's photosynthetic efficiency
388 (Weraduwege et al., 2015), while smaller leaves can increase water use efficiency by reducing
389 transpiration rates (Medrano et al., 2015). This adaptation is particularly important for survival in
390 harsh saline desert conditions.

391 Chlorophyll pigments serve as sensitive indicators of the metabolic state under salt stress
392 conditions (Chattopadhyay et al., 2011). In the present study, the least saline population SDK and
393 moderately saline population KHP showed an increase in chlorophyll *a*, chlorophyll *b*, total
394 chlorophyll, and carotenoid content. Similar findings have been reported by Amirjani (2011) on
395 rice *Triticum aestivum* L. and Sarabi et al. (2017) on melon (*Cucumis melo* L.). They noted an
396 augmentation in photosynthesis-related parameters under moderate salinity levels, but a decline
397 was observed under high salinity conditions. Conversely, the highly saline population RYK
398 exhibited lower amounts of chlorophyll pigments and carotenoids. This decrease in pigment
399 content aligns with other studies that have reported a significant reduction in photosynthetic
400 pigments under highly saline conditions, such as López-Millán et al. (2009) in *Lycopersicon*
401 *esculentum*, Peng et al. (2013) in *Elsholtzia splendens*, and Sytar et al. (2013) in various plant
402 species. In the present study, the BWP population showed an increasing trend in organic
403 osmolytes. The accumulation of osmolytes is an effective strategy employed by plants to endure
404 prevailing, which serves as a defensive mechanism for plants to maintain turgor pressure and
405 prevent tissue collapse due to desiccation (Kholodova et al., 2010; Sun et al., 2010). Elevated
406 levels of total antioxidant activity were observed in *P. hysterophorus* populations inhabiting
407 roadside areas, such as VEH and DGK. These findings align with previous studies conducted by
408 Nadgorska-Socha et al. (2013), Zemanova et al. (2013), and Almohisen (2014), which
409 demonstrated that dust pollution stimulates the production of various metabolites in plants. These
410 metabolites play a crucial role in mitigating stress by activating the plants' defense systems
411 (Sharma & Dietz, 2006).

412 The anatomical characteristics of plants have been recognized as highly responsive to climatic
413 conditions (Caemmerer & Evans, 2015; Iqbal et al., 2021). This adaptability enables plants to
414 thrive and survive in challenging environment (De Micco & Aronne, 2012). The size of the root
415 cross-sectional area is predominantly determined by the relative proportions of the cortical region
416 and the vascular bundle area (Table 4). An expansion in root area not only enhances the capacity
417 for water storage but also strengthens the mechanical integrity of the plant's soft tissues. This, in
418 turn, facilitates the efficient transportation of water and minerals from the roots to the aerial parts
419 of the plant, aided by physiological adjustments. The observed increase in root cross-sectional area
420 indicates better growth in the population inhabiting waste land (RYK). Roots, being underground
421 plant parts, are relatively less affected by environmental conditions compared to other plant organs

422 (Fitter & Hay, 2012). Epidermis is an outermost protective layer of roots, and under harsh
423 condition its strong friction of rhizospheric soil (McKenzie et al., 2013). In resulting, this may be
424 damaged, mainly in grasses and herbs (McCully, 1999). *P. hysterophorus* showed a significant
425 increase of this parameter in MUL population (along water channel). Thicker epidermal layers
426 play vital role in resisting the friction of soil compaction as well as impede the excessive water
427 and solute translocation inside root tissues (Chimungu et al., 2015). The water storage parenchyma
428 (cortex) and vascular region (metaxylem vessels and phloem) in the roots play a crucial role,
429 especially during water deficit or saline conditions. These adaptations are particularly significant
430 for the survival of arid zone species such as *P. hysterophorus* (Hsiao & Xu, 2000). A significantly
431 increased storage parenchyma and vascular region has been observed in populations of KHP (along
432 waste deposit) and MUZ (along agriculture field).

433 The plants growing in wastelands (KHP) demonstrated the highest values for the majority of stem
434 anatomical characteristics, as shown in Table 4. These characteristics encompass dermal, vascular,
435 and storage tissues, indicating favorable growth conditions and enhanced biomass production, as
436 evidenced by the shoot fresh weight. These findings are consistent with previous studies conducted
437 by Engloner (2009) and Guo & Miao (2010). The presence of sclerified tissues in the stems is a
438 notable adaptation to dry conditions (Nikolova & Vassilev, 2011). It was recorded in stems from
439 almost all habitats, but in populations from roadsides (VEH and DGK), there was higher lignin
440 deposition compared to the other populations. Under extreme dry and hot conditions, tissue
441 sclerification is beneficial for preventing from collapse of internally metabolically active tissues
442 during desiccation (da Cruz Maciel et al., 2015; Ahmad et al., 2016).

443 In arid zone species like *P. hysterophorus*, the leaf blade plays a vital role as it needs to withstand
444 harsh environmental conditions for the plant's survival. Among the studied populations, the plants
445 from roadside habitats (VEH) exhibited the highest values for various leaf anatomical
446 characteristics, including leaf thickness in terms of midrib and lamina thickness, as well as
447 mechanical and storage tissues such as cortical thickness and its cells area. These adaptations are
448 indicative of the plant's ability to protect the leaf blade from the challenging environmental
449 conditions encountered in roadside habitats (Ameer et al., 2023). Three populations namely KHP
450 (near wasteland), AHP (near Punjab Barrage) and LAP (along agriculture field) possessed thick
451 epidermis and sparse surface hairiness. Both are effective for evapo-transpiration loss when
452 population surviving in dry environmental condition (González et al., 2008; Sarwar et al., 2022).

453 Overall, these results suggest that phenotypic plasticity in structural and functional adaptations of
454 *P. hysterophorus* contribute to its resilience, competitive ability, and allowing it to adapt to a wide
455 range of environmental conditions, making it a successful and problematic invasive species.

456 **Conclusion**

457 In conclusion, *P. hysterophorus* displays significant variations in both structural and functional
458 attributes, enabling it to tolerate diverse environmental adversities. The wide distribution of this
459 species can be attributed to its specific adaptations along environmental gradients. It exhibits a
460 range of adaptations, including changes in growth parameters, microstructural features, and
461 functional traits. These adaptations, such as enhanced biomass production, long and numerous
462 roots, thicker epidermis, development of storage parenchyma tissues, lignification of cortical
463 region and vascular bundles, and increased levels of organic osmolytes and antioxidants. Overall,
464 the structural and functional adaptations of *P. hysterophorus* contribute to its resilience,
465 competitive ability, and ability to colonize a wide range of habitats, making it a successful and
466 problematic invasive species.

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470 ZU: The principal researcher responsible for conducting the experimental work.

471 UI: The principal supervisor of the second author, providing guidance in statistical analysis, data
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473 UI and KSA: Conducted a thorough review of the article to correct any language errors.

474 AA and HA: Contributed to the research by carrying out the practical aspects, including
475 biochemical analysis, anatomical photography, and data collection.

476 **Availability of Data and Material:**

477 All the data and relevant information is present in the manuscript.

478 **Declarations**

479 **Ethics Approval:** Since the study did not involve animal or human subjects, specific ethical
480 approval was not required. However, all necessary guidelines provided by The Islamia University
481 of Bahawalpur, Rahim Yar Khan Campus for handling plant material in the laboratory were strictly
482 adhered to. Following the completion of the study, proper measures were taken to dispose of all
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484 **Consent to Participate:** The contributions of all participants in this study have been duly
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Table 1 (on next page)

Table 1. Metrological record of differently adapted populations of star weed (*Parthenium hysterophorus* L.) collected from Punjab province

1 **Table 1. Habitat types and metrological record of differently adapted populations of star weed (*Parthenium hysterophorus* L.)**
 2 **collected from the Punjab province**

Ecological regions	Collection sites	Habitat types	Annual Temp. (°C)		Rainfall (mm)	Altitude (m.a.s.l)	Latitude (N)	Longitude (E)
			Max.	Min.				
Near wasteland	Rahim Yar Khan	Near the wasteland	44	13	115	88	28° 42' 12.29"	70° 29' 89.19"
	Sadiqabad	Along barren land	40	12	101	76	28° 09' 19.29"	70° 19' 12.99"
	Khanpur	Near waste deposit	43	15	110	184	32° 08' 51.27"	72° 38' 30.22"
Along water channel	Bahawalpur	Along the river Indus	44	13	179	149	31° 08' 41.23"	72° 08' 46.38"
	Liaquatpur	Along the water canal	34	14	119	237	32° 43' 19.02"	72° 58' 42.73"
	Ahmadpur	Near Punjab Barrage	40	16	142	212	30° 39' 31.63"	73° 23' 50.62"
	Multan	Along Chenab River	38	12	209	186	32° 17' 43.54"	72° 21' 03.24"
Along roadside	Vehari	Near the roadside	41	12	195	146	30° 55' 46.74"	71° 45' 41.90"
	DG Khan	Along railway track	40	11	143	198	28° 27' 42.58"	71° 03' 919.22"
	Rajanpur	Near M5 motorway	43	12	120	117	28° 46' 04.86"	71° 20' 03.13"
	Jhang	Near GT road	40	10	155	267	29° 58' 01.03"	70° 19' 36.63"
Near agriculture field	Muzaffargarh	Near cotton field	44	13	176	210	32° 25' 30.62"	37° 13' 31.40"
	Sargodha	Along sorghum field	38	9	246	192	31° 28' 42.68"	73° 12' 36.66"
	Faisalabad	Along Rice field	40	10	201	140	29° 20' 05.33"	71° 56' 04.29"
	Layyah	Wheat field	43	12	130	288	32° 24' 45.54"	71° 58' 00.51"

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Table 2 (on next page)

Soil physicochemical parameters of collection sites of differently adapted populations of star weed (*Parthenium hysterophorus* L.) collected from Punjab province

1 **Table 2. Soil physicochemical parameters of collection sites of differently adapted populations of star weed (*Parthenium***
 2 ***hysterophorus* L.) collected from the Punjab province**

Ecological regions	Collection sites	Soil texture	ECe (dS m ⁻¹)	pH	OM (%)	SP (%)	PO ₄ ³⁻ (mg UKg ⁻¹)	NO ₃ ⁻ (mg UKg ⁻¹)	Cl ⁻ (mg UKg ⁻¹)	Ca ²⁺ (mg UKg ⁻¹)	Na ⁺ (mg UKg ⁻¹)	K ⁺ (mg UKg ⁻¹)
Near wasteland	RYK	Loamy	6.73 ^a	6.2 ^j	0.35 ^e	36 ^c	3.1 ^b	3.3 ^c	567.8 ^a	156.1 ^a	398.9 ^a	64.4 ⁱ
	SDK	Sandy	0.76 ^h	8.4 ^g	0.42 ^d	16 ^{gh}	1.6 ^g	2.9 ^d	83.4 ^h	54.2 ^g	54.2 ⁱ	70.1 ⁱ
	KHP	Loamy	6.69 ^a	8.8 ^b	0.28 ^g	38 ^b	3.4 ^{ab}	4.0 ^b	434.5 ^b	67.7 ^{ef}	297.1 ^b	260.3 ^b
Along water channel	BWP	Sandy	0.96 ^g	8.7 ^c	0.28 ^g	17 ^{gh}	1.9 ^d	2.9 ^d	102.7 ^g	71.9 ^e	147.8 ^f	80.8 ^g
	LAP	Sandy	3.46 ^c	8.0 ^h	0.35 ^e	15 ^h	2.2 ^c	3.2 ^c	389.1 ^c	97.3 ^c	297.1 ^b	180.9 ^d
	AHP	Sandy	1.06 ^f	8.6 ^d	0.42 ^d	16 ^{gh}	1.9 ^d	3.5 ^c	130.5 ^f	63.5 ^f	164.1 ^e	148.5 ^e
	MUL	Loamy sand	4.33 ^b	7.8 ⁱ	0.56 ^a	22 ^d	2.2 ^c	3.2 ^c	72.1 ^j	60.2 ^f	266.1 ^c	276.3 ^a
Along roadside	VEH	Loamy	1.15 ^e	8.2 ^f	0.21 ^h	32 ^d	3.1 ^b	4.3 ^b	109.8 ^g	78.7 ^d	180.7 ^d	124.1 ^f
	DGK	Clayey loam	1.19 ^e	8.7 ^b	0.28 ^g	38 ^b	3.4 ^{ab}	4.0 ^b	72.1 ^j	66.7 ^{ef}	60.8 ^h	258.3 ^b
	RJP	Loamy sand	0.90 ^g	8.5 ^e	0.26 ^g	16 ^{gh}	1.8 ^d	2.8 ^d	100.7 ^g	70.9 ^e	145.8 ^f	79.8 ^g
	JHG	Loamy	3.01 ^c	8.0 ^h	0.35 ^e	15 ^h	2.2 ^c	3.2 ^c	389.1 ^c	97.3 ^c	61.8 ^h	180.9 ^d
Near agriculture field	MUZ	Sandy	1.33^d	8.2^f	0.28^g	17^{gh}	1.9^d	2.0^c	178.6^c	110.9^b	134.0^g	80.1^g
	SAR	Sandy	0.77^h	8.5^e	0.45^b	18^f	1.9^d	4.0^b	79.6ⁱ	77.1^d	175.0^d	75.2^h
	FSD	Sandy	1.08^f	8.7^c	0.43^{bd}	19^e	2.3^c	4.0^b	111.6^g	104.3^{bc}	147.1^f	196.6^c
	LYH	Loamy	1.20^{de}	8.9^a	0.31^f	42^a	3.6^a	5.1^a	198.3^d	94.3^c	88.9^g	276.8^a
	LSD		0.5	1.0	0.5	6.0	0.5	1.0	7.0	6.0	25.8	6.0

3 Means shearing similar letter in each row are not statistically significant.

4 * = Significant at $P < 0.05$, ** = Significant at $P < 0.01$, *** = Significant at $P < 0.001$, NS = not significant

5 **Abbreviations are given as footnote of Figure 7&8.**

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Table 3(on next page)

Growth and physiological attributes of differently adapted populations of star weed (*Parthenium hysterophorus* L.) collected from Punjab province

Abbreviations are given at start of manuscript. Means shearing similar letters in each row are statistically not significant *, significant at $P < 0.05$; **, significant at $P < 0.01$; ***, significant at $P < 0.001$; NS, not significant

1 **Table 3. Growth and physiological attributes of differently adapted populations of star weed (*Parthenium hysterophorus* L.)**
 2 **collected from the Punjab province**

Ecological regions	Near wasteland				Along water channel				Along roadside				Near agriculture field				
Collection sites	RYK	SDK	KHP	BWP	LAP	AHP	MUL	VEH	DGK	RJP	JHG	MUZ	SAR	FSD	LYH	LSD	F-value
Growth attributes																	
Plant height (cm)	37.0 ^d	46.0 ^b	40.3 ^c	56.5 ^a	45.0 ^b	31.0 ^c	39.5 ^c	41.2 ^c	37.3 ^d	20.3 ^f	37.7 ^d	36.3 ^d	43.0 ^c	16.3 ^g	30.0 ^e	11.6	72.6***
Shoot length (cm)	30.0 ^d	40.3 ^b	31.0 ^d	44.7 ^a	35.0 ^c	26.0 ^e	33.0 ^{cd}	36.3 ^c	30.4 ^d	16.3 ^f	28.0 ^d	30.0 ^d	33.0 ^{cd}	11.3 ^g	24.0 ^e	4.5	19.4***
Shoot fresh weight (g plant ⁻¹)	6.3 ^d	8.2 ^{bc}	11.5 ^a	9.4 ^b	8.2 ^{bc}	5.4 ^{de}	7.7 ^c	11.0 ^a	4.5 ^e	4.5 ^e	9.4 ^b	4.7 ^c	11.5 ^a	3.0 ^f	4.4 ^{de}	2.2	14.9***
Shoot dry weight (g plant ⁻¹)	3.1 ^c	4.1 ^b	5.8 ^a	4.7 ^b	4.1 ^b	2.7 ^{cd}	3.9 ^b	5.8 ^a	2.2 ^d	2.0 ^d	4.7 ^b	2.3 ^d	5.8 ^a	1.2 ^c	2.0 ^{cd}	1.4	11.8**
Root length (cm)	8.0 ^b	6.0 ^c	8.0 ^b	11.5 ^a	10.0 ^{ab}	7.7 ^b	5.7 ^c	4.5 ^d	7.3 ^{bc}	7.7 ^b	10.0 ^{ab}	6.0 ^c	10.0 ^{ab}	6.0 ^c	7.2 ^b	1.8	31.7***
Root fresh weight (g plant ⁻¹)	1.5 ^a	0.5 ^{bc}	1.5 ^a	0.7 ^b	1.5 ^a	0.6 ^c	0.7 ^b	0.7 ^b	0.7 ^b	0.4 ^d	0.6 ^c	0.8 ^b	1.5 ^a	0.5 ^{bc}	0.5 ^c	1.0	68.8***
Root dry weight (g plant ⁻¹)	1.2 ^a	0.3 ^c	1.0 ^{ab}	0.2 ^f	1.0 ^{ab}	0.2 ^f	0.3 ^c	0.4 ^d	0.5 ^c	0.2 ^f	0.3 ^c	0.5 ^c	1.0 ^{ab}	0.3 ^c	0.2 ^f	0.5	86.1***
Leaf number (per branch)	29.5 ^a	18.5 ^d	25.5 ^b	17.0 ^d	19.0 ^d	10.5 ^{ef}	14.0 ^c	22.0 ^c	15.5 ^c	14.5 ^c	18.5 ^d	11.0 ^{ef}	20.5 ^c	9.0 ^f	9.5 ^{ef}	4.3	25.7***
Leaf area (cm ²)	14.9 ^k	53.4 ^c	19.2 ⁱ	65.5 ^a	38.4 ^e	39.2 ^c	23.9 ^j	65.4 ^a	38.0 ^e	16.6 ^h	27.6 ^g	47.9 ^d	59.3 ^b	33.7 ^f	37.2 ^c	9.8	8.3**
Physiological attributes																	
Total soluble protein (µg g ⁻¹ d.wt.)	47.9 ^a	26.4 ^f	21.7 ^g	41.9 ^b	20.8 ^g	23.1 ^g	22.8 ^g	9.4 ⁱ	32.0 ^d	29.8 ^c	35.7 ^c	19.3 ^h	36.3 ^c	24.7 ^g	22.1 ^g	6.7	33.3***
Proline (µmol g ⁻¹ dwt.)	8.8 ^c	8.8 ^c	7.0 ^d	19.8 ^a	5.9 ^c	1.6 ^g	3.5 ^f	3.6 ^f	6.7 ^{de}	10.4 ^b	10.8 ^b	8.9 ^c	7.9 ^c	8.5 ^c	1.6 ^g	9.1	39.1***
Glycine betaine (µmol g ⁻¹ dwt.)	3.6 ^b	3.8 ^b	2.6 ^c	10.2 ^a	2.2 ^c	2.5 ^c	2.4 ^c	2.1 ^c	2.4 ^c	3.1 ^b	2.4 ^c	2.6 ^c	3.9 ^b	1.9 ^d	2.3 ^c	4.5	35.0***
Chlorophyll a (mg g ⁻¹ f. wt.)	1.9 ^c	2.4 ^a	1.9 ^c	1.9 ^c	1.7 ^{cd}	2.2 ^b	1.3 ^d	1.7 ^{cd}	1.3 ^d	1.3 ^d	1.3 ^d	1.7 ^{cd}	2.2 ^b	1.3 ^d	2.0 ^b	1.3	52.8***
Chlorophyll b (mg g ⁻¹ f. wt.)	0.3 ^f	2.0 ^a	0.7 ^c	1.8 ^{ab}	1.0 ^d	1.7 ^{ab}	1.8 ^{ab}	1.8 ^{ab}	0.8 ^e	1.5 ^c	2.0 ^a	2.0 ^a	1.3 ^c	2.0 ^a	1.6 ^{ab}	1.0	40.2***
Total chlorophyll (mg g ⁻¹ f. wt.)	2.1 ^f	4.4 ^a	2.6 ^d	3.7 ^{ab}	2.7 ^d	3.9 ^{ab}	3.1 ^c	3.5 ^b	2.1 ^f	2.8 ^c	3.3 ^b	3.7 ^{ab}	3.5 ^b	3.3 ^b	3.7 ^{ab}	1.5	60.5***
Carotenoids (mg g ⁻¹ f. wt.)	1.5 ^d	2.4 ^b	1.4 ^d	1.4 ^d	2.8 ^a	1.8 ^c	1.8 ^c	2.6 ^{ab}	1.8 ^c	1.7 ^c	1.9 ^c	1.0 ^c	2.5 ^b	1.7 ^c	1.6 ^c	1.1	18.8***
Chlorophyll a/b	6.3 ^a	1.2 ^d	2.7 ^b	1.0 ^e	1.7 ^c	1.2 ^d	0.7 ^f	0.9 ^f	1.6 ^c	0.8 ^f	0.6 ^g	0.8 ^f	0.3 ^h	0.6 ^g	1.0 ^d	0.5	73.1***
Total Chlorophyll/Carotenoid	1.4 ^e	3.1 ^{ab}	1.0 ^f	2.6 ^c	1.3 ^e	0.9 ^g	1.7 ^d	0.3 ^h	1.1 ^f	1.6 ^d	1.7 ^d	3.7 ^a	1.4 ^c	1.9 ^d	0.7 ^g	0.9	89.2***
Antioxidant activity (%)	5.0 ^d	5.4 ^d	4.2 ^c	5.2 ^d	3.5 ^f	6.5 ^c	9.9 ^a	9.9 ^a	9.9 ^a	6.1 ^c	9.3 ^{ab}	5.7 ^d	6.2 ^c	7.8 ^{bc}	6.4 ^c	3.3	36.4***

3 Abbreviations are given at start of manuscript. Means shearing similar letters in each row are statistically not significant

4 *, significant at $P < 0.05$; **, significant at $P < 0.01$; ***, significant at $P < 0.001$; NS, not significant

5 **Abbreviations are given as footnote of Figure 7&8.**

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Table 4(on next page)

Anatomical characteristics of differently adapted populations of star weed (*Parthenium hysterophorus* L.) collected from Punjab province

Abbreviations are given at the start of manuscript. Means shearing similar letters in each row are statistically not significant. *, significant at $P < 0.05$; **, significant at $P < 0.01$; ***, significant at $P < 0.001$; NS, not significant

1 **Table 4. Anatomical characteristics of differently adapted populations of star weed (*Parthenium hysterophorus* L.) collected**
 2 **from the Punjab province**

Ecological regions	Near wasteland				Along water channel				Along roadside				Near agriculture field				F-ratio
Collection sites	RYK	SDK	KHP	BWP	LAP	AHP	MUL	VEH	DGK	RJP	JHG	MUZ	SAR	FSD	LYH	LSD	F-ratio
Root anatomy																	
Root area (μm^2)	400.4 ^a	282.6 ^c	259.1 ^f	259.1 ^f	306.2 ^d	304.6 ^d	306.2 ^d	306.2 ^d	353.3 ^b	329.7 ^c	282.6 ^b	329.7 ^c	400.4 ^a	259.1 ^g	353.3 ^b	10.9	66.5 ^{***}
Epidermal thickness (μm)	18.8 ^c	17.3 ^c	12.6 ^{de}	18.8 ^c	22.0 ^b	17.3 ^c	31.4 ^a	17.3 ^c	14.1 ^d	15.7 ^d	22.0 ^b	15.7 ^d	22.0 ^b	14.1 ^d	9.4 ^e	5.4	47.3 ^{***}
Cortical thickness (μm)	94.2 ^a	37.7 ^e	51.8 ^d	45.5 ^d	55.0 ^c	37.7 ^e	55.0 ^c	65.9 ^b	55.0 ^c	67.5 ^b	55.0 ^c	36.1 ^e	55.0 ^c	31.4 ^f	65.9 ^b	8.9	98.6 ^{***}
Cortical cell area (μm^2)	14.1 ^a	9.4 ^c	14.1 ^a	11.0 ^b	9.4 ^c	9.4 ^c	9.4 ^c	11.0 ^b	9.4 ^c	9.4 ^c	9.4 ^c	7.4 ^d	7.4 ^d	9.4 ^c	9.4 ^c	2.5	86.4 ^{***}
Vascular bundle areas (μm^2)	70.7 ^c	94.2 ^b	70.7 ^c	121.3 ^a	69.1 ^c	94.2 ^b	55.0 ^c	70.7 ^c	65.9 ^d	67.5 ^d	70.7 ^c	70.7 ^c	67.5 ^d	70.7 ^c	69.1 ^c	20.3	72.4 ^{***}
Metaxylem area (μm^2)	12.6 ^c	11.0 ^d	15.7 ^a	15.7 ^a	12.6 ^c	11.0 ^d	12.6 ^c	9.4 ^e	12.6 ^c	11.0 ^d	14.1 ^b	15.7 ^a	9.4 ^e	12.6 ^c	14.1 ^b	4.3	85.8 ^{***}
Phloem area (μm^2)	1.0 ^c	1.8 ^b	2.5 ^a	0.5 ^d	2.5 ^a	1.9 ^b	1.8 ^b	1.0 ^c	1.0 ^c	1.9 ^b	0.5 ^d	2.5 ^a	1.7 ^b	2.5 ^a	1.9 ^b	1.1	19.9 ^{***}
Stem anatomy																	
Stem area (μm^2)	229.1 ^g	282.6 ^c	440.4 ^a	259.1 ^f	290.2 ^d	290.6 ^d	290.2 ^d	290.2 ^d	343.3 ^b	300.7 ^c	182.6 ^b	440.4 ^a	259.1 ^f	300.7 ^c	340.3 ^b	32.2	35.6 ^{***}
Epidermal thickness (μm)	9.4 ^d	14.1 ^c	23.6 ^a	14.1 ^c	14.1 ^c	18.8 ^b	9.4 ^d	18.8 ^b	14.1 ^c	9.4 ^d	14.1 ^c	9.4 ^d	23.6 ^a	14.1 ^c	14.1 ^c	6.5	21.4 ^{***}
Cortical thickness (μm)	18.8 ^b	23.6 ^g	70.7 ^a	33.0 ^f	55.0 ^c	47.1 ^d	18.8 ^b	47.1 ^d	67.5 ^{ab}	47.1 ^d	47.1 ^d	39.3 ^e	59.7 ^b	47.1 ^d	47.1 ^d	12.4	37.6 ^{***}
Cortical cell area (μm^2)	12.6 ^b	9.4 ^c	11.9 ^b	6.3 ^d	14.1 ^a	9.4 ^c	9.4 ^c	9.4 ^c	11.0 ^b	9.4 ^c	11.0 ^b	9.4 ^c	9.4 ^c	14.1 ^a	14.1 ^a	3.1	19.5 ^{***}
Vascular bundle area (μm^2)	94.2 ^b	94.2 ^h	164.9 ^b	131.9 ^e	108.3 ^g	164.9 ^a	146.0 ^c	117.8 ^f	146.0 ^c	149.2 ^b	128.7 ^e	117.8 ^f	133.5 ^d	94.2 ^b	133.5 ^d	12.7	49.4 ^{***}
Metaxylem area (μm^2)	12.6 ^b	15.7 ^b	17.3 ^a	9.4 ^c	14.1 ^b	14.1 ^b	18.8 ^a	9.4 ^c	14.1 ^b	14.1 ^b	14.1 ^b	9.4 ^c	14.1 ^b	14.1 ^b	9.4 ^c	4.4	87.3 ^{***}
Phloem area (μm^2)	20.4 ^f	14.1 ^g	36.1 ^e	40.8 ^d	47.1 ^c	48.7 ^c	45.5 ^c	47.1 ^c	58.1 ^b	47.1 ^c	47.1 ^c	42.4 ^d	47.1 ^c	58.1 ^b	69.1 ^a	15.8	52.3 ^{***}
Leaf anatomy																	
Midrib thickness (μm)	379.9 ^c	420.8 ^a	376.8 ^c	337.6 ^d	329.7 ^e	235.5 ^d	329.7 ^e	389.4 ^b	376.8 ^c	329.7 ^e	329.7 ^e	329.7 ^e	282.6 ^f	235.5 ^g	282.6 ^f	12.9	18.5 ^{***}
Lamina thickness (μm)	22.0 ^c	14.1 ^c	18.8 ^d	14.1 ^c	22.0 ^c	17.3 ^d	18.8 ^d	38.1 ^a	17.3 ^d	11.0 ^f	14.1 ^c	14.1 ^c	28.3 ^b	26.7 ^{bc}	14.1 ^c	6.4	73.8 ^{***}
Epidermal thickness (μm)	18.8 ^b	15.7 ^c	23.6 ^a	14.1 ^d	23.6 ^a	23.6 ^a	10.6 ^e	15.7 ^c	14.1 ^c	12.6 ^{de}	17.3 ^b	17.3 ^b	15.7 ^c	18.8 ^b	15.7 ^c	3.3	89.4 ^{***}
Cortical thickness (μm)	117.8 ^h	106.8 ⁱ	141.3 ^c	180.6 ^{ab}	139.7 ^f	117.8 ^h	150.7 ^d	185.3 ^a	153.9 ^d	158.6 ^c	127.2 ^g	122.5 ^g	139.7 ^f	100.1 ^j	119.3 ^h	17.6	36.3 ^{***}
Cortical cell area (μm^2)	14.1 ^b	14.1 ^b	15.7 ^b	15.7 ^b	12.6 ^c	11.0 ^c	15.7 ^b	18.8 ^a	11.0 ^c	14.1 ^b	14.1 ^b	18.8 ^a	14.1 ^b	9.4 ^d	9.4 ^d	4.4	72.7 ^{***}
Vascular bundle area (μm^2)	47.1 ^f	117.8 ^a	92.6 ^c	83.2 ^d	70.7 ^e	69.1 ^e	69.1 ^e	70.7 ^e	83.2 ^d	97.3 ^b	69.1 ^e	92.6 ^c	70.7 ^e	70.7 ^e	83.2 ^d	12.5	19.8 ^{***}
Metaxylem area (μm^2)	18.8 ^c	37.7 ^a	29.8 ^b	10.9 ^g	17.3 ^e	16.2 ^{ef}	14.8 ^f	22.0 ^d	22.0 ^d	20.4 ^e	25.1 ^c	17.3 ^e	18.8 ^c	14.1 ^f	22.0 ^d	8.0	14.5 ^{**}
Phloem area (μm^2)	15.7 ^c	23.6 ^b	20.4 ^{bc}	16.0 ^c	14.1 ^c	11.8 ^d	16.0 ^c	20.4 ^{bc}	11.8 ^d	22.0 ^{bc}	23.6 ^b	20.4 ^{bc}	28.3 ^a	15.7 ^c	20.4 ^{bc}	5.2	11.3 ^{**}

3 Abbreviations are given at the start of manuscript. Means shearing similar letters in each row are statistically not significant.

4 *, significant at $P < 0.05$; **, significant at $P < 0.01$; ***, significant at $P < 0.001$; NS, not significant

5 Abbreviations are given as footnote of Figure 7&8.

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Figure 1

Map of Punjab showing collection sites of *Parthenium hysterophorus* L. sampled from different districts

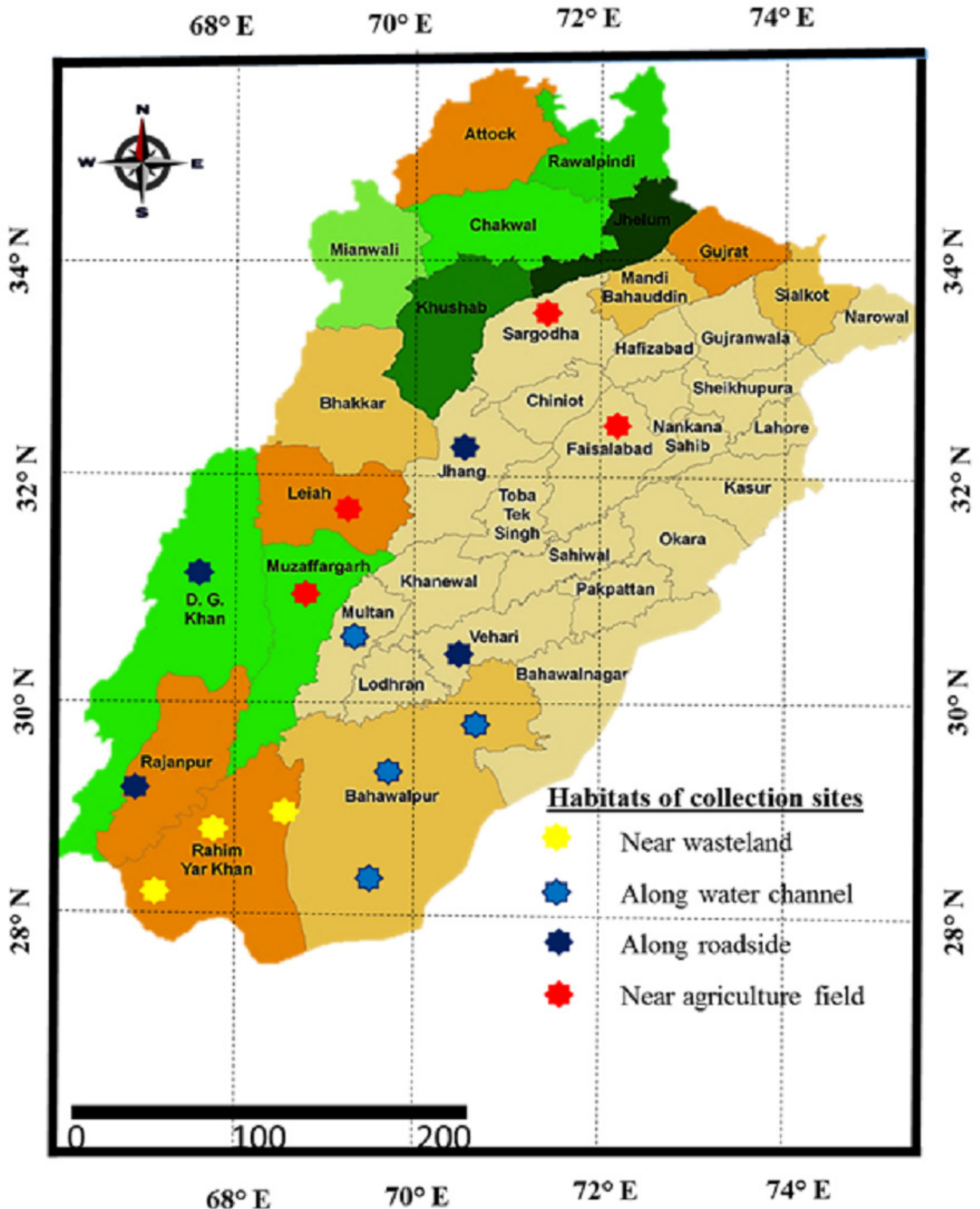


Figure 2

Habitat view and description of *Parthenium hysterophorus* L. populations collected from different ecological regions

a. RYK-Rahim Yar Khan: Saline patches in the Cholistan Desert with hot and dry environment, **b. SDK-Sadiqabad:** Open and barren land characterized by saline soil, **c. KHP-Khanpur:** Soil is sandy, climate dry and hot. *Eucalyptus* is common genus, **d. BWP-Bahawalpur:** Dominated along the bank of Indus River. Soil is sandy and hot climate, **e. LAP-Liaquatpur:** Growing along the water channel in patches form, soil is sandy, **f. AHP-Ahmadpur:** Vicinity of Punjab Barrage, sandy soil dominated with hydrophytes, **g. MUL-Multan:** Green belt of Punjab covered by various bushes and trees like Mango, **h. VEH-Vehari:** Dry and hot region characterized by loamy soil, **i. DGK-Dera Ghazi Khan:** Foothills of Suleiman Mountains, climate cool in winters and very hot in summers, **j. RJP-Rajanpur:** Desert flats characterized by hot climate and loamy sand. *Capparis* and *Salvadora* are commonly found, **k. JHG-Jhang:** Sandy soil of the Thal Desert characterized by small sand dunes, climate very hot and dry, **l. MUZ-Muzaffargarh:** Lush green region of the Punjab province dominated by various crops and vegetation, **m. SAR-Sargodha:** Hot and dry climate having sandy soil. *Calotropis*, *Prosopis* and *Acaica* are the common species, **n. FSD-Faisalabad:** It is characterized by sandy soil and hot climate, where rice and wheat are commonly cultivated crops, **o. LYH-Layyah:** Flats of Thal Desert, characterized by chickpea plantation on large scale, **p. Inflorescence:** The inflorescences are grouped into 4 or 5, in small globular heads.



Figure 3

Root transvers sections of *Parthenium hysterophorus* L. populations collected from different ecological regions

a. RYK-Rahim Yar Khan. Thicker epidermis, enlarge cortical region and metaxylem vessels, **b. SDK-Sadiqabad.** Reduced root cellular area, cortical thickness, metaxylem vessels and slightly crushed, **c. KHP-Khanpur.** Reduced root area and epidermal thickness, enhanced metaxylem vessels **d. BWP-Bahawalpur.** Greatly enhanced epidermal thickness, cortical thickness and metaxylem area, **e. LAP-Liaquatpur.** Extraordinarily thicker epidermis, cortical region and metaxylem vessels, **f. AHP-Ahmadpur.** Extraordinarily thick cortical region and enlarge metaxylem vessels, **g. MUL-Multan.** Thick epidermis and cortical region, enhanced metaxylem area, **h. VEH-Vehari.** Thicker epidermis, partially crushed cortical region and enlarge xylem vessels, **i. DGK- Dera Ghazi Khan.** Thick epidermis and cortical region, reduced xylem vessels, **j. RJP-Rajanpur.** Greatly reduced root cellular region and cortical thickness and metaxylem area, **k. JHG-Jhang.** Reduced root area, cortical region and metaxylem vessels, **l. MUZ-Muzaffargarh.** Reduced cortical thickness and partially crushed cortical region, **m. SAR-Sargodha.** Thick epidermis, enlarge metaxylem vessels and cortical region, **n. FSD-Faisalabad.** Thick cortical region and reduced xylem vessels, **o. LYH-Layyah.** Reduced cortical region and metaxylem area.

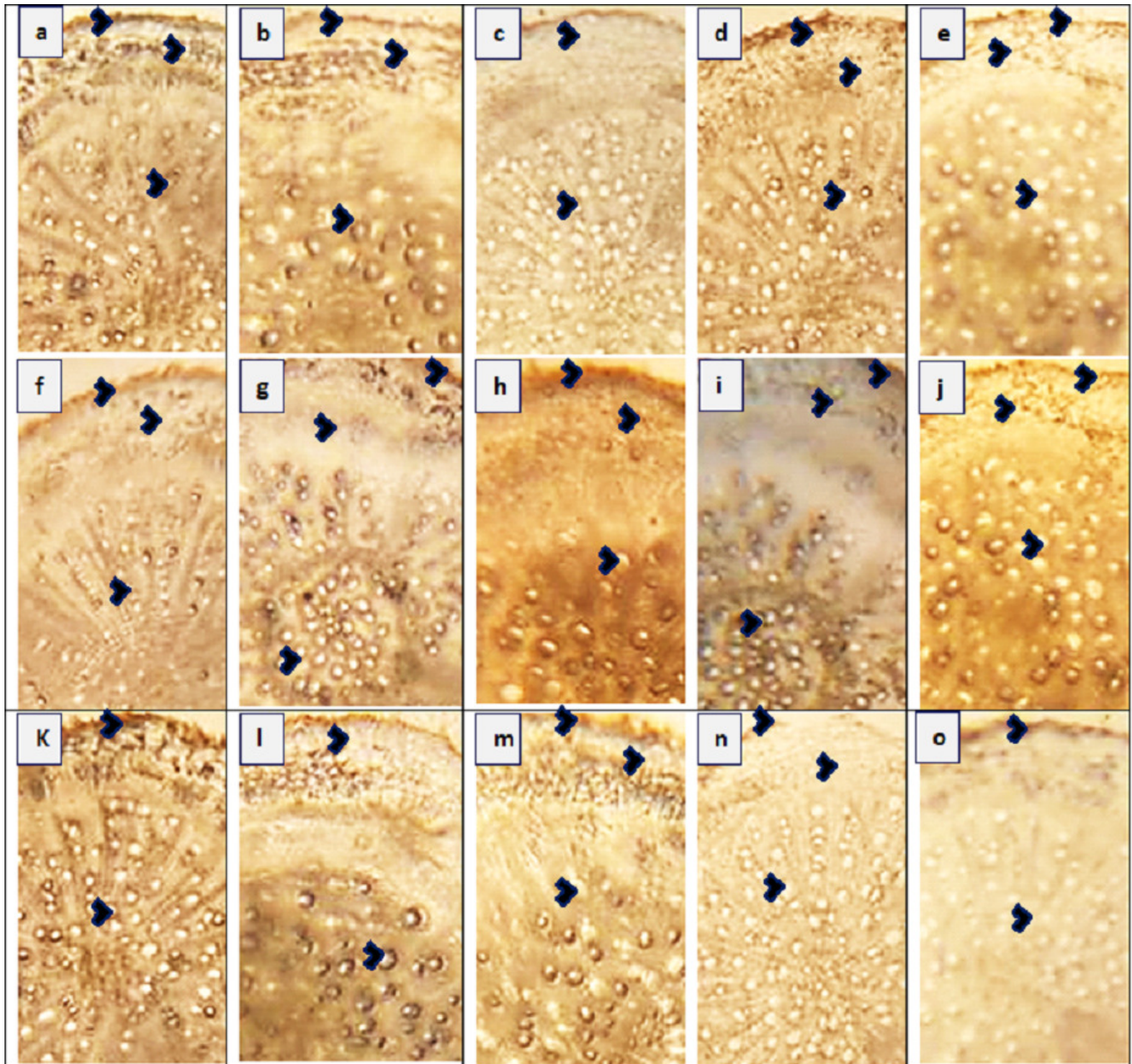


Figure 4

Stem transvers sections of *Parthenium hysterophorus* L. populations collected from different ecological regions

Description: **a. RYK-Rahim Yar Khan.** Thicker epidermis, enlarge cortical region and vascular bundles, **b. SDK-Sadiqabad.** Reduced stem cellular area, cortex thickness, metaxylem vessels and vascular bundle area, **c. KHP-khanpur.** Enlarge stem area, enhanced cortical and epidermal thickness, sparse hairiness on surface **d. BWP-Bahawalpur.** Greatly enhanced epidermal thickness, cortical and pith thickness, and vascular bundle area, **e. LAP-Liaqatpur.** Extraordinarily thick cortical region, vascular and pith region, thick surface pubescence, **f. AHP-Ahmadpur.** Extraordinary, reduced stem area, pith region and enlarge surface hairs, **g. MUL-Multan.** Thick cortical region reduced vascular bundles and enhanced pith area, **h. VEH-Vehari.** Thicker epidermis, partially crushed cortical region and reduced pith and vascular region, **i. DGK-DG Khan.** Thicker cortical region enhanced vascular bundles and pith region, **j. RJP-Rajanpur.** Greatly reduced stem area, pith thickness and vascular bundle area, **k. JHG-Jhang.** Reduced stem area, vascular region and pith region, **l. MUZ-Muzaffargarh.** Enhanced cortical thickness, vascular region and pith area, **m. SAR-Sargodha.** Thicker epidermis, enlarge vascular bundles and pith region, **n. FSD-Faisalabad.** Thick cortical region, surface hairiness, enlarge vascular bundles and xylem vessels, **o. LYH-Layyah.** Reduced stem area, pith thickness and vascular area.

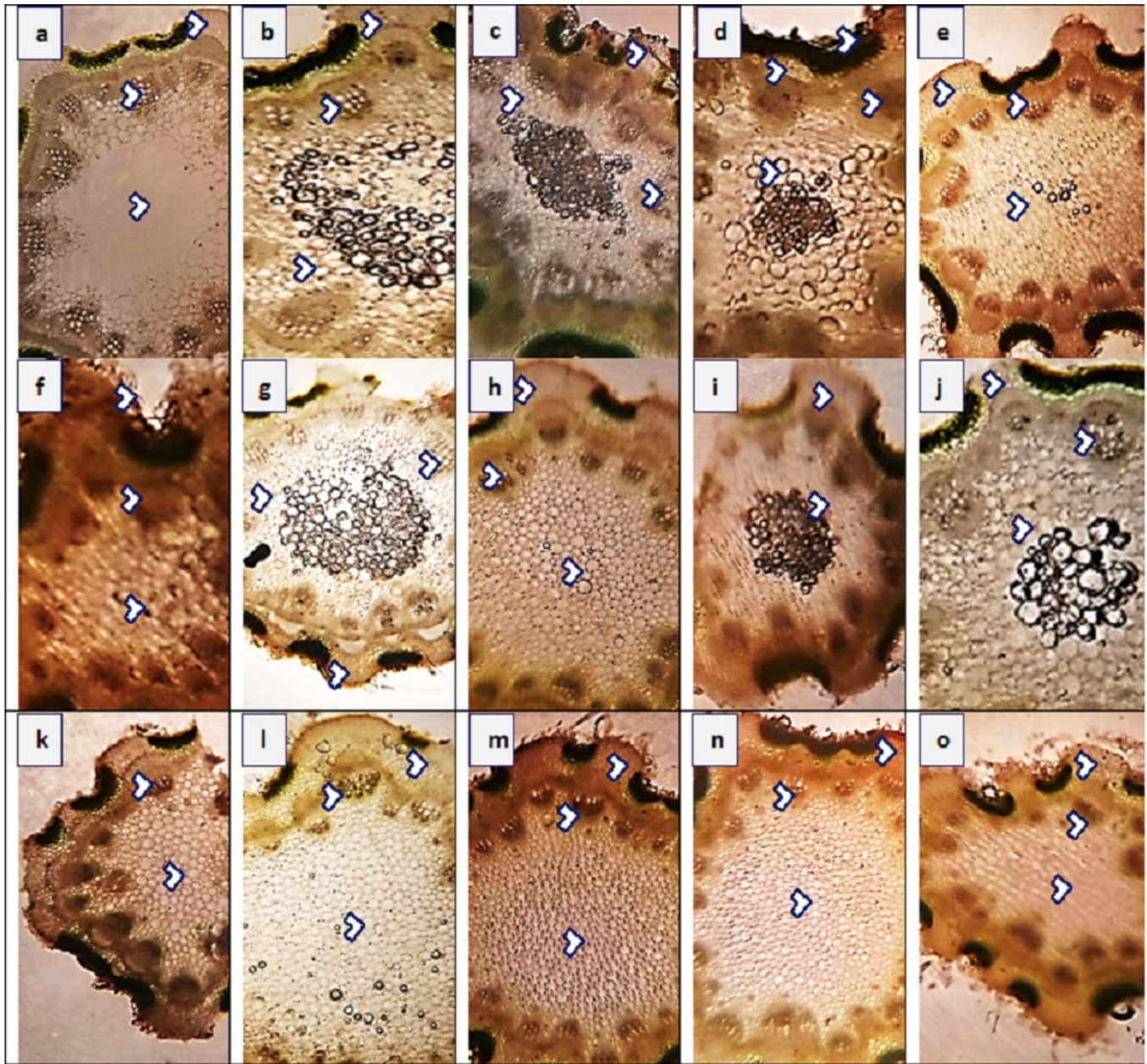


Figure 5

Leaf transvers sections of *Parthenium hysterophorus* L. populations collected from different ecological regions

Description: **a. RYK-Rahim Yar Khan.** Thicker lamina, enlarge proportion of cortical region and reduced vascular bundles, **b. SDK-Sadiqabad.** Thick leaf in terms of midrib and lamina thickness, enhanced cortex thickness and vascular region, **c. KHP-khanpur.** Reduced leaf thickness, enlarge cortical region and vascular bundle area, **d. BWP-Bahawalpur.** Greatly enhanced epidermal thickness, lamina thickness, cortical thickness and vascular area, **e. LAP-Liaquatpur.** Extraordinarily thick leaf, cortical region and vascular bundles, **f. AHP-Ahmadpur.** Extraordinarily thick cortical region, surface hairiness and reduced vascular bundles, **g. MUL-Multan.** Reduced lamina thickness and epidermal thickness enhanced cortical region and vascular area, **h. VEH-Vehari.** Thicker leaf, epidermis, enhanced cortical region and vascular bundle area, **i. DGK-DG Khan.** Sparse surface hairiness, Thick cortical region, enhanced vascular region, **j. RJP-Rajanpur.** Greatly reduced leaf thickness, cortical thickness and enlarged vascular bundle area, **k. JHG-Jhang.** Thick leaf area, vascular bundles and cortical region, **l. MUZ-Muzaffargarh.** Reduced lamina, cortical thickness and large vascular bundles, **m. SAR-Sargodha.** Thick epidermis, enlarge vascular bundles and cortical region, **n. FSD-Faisalabad.** Reduced leaf area, thick cortical region and reduced vascular bundles, **o. LYH-Layyah.** Enhanced surface hairiness, thickness of cortical region and vascular bundles.

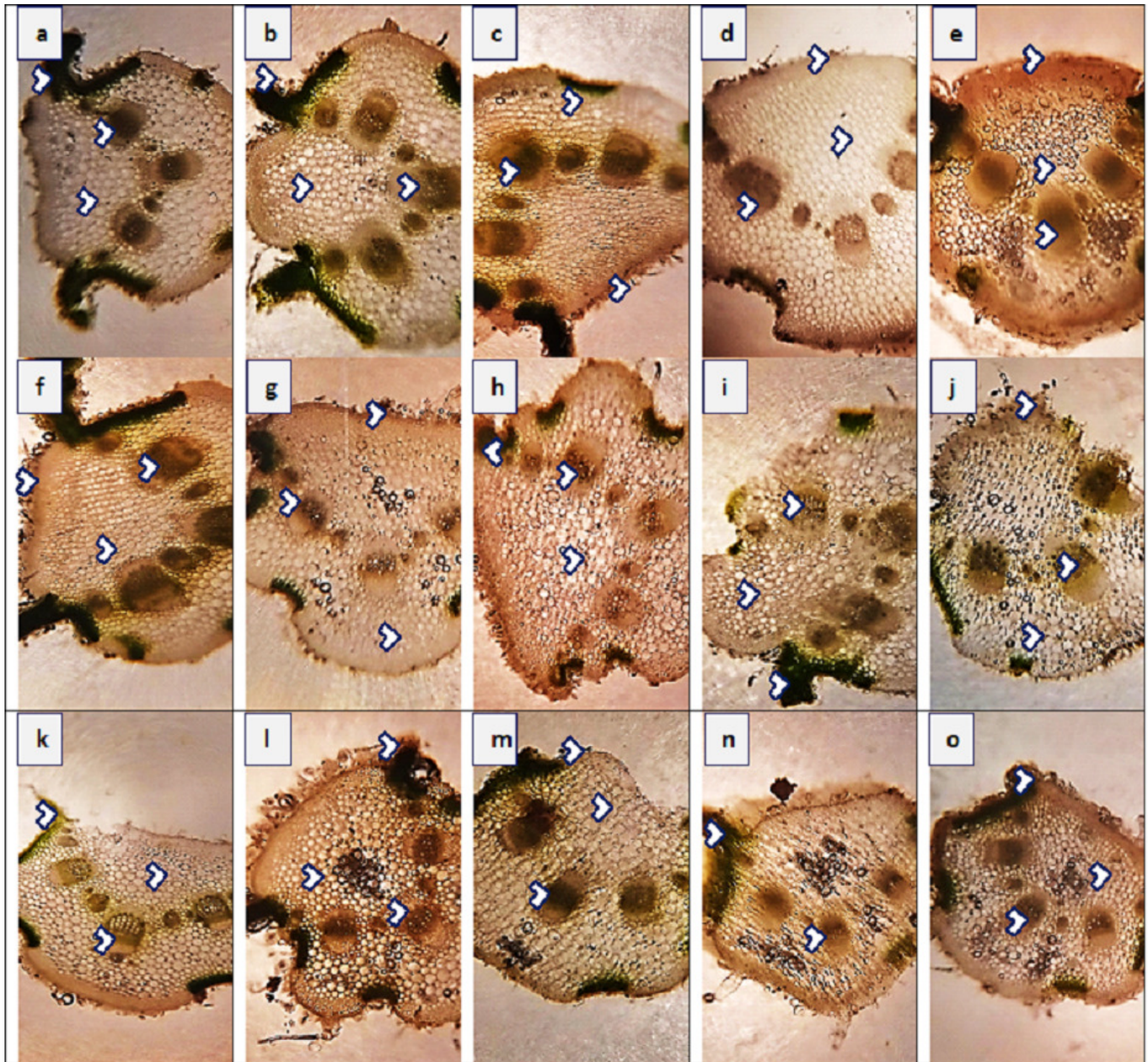


Figure 6

Principal component analysis (PCA) showing influence of soil physicochemical characteristics on A) growth and physiological features, B) root anatomy, C) stem anatomy, D) leaf anatomy of *Parthenium hysterophorus* collected from Punjab province

RYK-Rahim Yar Khan, SDK-Sadiq abad, KHP-Khanpur BWP Bahawalpur, LAP-Liaquatpur, AHP-Ahmadpur, MUL-Multan, VEH-Vehari, MUZ-Muzaffargarh, SAR-Sargodha, FSD-Faisalabad, DGK-Dera Ghazi Khan, RJP-Rajanpur, JHG-JhangLYH-Layyah. **Soil:** ECe-electrical conductivity, pH-soil pH, OM-organic matter, SP-saturation percentage, PO₄-phosphate, NO₃-nitrate, Cl-chloride, Ca-calcium, Na-sodium, K-potassium. **Physiology:** TSP-total soluble proteins, GB-glycine betaine, Chl a-chlorophyll a, Chl b-chlorophyll b, TChl-total chlorophyll, Car-carotenoids, Chl a/b-chlorophyll a/b ratio, TChl/Car-total chlorophyll/carotenoid ratio. **Morphology:** PH-plant height, SFW-shoot fresh weight, SDW-shoot dry weight, LN-leaf number, RL-root length, RFW-root fresh weight, RDW-root dry weight, LA-leaf area, SL-shoot length. **Root anatomy:** RA-root area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area. **Stem anatomy:** SA-stem area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area. **Leaf anatomy:** MrT-midrib thickness, LMT-lamina thickness, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area.

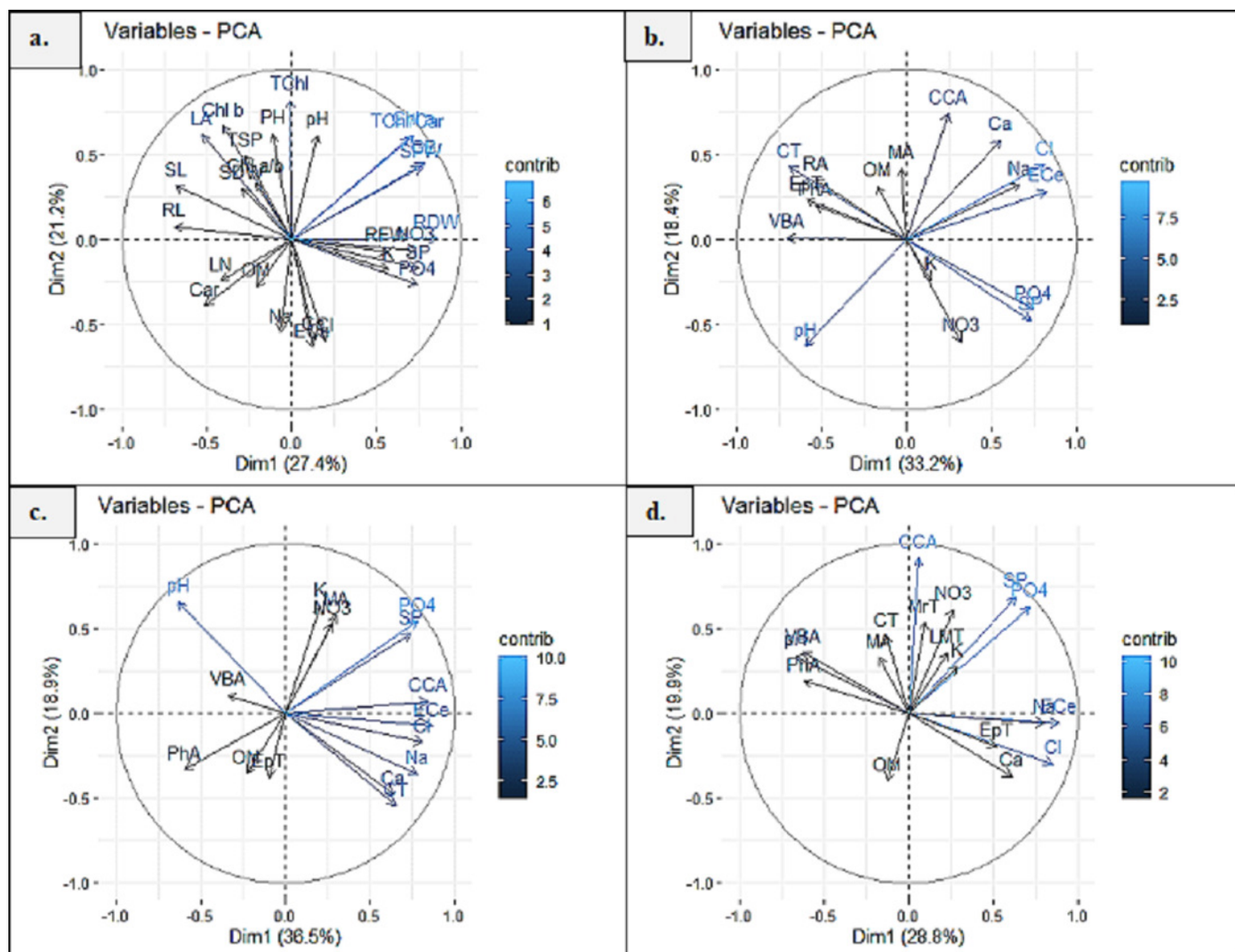


Figure 7

Heatmap showing association of soil physiochemical characteristics on a) growth and physiological characteristics, b) root, c) stem, and d) leaf anatomical features of *Parthenium hysterophorus* collected from the Punjab province

RYK-Rahim Yar Khan, SDK-Sadiq abad, KHP-Khanpur BWP-Bahawalpur, LAP-Liaquatpur, AHP-Ahmadpur, MUL-Multan, VEH-Vehari, MUZ-Muzaffargarh, SAR-Sargodha, FSD-Faisalabad, DGK-Dera Ghazi Khan, RJP-Rajanpur, JHG-Jhang LYH-Layyah. **Soil:** ECe-electrical conductivity, pH-soil pH, OM-organic matter, SP-saturation percentage, PO₄-phosphate, NO₃-nitrate, Cl-chloride, Ca-calcium, Na-sodium, K-potassium. **Physiology:** TSP-total soluble proteins, GB-glycine betaine, Chl a-chlorophyll a, Chl b-chlorophyll b, TChl-total chlorophyll, Car-carotenoids, Chl a/b-chlorophyll a/b ratio, TChl/Car-total chlorophyll/carotenoid ratio. **Morphology:** PH-plant height, SFW-shoot fresh weight, SDW-shoot dry weight, LN-leaf number, RL-root length, RFW-root fresh weight, RDW-root dry weight, LA-leaf area, SL-shoot length. **Root anatomy:** RA-root area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area. **Stem anatomy:** SA-stem area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area. **Leaf anatomy:** MrT-midrib thickness, LMT-lamina thickness, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area

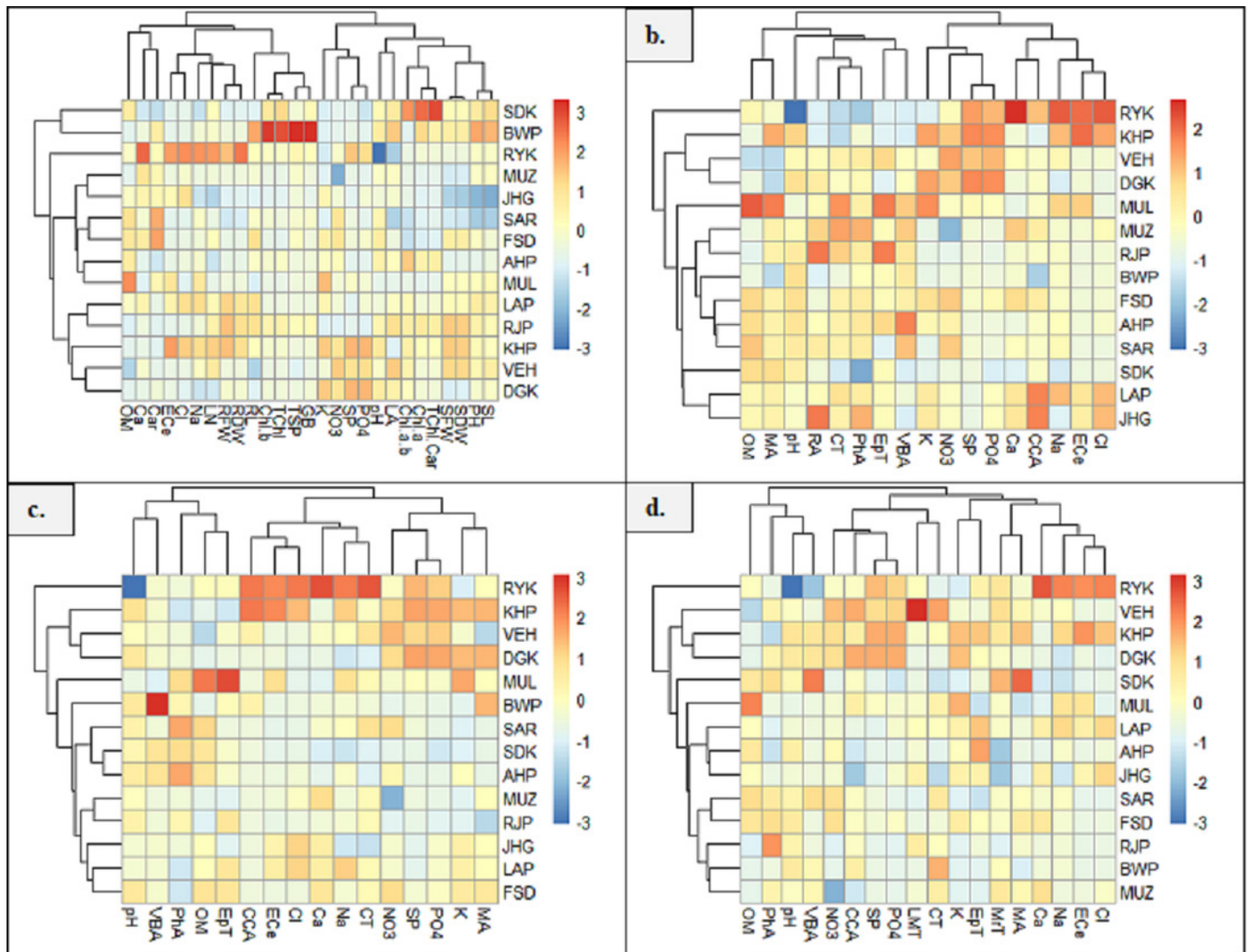


Figure 8

Prominent structural and functional adaptations in different populations of *Parthenium hysterophorus* collected from the Punjab province

RYK-Rahim Yar Khan, SDK-Sadiq abad, KHP-Khanpur, BWP-Bahawalpur, LAP-Liaquatpur, AHP-Ahmadpur, MUL-Multan, VEH-Vehari, MUZ-Muzaffargarh, SAR-Sargodha, FSD-Faisalabad, DGK-Dera Ghazi Khan, RJP-Rajanpur, JHG-JhangLYH-Layyah. **Soil:** ECe-electrical conductivity, pH-soil pH, OM-organic matter, SP-saturation percentage, PO₄-phosphate, NO₃-nitrate, Cl-chloride, Ca-calcium, Na-sodium, K-potassium. **Physiology:** TSP-total soluble proteins, GB-glycine betaine, Chl a-chlorophyll a, Chl b-chlorophyll b, TChl-total chlorophyll, Car-carotenoids, Chl a/b-chlorophyll a/b ratio, TChl/Car-total chlorophyll/carotenoid ratio. **Morphology:** PH-plant height, SFW-shoot fresh weight, SDW-shoot dry weight, LN-leaf number, RL-root length, RFW-root fresh weight, RDW-root dry weight, LA-leaf area, SL-shoot length. **Root anatomy:** RA-root area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vasculer bundle area, MA-metaxyelm area, PhA-phloem area. **Stem anatomy:** SA-stem area, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxyelm area, PhA-phloem area. **Leaf anatomy:** MrT-midrib thickness, LMT-lamina thickness, EpT-epidermal thickness, CT-cortical thickness, CCA-cortical cell area, VBA-vascular bundle area, MA-metaxylem area, PhA-phloem area

