

1 Environment as a limiting factor of the historical global spread of mungbean

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26 **Abstract**

27 While the domestication history has been investigated in many crops, the process of cultivation
28 range expansion and factors governing this process received relatively little attention. Here using
29 mungbean (*Vigna radiata* var. *radiata*) as a test case, we investigated the genomes of more than
30 one thousand accessions to illustrate climatic adaptation's role in dictating the unique routes of
31 cultivation range expansion. Despite the geographical proximity between South and Central Asia,
32 genetic evidence suggests mungbean cultivation first spread from South Asia to Southeast, East,
33 and finally reached Central Asia. Combining evidence from demographic inference, climatic
34 niche modeling, plant morphology, and records from ancient Chinese sources, we showed that
35 the specific route was shaped by the unique combinations of climatic constraints and farmer
36 practices across Asia, which imposed divergent selection favoring higher yield in the south but
37 short-season and more drought-tolerant accessions in the north. Our results suggest that
38 mungbean did not radiate from the domestication center as expected purely under human activity,
39 but instead the spread of mungbean cultivation is highly constrained by climatic adaptation,
40 echoing the idea that human commensals are more difficult to spread through the south-north axis
41 of continents.

42 **Main Text**

43 **Introduction**

44 Domestication is a process where plants or animals were cultivated by humans, leading to
45 associated genetic and morphological changes. These changes may be intentional from human
46 selection or unintentional as a result of adaptation to the environments of cultivation (Fuller,
47 2007). Later, the cultivated plants spread out from their initial geographical range (Meyer and
48 Purugganan, 2013), and elucidating the factors affecting the range expansion of crops is another
49 focus of active research (Gutaker et al., 2020). In the old world, during the process of “prehistory
50 food globalization” (Jones et al., 2011), crops originated from distinct regions were transported
51 and grown in Eurasia. Archeological evidence has shown that such “trans-Eurasian exchange”
52 had happened by 1500 BC (Liu et al., 2019), and the proposed spread routes from archeological
53 studies were supported by modern genetic evidence especially in rice (Gutaker et al., 2020) and
54 barley (Lister et al., 2018). Interestingly, the spread may accompany genetic changes for the
55 adaptation to novel environments. For example, in barley, variations in the gene *Photoperiod-H1*
56 (*Ppd-H1*) resulting in the non-responsiveness to longer daylengths were likely associated with
57 the historical expansion to high-latitude regions (Jones et al., 2008; Jones et al., 2016). While
58 these mid-latitude cereals have been extensively studied, investigations of crops originated from
59 other climate zones are needed. Using the South Asian legume mungbean as a test case, here we
60 investigate how climatic adaptation might affect crop spread route and the evolutionary changes
61 making such spread possible.

62 Mungbean (*Vigna radiata* (L.) Wilczek var. *radiata*), also known as green gram, is an
63 important grain legume in Asia (Nair and Schreinemachers, 2020), providing carbohydrates,
64 protein, folate, and iron for local diets and thereby contributing to food security (Kim et al., 2015).
65 Among pulses, mungbean is capable of tolerating moderate drought or heat stress and has a
66 significant role in rainfed agriculture across arid and semi-arid areas (Pratap et al., 2019), which
67 are likely to have increased vulnerabilities to climate change. Although there have been studies
68 about the genetic diversity of cultivated and wild mungbean (Ha et al., 2021; Kang et al., 2014;
69 Noble et al., 2018; Sangiri et al., 2007), the evolutionary history of cultivated mungbean after
70 domestication still lacks genetic studies. Existing archeological evidence suggests that South Asia

71 is the probable area of mungbean domestication, and at least two independent domestication
72 events have been suggested, including Maharashtra and the eastern Harappan zone (Fuller and
73 Harvey, 2006). The early archeological records suggest that the selection of large seed sizes
74 occurred in the eastern Harappan zone by the 3rd millennium BC and in Maharashtra, dating to
75 the late 2nd to early 1st millennium BC (Fuller and Harvey, 2006). This pulse later spread to
76 mainland Southeast Asia and has been reported in southern Thailand dating to the late first
77 millennium BC (Castillo et al., 2016). Further north, the earliest record of mungbean in China
78 was from the book *Qimin Yaoshu* (齊民要術, 544 AD). While mungbean is also cultivated in
79 Central Asia today, it was not identified in archaeobotanical evidence ranging from several
80 millennium BC to the medieval period (Miller, 1999; Spengler et al., 2018b; Spengler et al., 2017),
81 suggesting later arrival. While the archaeobotanical studies elucidated the route of mungbean
82 cultivation range expansion, researches are still needed to identify the genetic evidence and
83 factors shaping such spread route.

84 A recent genetic study revealed that present-day cultivated mungbeans have the same
85 haplotype in the promoter region reducing the expression of *VrMYB26a* (Lin et al., 2022), a
86 candidate gene controlling the important domestication trait, pod shattering, in several *Vigna*
87 species (Takahashi et al., 2020). This suggests the loss of pod shattering phenotype in cultivated
88 mungbean may have a common origin, and despite the archaeobotanical findings of several
89 independent early cultivations of mungbean in South Asia (Fuller and Harvey, 2006), descendants
90 from one of these cultivation origins might have dominated South Asia before the pan-Asia
91 expansion. Since large regions remain archaeologically unexplored, utilization of genetic data
92 can be a crucial complementation to reconstruct crop evolutionary history. Using seed proteins
93 (Tomooka et al., 1992) and isozymes (Dela Vina and Tomooka, 1994), previous studies proposed
94 two expansion routes out of India, one in the south to Southeast Asia and the other in the north
95 along the silk road to China. While later studies used DNA markers to investigate mungbean
96 population structure (Breria et al., 2020; Gwag et al., 2010; Islam and Blair, 2018; Noble et al.,
97 2018; Sandhu and Singh, 2021; Sangiri et al., 2007), few have examined these hypothesized
98 routes in detail. Therefore, genomic examination of the cultivation range expansion proposed by
99 archaeobotanical studies and the elucidation of its contributing factors are strongly needed.

100 In this study, we compiled an international effort, reporting a global mungbean diversity
101 panel of more than 1,100 accessions derived from (i) the mungbean mini-core collection of the
102 World Vegetable Centre (WorldVeg) genebank, (ii) the Australian Diversity Panel (ADP), and
103 (iii) the Vavilov Institute (VIR), which hosts a one-century-old collection enriched with mid-
104 latitude Asian accessions that are underrepresented in other genebanks, many of which were old
105 landraces collected by Nikolai I. Vavilov and his teams in the early 20th century (Burlyueva et al.,
106 2019). These germplasms harbor a wide range of morphological variations (Figure 1A) and
107 constitute the most comprehensive representation of worldwide mungbean genetic variation. We
108 used this resource to investigate the global history of mungbean after domestication to reveal a
109 spread route highly affected by climatic constraints across Asia, eventually shaping the
110 phenotypic characteristics for local adaptation to distinct environments.

111 **Results**

112 **Population structure and spread of mungbean**

113 Using DArTseq, we successfully obtained new genotype data of 290 mungbean accessions from
114 VIR (Supplementary file 1). Together with previous data (Breria et al., 2020; Noble et al., 2018),
115 our final set included 1,108 samples with 16 wild and 1,092 cultivated mungbean. A total of
116 40,897 single nucleotide polymorphisms (SNPs) were obtained. Of these, 34,469 bi-allelic SNPs,
117 with a missing rate less than 10%, were mapped on 11 chromosomes and retained for subsequent
118 analyses.

119 The genetic structure was investigated based on the 10,359 LD-pruned SNPs. Principal
120 component analysis (PCA, Figure 1C) showed a triangular pattern of genetic variation among
121 cultivated mungbeans, consistent with previous studies (Breria et al., 2020; Noble et al., 2018;
122 Sokolkova et al., 2020) and ADMIXTURE $K = 3$ (Figure 1B). The geographic distribution of
123 these genetic groups is not random, as these three groups are distributed in South Asia (India and
124 Pakistan), Southeast Asia (Cambodia, Indonesia, Philippines, Thailand, Vietnam, and Taiwan),
125 and more northerly parts of Asia (China, Korea, Japan, Russia, and Central Asia). As K
126 increased, the cross-validation (CV) error decreased a little after $K = 4$ (Figure 1-figure
127 supplement 1), where the north group could be further divided (Figure 1B). Therefore, worldwide
128 diversity of cultivated mungbean could be separated into four major genetic groups corresponding

129 to their geography: South Asian (SA), Southeast Asian (SEA), East Asian (EA), and Central
130 Asian (CA) groups. Note that the genetic groups were named after the region where most of their
131 members distribute, and exceptions exist. For example, many EA accessions also distribute in
132 Central Asia, and some SEA accessions were found near the eastern and northeastern coasts of
133 India. Throughout this work, we make clear distinction between genetic group names (e.g., SA)
134 and a geographic region (e.g., South Asia). Therefore, unlike any other previous work in this
135 species, this study incorporates global genetic variation among cultivated mungbean of this
136 important crop.

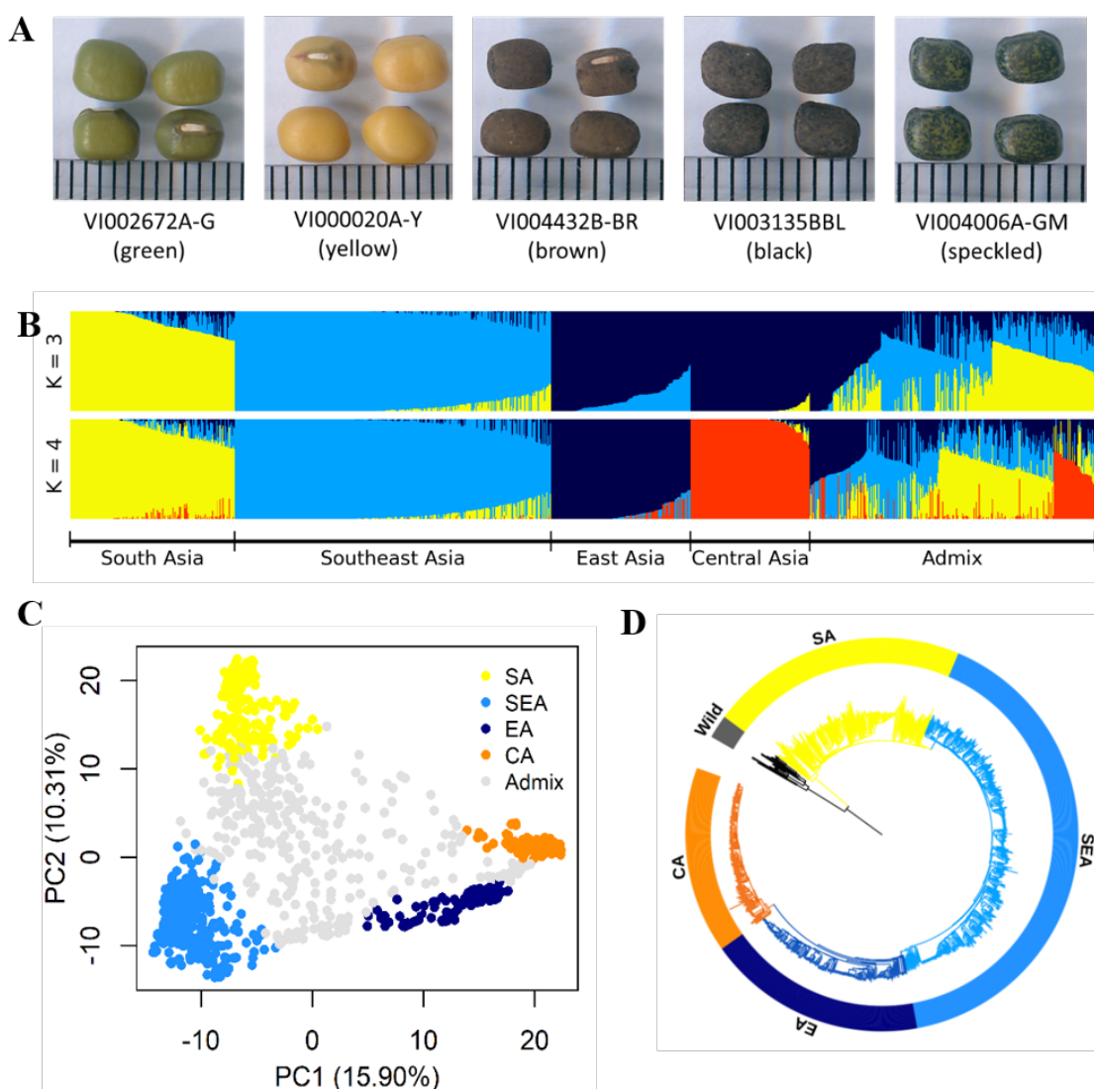


Figure 1. Diversity of worldwide mungbean. (A) Variation in seed colour. (B) ADMIXTURE ancestry coefficients, where accessions were grouped by group assignments ($Q \geq 0.7$). (C) Principal component analysis (PCA) plot of 1,092 cultivated mungbean accessions. Accessions were coloured based on their assignment to four inferred genetic groups ($Q \geq 0.7$), while accessions with $Q < 0.7$ were coloured gray. (D) Neighbor-joining (NJ) phylogenetic tree of 788 accessions with $Q \geq 0.7$ with wild mungbean as outgroup (black colour).

138 Using wild progenitor *V. radiata* var. *sublobata* (Wild hereafter) as the outgroup, the
139 accession-level (Figure 1D) and population-level (Figure 2A) phylogenies both suggest CA to be
140 genetically closest to EA. The SEA group is more distant, and SA is the most diverged. This
141 relationship is supported by the outgroup f_3 tests showing EA and CA share the most genetic drift,
142 followed by SEA and SA (Supplementary file 2). Pairwise F_{ST} and d_{xy} also give the same
143 conclusion (Figure 2B). Similarly, the f_4 tests (Figure 2C) strongly reject the cases where SEA
144 and CA form a clade relative to SA and EA ($f_4(\text{SA,EA;SEA,CA}) = 0.016$, $Z = 9.519$) or SEA and
145 EA form a clade relative to SA and CA ($f_4(\text{SA,CA;SEA,EA}) = 0.021$, $Z = 13.956$), again
146 suggesting EA and CA to be closest. With regards to the relationship among Wild, SA, SEA, and
147 EA, f_4 tests suggest SEA and EA form a clade relative to Wild and SA (non-significant results in
148 $f_4(\text{Wild,SA;EA,SEA})$ but opposite in other combinations). Notably, both TreeMix (Figure 2A)
149 and the f_4 test (Figure 2C, $f_4(\text{SA,SEA;CA,EA}) = 0.005$, $Z = 6.843$) suggest gene flow between
150 SEA and EA. Consistent with archeological evidence of South Asian domestication, the
151 nucleotide diversity (π) decreased from SA (1.0×10^{-3}) to SEA (7.0×10^{-4}) and EA (5.0×10^{-4}),
152 while the CA group has lowest diversity (3.0×10^{-4}) (Figure 2B). LD also decays the fastest in
153 Wild and then the SA group (Figure 2D), followed by other genetic groups. In summary, all
154 analyses are consistent with our proposed order of cultivated mungbean divergence.

155 Our proposed demographic history could be confounded by factors such as complex
156 hybridization among groups. For example, SEA and CA might have independently originated
157 from SA and later generated a hybrid population in EA (Figure 2-figure supplement 1A). Other
158 possibilities are that either SEA or CA are the hybrid of other populations (Figure 2-figure
159 supplement 1B and C). We examined these possibilities using f_3 statistics for all possible trios
160 among the four groups. None of the tests gave a significantly negative f_3 value (Supplementary
161 file 3), suggesting the lack of a strong alternative model to our proposed relationship among these
162 four groups.

163 Based on the solid relationship among these genetic groups, we used fastsimcoal2 to
164 model their divergence time, allowing population size change and gene flow at all time points
165 (Figure 2-figure supplement 2A-D). According to this model, after initial domestication, the out-
166 of-India event (when other groups diverged from SA) happened about 8.3 thousand generations
167 ago (kga) with 75% parametric bootstrap range between 4.7 and 11.3 kga. Not until more than

168 five thousand generations later (2.7 kga, 75% range 1.1-4.6 kga) did SEA diverge from the
169 common ancestor of present-day EA and CA. CA diverged from EA only very recently (0.2 kga,
170 75% range 0.1-0.8 kga). Note that the divergence time was estimated in the number of generations,
171 and the much longer growing seasons in the southern parts of Asia may allow more than one
172 cropping season per year (Mishra et al., 2022; Vir et al., 2016).

173 Our results suggest the non-South-Asian accessions have a common origin out of India
174 (otherwise these groups would branch off independently from the SA group). Given this, the
175 phylogenetic relationship (Figure 2A) is consistent with the following hypotheses. (1) The east
176 hypothesis: Mungbean expanded eastwards and gave rise to the SEA group. This group might
177 initially occupy northeast South Asia and later expanded to Southeast Asia either through the land
178 or maritime route (Castillo et al., 2016; Fuller et al., 2011). The group later expanded northwards
179 as EA. EA expanded westwards into Central Asia and gave rise to the CA group. (2) The north
180 hypothesis: The group leaving South Asia first entered Central Asia as the EA group. EA
181 expanded eastwards into East Asia through the Inner Asian Mountain Corridor (Stevens et al.,
182 2016). The eastern population of EA expanded southwards as the SEA group, and later the
183 western population of EA diverged as the CA group. (3) The northeast hypothesis: The group
184 leaving South Asia (through either of the above-mentioned routes) was first successfully
185 cultivated in northern East Asia without previously being established in Southeast Asia or Central
186 Asia. The EA group then diverged southwards as SEA and later expanded westwards, giving rise
187 to CA. Consistent with this model, the genetic variation of the EA group gradually declines from
188 east to west, accompanied by the gentlest decline of precipitation per unit geographic distance
189 across Asia (Figure 2F).

190 While all three hypotheses are consistent with the phylogeny (Figure 2A), the SEA group
191 originated earlier than EA in the east hypothesis but later in the two other hypotheses. The former
192 case predicts higher nucleotide diversity and faster linkage disequilibrium decay in SEA than EA,
193 which is supported by our results (Figure 2B and D). While populations that were established in
194 a region for an extended time could accumulate genetic differentiation, generating patterns of
195 isolation by distance, rapid-spreading populations in newly colonized regions could not (Lee et
196 al., 2017; The 1001 Genomes Consortium, 2016). Using this idea, Mantel's test revealed a
197 significantly positive correlation between genetic and geographic distances for the SA genetic

198 group ($r = 0.466$, $P = 0.010$), followed by SEA ($r = 0.252$, although not as significant, $P = 0.069$).
199 No such association was found for EA ($r = 0.030$, $P = 0.142$) or CA ($r = 0.087$, $P = 0.172$). In
200 addition, the southern groups (SA and SEA) together ($r = 0.737$, $P = 0.001$) have a much stronger
201 pattern of isolation by distance than the northern groups (EA and CA, $r = 0.311$, $P = 0.001$)
202 (Figure 2E). Using $Q \geq 0.5$ instead of $Q \geq 0.7$ to assign individuals into genetic groups generated
203 results that are largely consistent (Supplementary file 4). These results are again consistent with
204 the “east hypothesis” that local accessions from the SA and SEA groups were established much
205 earlier than those from EA and CA. Finally, the genetic variation of the EA group is highest in
206 the eastern end and declines westwards (Figure 2F). This does not support the north hypothesis
207 where EA first existed in Central Asia and expanded eastwards.

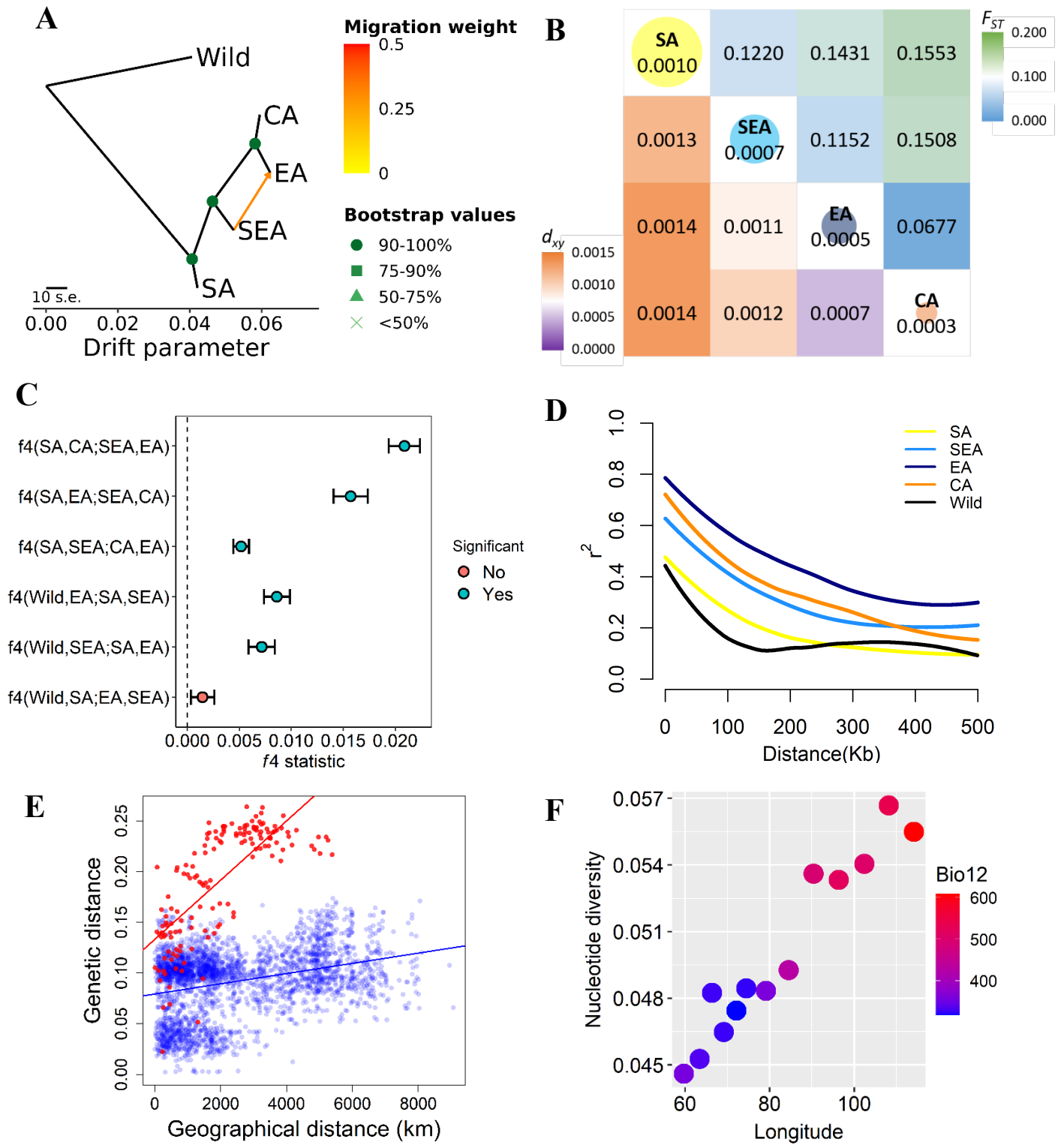


Figure 2. Fine-scale genetic relationship and admixture among four inferred genetic groups. (A) TreeMix topologies with one suggested migration event. Colours on nodes represent support values after 500 bootstraps. (B) Diversity patterns within and between inferred genetic groups as estimated using nucleotide diversity (π in diagonal, where the size of the circle represents the level of π) and population differentiation (F_{ST} in upper diagonal and d_{xy} in lower diagonal). (C) f_4 statistics. Points represent the mean f_4 statistic and lines are the standard error. Only f_4 statistics with Z-score $> |3|$ are considered statistically significant. The dashed line denotes $f_4 = 0$. (D) Linkage disequilibrium (LD) decay. (E) Isolation by distance plot of genetic distance versus geographic distance, with the southern group in red circles and the northern group in blue circles. (F) Relationship between Bio12 (annual precipitation) and nucleotide diversity (π) of the EA genetic group across the east-west axis of Asia. Dot colors represent the annual precipitation of each population.

210 **Environmental differentiation of the inferred genetic groups**

211 We further examined the possible causes governing the expansion of mungbean cultivation
212 ranges. For a crop to be successfully cultivated in a new environment, dispersal and adaptation
213 are both needed. Being a crop that has lost the ability of pod shattering, the spread of mungbean
214 was governed by commerce or seed exchange. While barriers such as the Himalayas or Hindu
215 Kush may limit human activity, South and Central Asia was already connected by a complex
216 exchange network linking the north of Hindu Kush, Iran, and the Indus Valley as early as about
217 4 thousand years ago (kya) (Dupuy, 2016; Kohl, 2007; Kohl and Lyonnet, 2008; Lamberg-
218 Karlovsky, 2002; Lombard, 2020; Lyonnet, 2005), and some sites contain diverse crops
219 originated across Asia (Spengler et al., 2021). Similarly, other ancient land or maritime exchange
220 routes existed among South, Southeast, East, and Central Asia (Stevens et al., 2016). This
221 suggests that mungbean could have been transported from South to Central Asia, but our genetic
222 evidence suggests that the present-day CA group did not descend directly from the SA group.
223 Therefore, we investigated whether climatic adaptation, that is, the inability of mungbean to
224 establish in a geographic region after human-mediated long-range expansion, could be a
225 contributing factor.

226 Multivariate analysis of variance (MANOVA) of eight bioclimatic variables (after
227 removing highly-correlated ones; Supplementary file 5 and 6) indicated strong differentiation in
228 the environmental niche space of the four genetic groups (Supplementary file 7 and 8). PCA of
229 climatic factors clearly reflects geographic structure, where the axis explaining most variation
230 (PC1, 42%) separates north and south groups and is associated with both temperature- and
231 precipitation-related factors (Figure 3A and Supplementary file 9). Consistent with their
232 geographic distribution, overlaps between EA and CA and between SA and SEA were observed.
233 While these analyses were performed using bioclimatic variables from year-round data, we
234 recognized that summer is the cropping season in the north. Parallel analyses using the
235 temperature and precipitation of May, July, and September yielded similar results
236 (Supplementary file 10; Figure 3-figure supplement 1).

237 Based on the Köppen climate classification (Köppen, 2011), we categorized the Asian
238 mungbean cultivation range into six major climate zones (Figure 3-figure supplement 2): dry hot
239 (BSh and BWh), dry cold (BSk and BWk), temperate dry summer (Csa), tropical savanna (Aw),

240 Continental (Dwb and Dfb), and temperate wet summer (Cfa and Cwa). The former three are
241 relatively drier than the latter three zones. While SEA and CA are relatively homogeneous, SA
242 and EA have about half of the samples in the dry and non-dry zones (Figure 3-figure supplement
243 2). We therefore separated SA into SAe and SAw and EA into EAe and EAw, corresponding to
244 the wetter eastern and drier western regions within the SA and EA ranges. Environmental niche
245 modeling revealed distinct suitable regions of these six groups except for CA and EAw, whose
246 geographical ranges largely overlap (Figure 3B). Consistent with PCA, pairwise Schoener's D
247 values are smallest between the northern and southern groups while largest (suggesting overlaps
248 of niche space) between the eastern and western subsets within north and south (Figure 3B),
249 consistent with PCA that the major axis of climatic difference is between the northern and
250 southern parts of Asia. Analyses using temperature and precipitation from May, July, and
251 September yielded similar results (Figure 3-figure supplement 3). Given a single out-of-India
252 event (Figure 2A), the results suggest it might be easier to first cultivate mungbean in Southeast
253 rather than Central Asia, supporting the east hypothesis.

254 While both temperature and precipitation variables differ strongly between north and
255 south, one should note that these year-round temperature variables do not correctly reflect
256 conditions in the growing seasons. In the north, mungbean is mostly grown in summer where the
257 temperature is close to the south (Figure 3-figure supplement 4A-C). On the other hand,
258 precipitation differs drastically between the north and south, especially for the CA group, where
259 the summer growing season is the driest of the year (Figure 3-figure supplement 4D). By
260 estimating the regression slope of annual precipitation on geographical distance, we obtained a
261 gradient of precipitation change per unit geographic distance between pairs of genetic groups
262 (Figure 3C). Despite the SA-SEA transect having the steepest gradient (slope = 0.21), the spread
263 from SA to SEA has been accompanied by an increase of precipitation and did not impose drought
264 stress. However, the second highest slope (0.18) is associated with a strong precipitation decrease
265 if the SA group were to disperse to Central Asia. Results from the precipitation of May, July, and
266 September yielded similar conclusion (Figure 3-figure supplement 5). This likely explains why
267 no direct historic spread is observed from South to Central Asia.

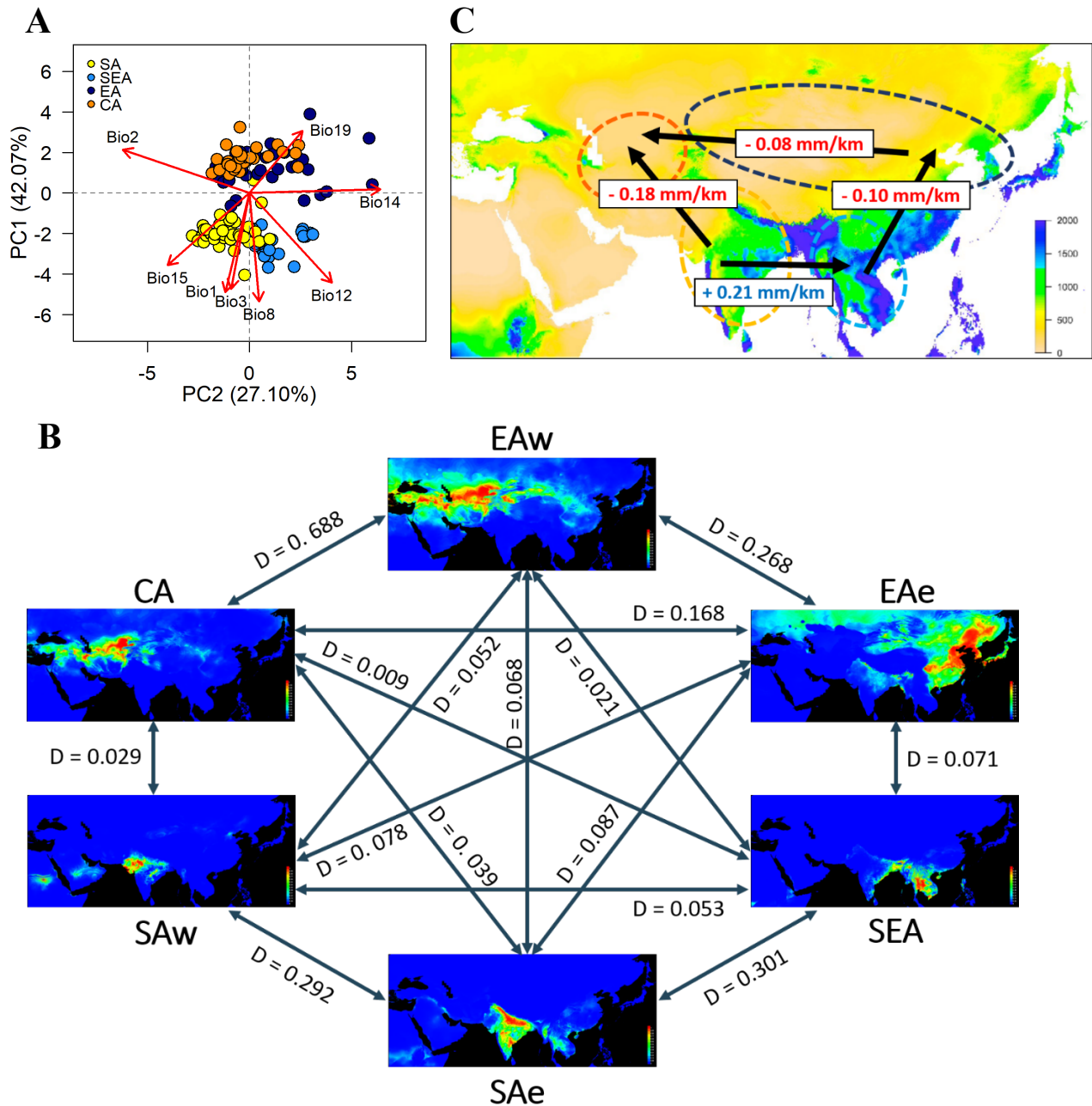


Figure 3. Environmental variation among genetic groups of mungbean. (A) Principal component analysis (PCA) of the eight bioclimatic variables. Samples are coloured according to four inferred genetic groups as indicated in the legend. (B) Predicted distribution at current climate conditions. Red colour indicates high suitability, and blue indicates low suitability. Values between pairs represent niche overlap measured using Schoener's D, and higher values represent higher overlaps. Abbreviations: SAw: South Asia (west), SAe: South Asia (east); SEA: Southeast Asia; EAe: East Asia (east); EAw: East Asia (west) and CA: Central Asia. (C) Environmental gradient across potential directions of expansion. The value on each arrow indicates a change in annual precipitation per kilometer. The background map is colored according to annual precipitation (Bio12, in mm).

269 **Trait variation among genetic groups**

270 If environmental differences constrained the spread route of mungbean, the current cultivated
271 mungbean accessions occupying distinct environments should have locally adaptive traits for
272 these environments. Indeed, PCA of four trait categories show substantial differences among
273 genetic groups (phenology, reproductive output, and size in field trials, as well as plant weight in
274 lab hydroponic systems, Figure 4A). In the field, CA appears to have the shortest time to
275 flowering, the lowest yield in terms of seed size and pod number, and the smallest leaf size (Figure
276 4B and Supplementary file 11). On the other hand, SEA accessions maximize seed size, while
277 SA accessions specialize in developing the largest number of pods (Figure 4B). These results
278 suggest that CA has a shorter crop duration, smaller plant size, and less yield, consistent with
279 drought escape phenotypes. This is consistent with the northern short growing season constrained
280 by temperature and daylength (below), as well as the low precipitation during the short season.

281 In terms of seedling response to drought stress, the Q_{ST} values of most traits (root, shoot,
282 and whole plant dry weights under control and drought treatments) are higher than the tails of
283 SNP F_{ST} , suggesting trait evolution driven by divergent selection (Figure 4C; Figure 4-figure
284 supplement 6). Significant treatment, genetic group, and treatment by group interaction effects
285 were observed except on a few occasions (Table 1). Consistent with field observation, SEA has
286 the largest seedling dry weight (Figure 4D). While simulated drought significantly reduced shoot
287 dry weight for all groups, the effect on SEA is especially pronounced (treatment-by-group
288 interaction effect, $F_{2,575} = 23.55$ $P < 0.001$, Table 1 and Figure 4D), consistent with its native
289 habitats with abundant water supply (Figure 3-figure supplement 4D and Supplementary file 12).
290 All groups react to drought in the same way by increasing root:shoot ratio (Figure 4D), suggesting
291 such plastic change may be a strategy to reduce transpiration. Despite the lack of treatment-by-
292 group interaction ($F_{2,575} = 1.39$, $P > 0.05$), CA consistently exhibits a significantly higher
293 root:shoot ratio, a phenotype that is potentially adaptive to its native environment of lower water
294 supply (Figure 3-figure supplement 4D and Supplementary file 12).

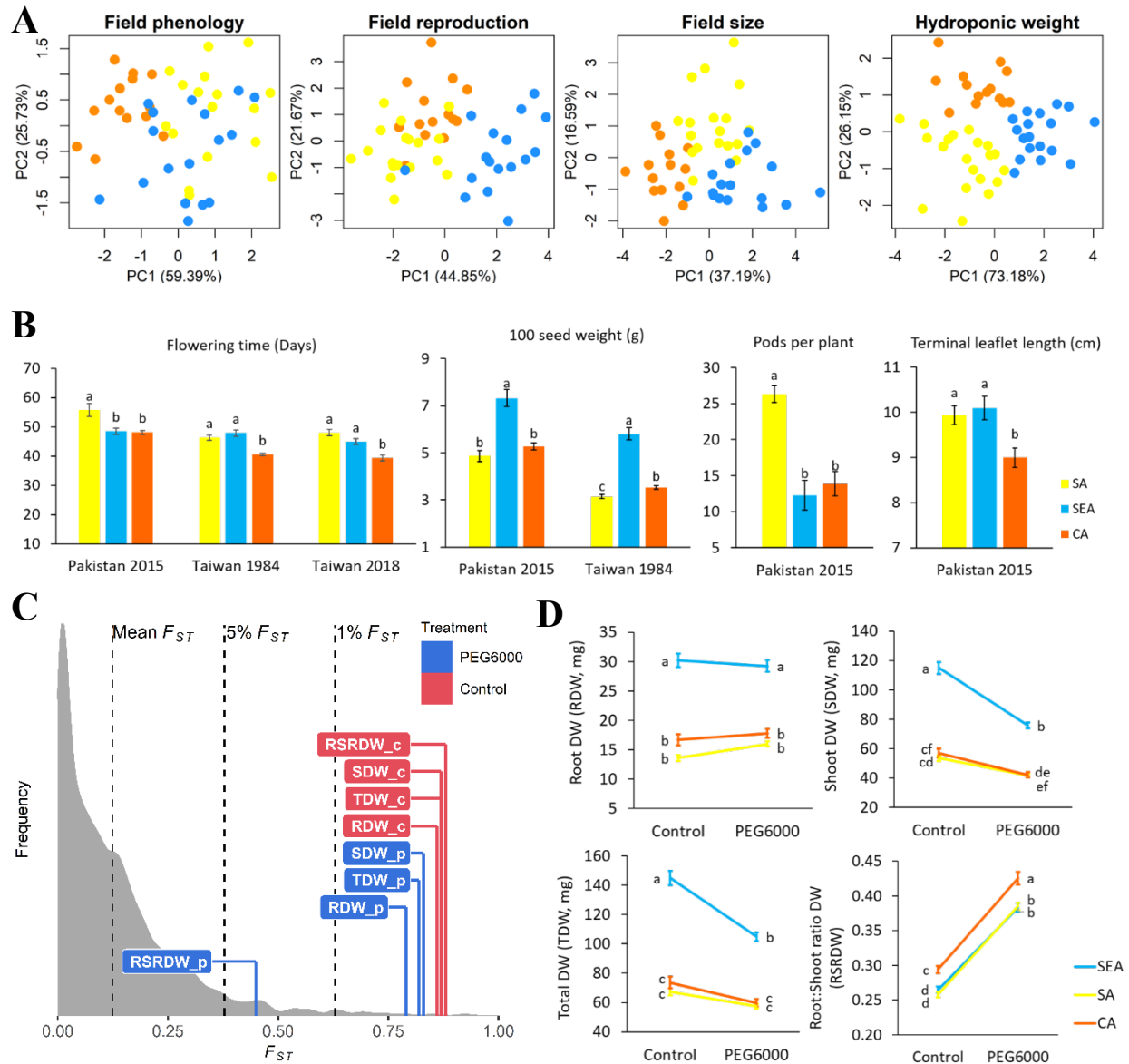


Figure 4. Quantitative trait differentiation among genetic groups. (A) Principal component analysis (PCA) of four trait categories. (B) Trait variability from common gardens in field experiments. (C) Comparison of Q_{ST} - F_{ST} for four drought-related traits under two environments. F_{ST} values (mean, 5% and 1%) were indicated by black dashed lines. The Q_{ST} for each trait was colored according to treatment and was calculated as Equation 2 in Materials and Methods. Abbreviations: RDW: root dry weight; SDW: shoot dry weight; TDW: total dry weight; RSRDW: root:shoot ratio dry weight; c: control; p: PEG6000. (D) Effect of PEG6000 (-0.6 MPa) on root dry weight (RDW), shoot dry weight (SDW), total dry weight (TDW), and root:shoot ratio dry weight (RSRDW) among genetic groups. Data were expressed as the mean \pm standard error. Lowercase letters denote significant differences under Tukey's HSD in (B) and (D).

Table 1. Analysis of variance (ANOVA) *F* values for the dry weight (mg) of mungbean seedlings across three different genetic groups

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Source of variation	df	RDW	SDW	TDW	RSRDW
Treatment	1	2.65 ^{n.s.}	133.26 ^{***}	72.26 ^{***}	978.76 ^{***}
Genetic group	2	60.63 ^{***}	79.62 ^{***}	76.54 ^{***}	13.27 ^{***}
Treatment x Genetic group	2	3.29 [*]	23.55 ^{***}	17.79 ^{***}	1.39 ^{n.s.}

(df: degrees of freedom; RDW: root dry weight; SDW: shoot dry weight; TDW: total dry weight; RSRDW: root:shoot ratio dry weight. Significance level * $P < 0.05$, *** $P < 0.001$, n.s. non-significant)

297

298 **Support from ancient Chinese sources**

299 Mungbean has been occasionally mentioned in ancient Chinese sources. Here we report the
300 records associated with our proposed mungbean spread route and the underlying mechanisms.
301 The “Classic of Poetry” (Shijing 詩經) contains poems dating between the 11th to 7th centuries
302 BCE near the lower and middle reaches of the Yellow River. While crops (especially soy bean,
303 菽), vegetables, and many other plants have been mentioned, mungbean was not recorded. This
304 is consistent with our results that mungbean had not reached the northern parts of East Asia at
305 that time (the EA group diverged from the SEA group at around 2.7 kya). The first written record
306 of mungbean in China is in an agricultural encyclopedia Qimin Yaoshu (齊民要術, 544 AD,
307 Chinese text and translation in Supplementary note), whose spatiotemporal background (~1.5 kya
308 near the lower reaches of Yellow River) is again consistent with our estimated origin of the EA
309 group.

310 Our results suggest that the expansion of the mungbean cultivation range may be
311 associated with the novel phenotypic characteristics potentially adaptive to the new environments.
312 This proposal would be rejected if the novel phenotypic characteristics appeared very recently.
313 In support of our proposal, Xiangshan Yelu (湘山野錄, an essay collection during 1068-1077
314 AD) recorded that mungbean from the southern parts of Asia had higher yield and larger grains
315 than those in northern China (Chinese text and translation in Supplementary note). Similarly,
316 Tiangong Kaiwu (天工開物, 1637 AD) mentioned that mungbean must be sown during July and
317 August (Chinese text and translation in Supplementary note). The record suggests that the
318 daylength requirement restricts the sowing period of mungbean in the north. Together with the

319 dry summer (Figure 3-figure supplement 4D) and soon-arriving autumn frost, there might be a
320 strong selection favoring accessions with the rapid life cycle. These records suggest the
321 phenotypic characteristics of northern accessions did not originate very recently, and the unique
322 distribution of climatic zones in Asia resulted in not only the specific patterns of expansion but
323 also the evolution of novel phenotypic characteristics in mungbean.

324 **Discussion**

325 Using mungbean as a test case, we combined population genomics, environmental niche
326 modeling, empirical field and laboratory investigation, and ancient Chinese text analyses to
327 demonstrate the importance of climatic adaptation in dictating the unique patterns of cultivation
328 range expansion after domestication. In this study, we focus on how or when mungbean could be
329 established as part of local agriculture throughout Asia. We showed that after leaving South Asia,
330 mungbean was likely first cultivated in Southeast Asia, East Asia, and finally Central Asia. We
331 acknowledge that our data do not allow us to specify the number of previous out-of-India events
332 that did not leave traces in modern genetic data or their exact routes (for example, whether
333 mungbean expanded from South to Southeast Asia through the land or maritime routes). While
334 there might be multiple attempts to bring mungbean out of India as a commodity for consumption,
335 our results suggest all present-day non-South-Asian accessions have a common out-of-India
336 origin.

337 **The climate-driven spread route despite historical human activities**

338 Combining archeological records, population genetics, and niche modeling (Figure 2 and 3), our
339 results suggest that after the early cultivation of mungbean in northwestern or southern South
340 Asia (Fuller, 2007; Kingwell-Banham et al., 2015), the large environmental difference may
341 restrict its northwards spread to Central Asia. Mungbean may first spread to eastern South Asia,
342 and the subsequent expansion to Southeast Asia might be facilitated by the environmental
343 similarity between these two regions. This is supported by archaeobotanical remains from the
344 Thai-Malay Peninsula date to ca. 400-100 BCE (Castillo et al., 2016). It took more than five
345 thousand generations until mungbean further spread to northeast Asia, again likely due to the
346 environmental difference. The later appearance of mungbean in northern China is also supported
347 by historical records. After that, the EA group spread across the northern part of Asia within a

348 few thousand generations. Our proposed route suggests that mungbean reached Central Asia at
349 the latest, consistent with its absence from archeological sites in Central Asia, including
350 Turkmenistan and Uzbekistan in the Chalcolithic and Bronze ages (5th to 2nd millennium BC)
351 (Miller, 1999), Southeastern Kazakhstan in the Iron age dating 1st millennium BC (Spengler et
352 al., 2017), and eastern Uzbekistan during the medieval period (800-1100 AD) (Spengler et al.,
353 2018b). In addition, mungbean was only mentioned later by the 18th and early 19th centuries as a
354 pulse grown in the Khiva region of Uzbekistan (Annanepesov and Bababekov, 2003).

355 In this study, we suggest that the ability to disperse may not be an essential factor
356 restricting mungbean spread from South to Central Asia. Cultivated mungbean has lost the natural
357 ability of pod shattering to disperse seeds, and they mostly traveled through landscapes by
358 human-mediated seed exchange or commerce. Evidence of long-distance human-mediated
359 dispersal of mungbean was available. For example, mungbean seeds have been found near the
360 Red Sea coast of Egypt during the Roman (AD 1–250) period (Van der Veen and Morales, 2015).
361 As early as about four kya, the Bactria–Margiana Archaeological Complex (BMAC) civilization
362 north of the Hindu Kush had extensive contact with the Indus Valley Civilization (Dupuy, 2016;
363 Kohl, 2007; Kohl and Lyonnet, 2008; Lamberg-Karlovsky, 2002; Lombard, 2020; Lyonnet,
364 2005). By 1,500 BC, the “Trans-Eurasian Exchanges” of major cereal crops has happened (Liu
365 et al., 2019). The frequent crop exchange is evidenced by archaeobotanical findings in the Barikot
366 site (ca. 1200 BC-50 AD) in northern Pakistan (Spengler et al., 2021), where diverse crops were
367 cultivated, including those from West Asia (wheat, barley, pea, and lentil), South Asia
368 (urdbean/mungbean), and likely East Asia (rice). Despite this, in Bronze-age archeological sites
369 north of Hindu Kush, legumes (such as peas and lentils) were observed to a lesser extent than
370 cereals, and South Asian crops were not commonly found (Jeong et al., 2019; Spengler, 2015;
371 Spengler et al., 2014a; Spengler et al., 2018a; Spengler et al., 2014b). Interestingly, archeologists
372 suggested legume’s higher water requirement than cereals may be associated with this pattern,
373 and pea and lentil’s role as winter crops in Southwest Asia may be associated with their earlier
374 appearance in northern Central Asia than other legumes (Spengler et al., 2014a; Spengler et al.,
375 2018a; Spengler et al., 2014b). Therefore, despite the possibility of human-mediated seed
376 dispersal between South and Central Asia, our results and archeological evidence concurred that
377 mungbean arrived in Central Asia at the latest, likely restricted by environmental adaptation.

378 **Local adaptation of mungbean genetic groups**

379 Despite the profound impact of human-mediated dispersal on the spread of these and many other
380 crops (Herniter et al., 2020; Kistler et al., 2018), in mungbean we suggest adaptation to distinct
381 climatic regimes to be an important factor in the establishment after dispersal. Mungbean is
382 commonly grown under rainfed cultivation and depends on the residual moisture in the fields
383 after the primary crop, thus responding to water stress (Douglas et al., 2020). In the south, a
384 temperature range of 20-30°C and annual precipitation of 600-1,000 mm is optimal for mungbean
385 (Ha and Lee, 2019). In Central Asia, however, the annual precipitation could be as low as 286
386 mm, greatly below the lower limit required for the southern mungbean. This situation could be
387 further exacerbated by the fact that mungbean might not be a highly valued crop under extensive
388 care during cultivation. Indeed, the earliest record of mungbean in China (Qimin Yaoshu 齊民要
389 術, 544 AD) emphasizes its use as green manure. In Central Asia, mungbean is a minor crop
390 (Rani et al., 2018) grown with little input, only in the short duration between successive planting
391 of main crops (which is also the dry season in Central Asia, Supplement file 12 and Figure 3-
392 figure supplement 4) and using residual soil moisture with little irrigation. We suggest that the
393 lack of extensive input subjects mungbean to more substantial local climatic challenges than
394 highly valued high-input crops that receive intensive management, including irrigation. Therefore,
395 the combination of climatic constraints and cultural usage, instead of physical barriers, may have
396 shaped the historical spread route of the mungbean despite extensive human activities across the
397 continent.

398 In addition to the constraint of soil moisture, other factors may have contributed to the
399 selection of short-season accessions in the north. In the short summer seasons of much of Central
400 Asia, short crop cycling is a requirement. In Uzbekistan, mungbean is often sown in early July
401 after the winter wheat season and harvested before mid-October to avoid delays in the next round
402 of winter wheat and escape frost damage. Therefore, fast-maturing accessions are essential for
403 this production system (Rani et al., 2018). Similar rotation systems using mungbean to restore
404 soil fertility during the short summer season after the harvest of the main crop were also
405 mentioned in ancient Chinese sources (Chen, 1980). Mungbean is a short-day species from the
406 south, and daylength likely limits the window when mungbean could be grown in the north:

407 Chinese texts during the 17th century (Tiangong Kaiwu 天工開物, 1637 AD) specifically
408 mentioned the suitable duration to sow mungbean to control the flowering behavior for maximum
409 yield (Supplementary note). Therefore, unlike in the south where yield appears to be an important
410 selection target, the unique combination of daylength, agricultural practices, soil water
411 availability, and frost damage in the north requires the selection for short-season accessions,
412 likely limiting the direct adoption of southern accessions in the north. Consistent with this,
413 Central Asian accessions have a faster life cycle potentially adaptive to both short growing season
414 and reduced soil water availability, with reduced plant size and lower yield as tradeoffs. These
415 accessions also have increased root:shoot ratio for drought adaptation, similar to findings in rice
416 (Xu et al., 2015), alfalfa (Zhang et al., 2018), and chickpea (Kumar et al., 2012).

417 About accession sampling and climatic niche modeling, we recognize that not all samples
418 have available spatial data, and we do not have samples from some parts of Asia. For example,
419 while most samples of the SEA group were collected from Taiwan, Thailand, and Philippines,
420 we do not have many samples from the supposed contact zone between SA and SEA (Bangladesh
421 and Myanmar) or between SEA and EA (southern China). If more samples were available from
422 these contact zones, the modeled niche space between SA and SEA and between SEA and EA
423 would be even more similar than the current estimate, strengthening our hypothesis that niche
424 similarity might facilitate the cultivation expansion. On the other hand, clear niche differentiation
425 between SA and CA was evident despite the dense sampling near their contact zone. Based on
426 the Köppen climate classification, South Asia could be roughly separated into two major zones,
427 with the eastern zone slightly more similar to Southeast Asia (Figure 3-figure supplement 2). This
428 partially explained the existence of some SEA accessions in the northeastern coast of India. While
429 the SEA genetic group was named after the geographic region where most of its members were
430 found in the present time, we recognize the possibility that it first occupied northeastern South
431 Asia when it diverged from SA. In that case, the SA-SEA divergence time (4.7 to 11.3 thousand
432 generations ago) might indicate the divergence between the two climate zones within South Asia
433 rather than the expansion of mungbean into Southeast Asia, which may occur much later.

434 **Conclusion**

435 Our study demonstrates that mungbean's cultivation range expansion is associated with climatic
436 conditions, which shaped the genetic diversity and contributed to adaptive differentiation among
437 genetic groups. The climatic differences likely also resulted in farmers' differential emphasis on
438 using it mainly as a grain or green manure crop, further intensifying the phenotypic diversification
439 among regional mungbean accessions that could be used as an invaluable genetic resource for
440 genetic improvement in the future.

441 **Materials and Methods**

442 **Plant materials and SNP genotyping**

443 A total of 290 cultivated mungbean (*V. radiata* var. *radiata*) accessions were provided by the
444 Vavilov Institute (VIR). Most of the accessions are mainly landraces collected during 1910-1960
445 are considered these accessions as oldest cultivated mungbean collection from VIR (Burlyaeva
446 et al., 2019). The term landrace, as we use it here, refers to locally adaptive accessions coming
447 from the countries traditionally cultivating them, which also lacks modern genetic improvement.
448 The complete list of materials can be found in Supplementary file 1. Genomic DNA was extracted
449 from a single plant per accession using the QIAGEN Plant Mini DNA kit according to the
450 manufacturer's instruction with minor modification of pre-warming the AP1 buffer to 65°C and
451 increasing the incubation time of the P3 buffer up to 2 hours on ice to increase DNA yield. DNA
452 samples were sent to Diversity Arrays Technology Pty Ltd, Canberra, Australia for diversity array
453 technology sequence (DArTseq) genotyping.

454 DArTseq data of 521 accessions from the Australian Diversity Panel (ADP) (Noble et al.,
455 2018) and 297 accessions from the World Vegetable Center (WorldVeg) mini-core (Breria et al.,
456 2020) were also included in this study. In total, our dataset contains more than one thousand
457 accessions (1,092) and covers worldwide diversity of cultivated mungbean representing a wide
458 range of variation in seed colour (Figure 1A). Sixteen wild mungbean (*V. radiata* var. *sublobata*)
459 accessions were included as an outgroup. While all accessions used in this study have the country
460 of origin information, only those from VIR have detailed longitude and latitude information.
461 Therefore, for analyses connecting genetic information and detailed location (the isolation by
462 distance analyses), only the VIR samples were used.

463 The major goal of this study is to investigate the patterns of population expansion and the
464 underlying ecological causes instead of detailed haplotype analyses of specific genomic regions.
465 For this goal, genomewide SNPs provide similar information as whole-genome sequencing, as
466 have been shown in other species. Compared to other genotyping-by-sequencing technologies,
467 DArTseq has the additional advantage of less missing data among loci or individuals, providing
468 a more robust estimation of population structure.

469 **SNP calling**

470 Trimmomatic version 0.38 (Bolger et al., 2014) was used to remove adapters based on the
471 manufacturer's adapter sequences. Reads for each accession were trimmed for low-quality bases
472 with quality scores of $Q \leq 10$ using SolexaQA version 3.1.7.1 (Cox et al., 2010) and mapped to
473 the mungbean reference genome (Vradiata_ver6, (Kang et al., 2014) using the Burrows-Wheeler
474 Aligner (BWA) version 0.7.15 (Li and Durbin, 2009). Reads were then sorted and indexed using
475 samtools version 1.4.1 (Li et al., 2009). We used Genome Analysis Toolkit (GATK) version 3.7-
476 0-gcfedb67 (McKenna et al., 2010) to call all sites, including variant and invariant sites. We
477 obtained 1,247,721 sites with a missing rate of $< 10\%$ and a minimum quality score of 30. SNP
478 calling was performed using GATK (McKenna et al., 2010). Finally, we used VCFtools version
479 0.1.13 (Danecek et al., 2011) to remove SNPs with more than two alleles and 10% missing data,
480 resulting in 34,469 filtered SNPs. To reduce non-independence caused by linkage disequilibrium
481 (LD) among SNPs, SNPs were pruned based on a 50-SNP window with a step of 5 SNPs and r^2
482 threshold of 0.5 in PLINK (Purcell et al., 2007). This dataset of 10,359 LD-pruned SNPs (10%
483 missing data) was applied for all analyses related to population genomics unless otherwise noted.
484 For TreeMix that require LD-pruned SNPs with no missing dataset, we used 4,396 LD-pruned
485 SNPs with no missing data.

486 **Population genetics and differentiation analyses**

487 Population structure was investigated based on 10,359 LD-pruned SNPs using ADMIXTURE
488 (Alexander et al., 2009) with the number of clusters (K) ranging from 1 to 10. The analyses were
489 run ten times for each K value, and cross-validation (CV) error was used to obtain the most
490 probable K value for population structure analysis. ADMIXTURE plots were generated using
491 "Pophelper" in R (Francis, 2017). Genetic groups of accessions were assigned based on ancestry
492 coefficient $Q \geq 0.7$, otherwise the accession was considered admixed. The population structure

493 was also examined with principal component analysis (PCA). The neighbor-joining phylogenetic
494 tree was calculated using TASSEL (Trait Analysis by aSSociation, Evolution and Linkage)
495 software version 5.2.60 (Bradbury et al., 2007) and visualized using FigTree version 1.4.4
496 (<http://tree.bio.ed.ac.uk/software/figtree/>).

497 The relationships and gene flow among the four inferred genetic groups were further
498 assessed by TreeMix version 1.12 (Pickrell and Pritchard, 2012) using 4,396 LD-pruned SNPs
499 with no missing data. The analysis was run for 0 to 3 migration events with *V. radiata* var.
500 *sublobata* as an outgroup with a block size of 20 SNPs to account for the effects of LD between
501 SNPs. We estimated one as the optimal number of migration events using the “OptM” in R (Fitak,
502 2021). Bootstrap support for the resulting observed topology was obtained using 500 bootstrap
503 replicates.

504 Nucleotide diversity (π) and genetic differentiation (d_{xy} and F_{ST}) were estimated in 10 kb
505 windows with pixy version 1.2.7.beta1 (Korunes and Samuk, 2021) using all 1,247,721 invariant
506 and variant sites. LD decay for each genetic group was estimated based on 34,469 non-LD-pruned
507 SNPs using PopLDdecay (Zhang et al., 2019). The curves were fitted by a LOESS function and
508 an LD decay plot was drawn using R.

509 To investigate the relation among inferred genetic groups, f_3 and f_4 statistics were
510 computed based on filtered SNPs using ADMIXTOOLS version 7.0 (Patterson et al., 2012). The
511 f_3 statistic compares allele frequencies in two populations (A, B) and a target population C. In
512 “outgroup f_3 statistic”, C is the outgroup and positive values represent the shared genetic drift
513 between A and B. In “admixture f_3 statistic”, negative values indicate that the C is admixed from
514 A and B. For f_4 statistics, $f_4(A, B; C, D)$ measures the shared genetic drift between B populations
515 and C and D after their divergence from outgroup A. A positive value indicates that the B
516 population shares more alleles with D, and a negative value indicates that the B population shares
517 more alleles with C. We used two Mb as a unit of block-jackknife resampling to compute standard
518 errors. The Z-scores with absolute values greater than three are considered statistically significant.

519 To examine the role of geographic distance in shaping spatial genetic differentiation,
520 Mantel tests with 1,000 permutations were performed for each of the ADMIXTURE-inferred
521 genetic groups (separately for the groups defined by $Q \geq 0.7$ or $Q \geq 0.5$) using “ade4” in R.

522 Pairwise genetic distance between accessions was estimated based on all sites while the great
523 circle geographic distance was determined using “fields” in R. In addition, the same analysis was
524 conducted for southern and northern groups to examine if there was a south-north pattern of
525 differentiation.

526 Based on the shape of the phylogenetic tree, we used fastsimcoal2 (Excoffier et al., 2021),
527 which does not rely on whole-genome sequencing, to estimate the split time among genetic
528 groups. Fifty accessions were randomly picked from each genetic group. Population size was
529 allowed to change, and gene flow was allowed among populations. This analysis used all sites
530 covered by the DArT tags (including monomorphic sites), and the mutation rate was set to 1×10^{-8}
531 which was within the range of mutation rates used in eudicots (Barrera-Redondo et al., 2021;
532 Zheng et al., 2022). The models were run using unfolded site frequency spectrum using the major
533 allele in the wild progenitor population (*V. radiata* var. *sublobata*) as the ancestral allele. The
534 model was run independently 100 times, each with 100,000 simulations. After obtaining the run
535 with the highest likelihood, we performed parametric bootstrapping 100 times to obtain the 75%
536 confidence intervals of each parameter based on the previous study of Gutaker et al. (2020).

537 **Ecological niche modelling (ENM)**

538 To understand whether the habitats of genetic groups are differentiated, 248 sampling sites (82
539 for East Asia, 45 for Southeast Asia, 49 for South Asia and 72 for Central Asia genetic groups),
540 in combination with additional presence records obtained from the Global Biodiversity
541 Information Facility (GBIF, <https://www.gbif.org/>), were used for the analysis. Using the
542 longitude and latitude information, we extracted the Köppen climate zones (Köppen, 2011) using
543 “kgc” in R (Bryant et al., 2017). After excluding zones with less than five samples, the remaining
544 ten zones were grouped into six categories based on climate similarity: dry hot (BSh and BWh),
545 dry cold (BSk and BWk), temperate dry summer (Csa), tropical savanna (Aw), Continental (Dwb
546 and Dfb), and temperate wet summer (Cfa and Cwa). The former three are relatively dry
547 environments.

548 Climate layers comprising monthly minimum, maximum, mean temperature,
549 precipitation, and 19 bioclimatic variables were downloaded from the WorldClim database
550 version 1.4 (Hijmans et al., 2005). All climate layers available from WorldClim were created

551 based on climate conditions recorded between 1960 and 1990 at a spatial resolution of 30 arc-
552 seconds (approximately 1 km²). To minimize redundancy and model overfitting, pairwise Pearson
553 correlations between the 19 bioclimatic variables were calculated using ENMTools version 1.4.4
554 (Warren et al., 2010), excluding one of the two variables that has a correlation above 0.8
555 (Supplementary file 6). As a result, eight bioclimatic variables were used for all further analyses,
556 including Bio1 (annual mean temperature), Bio2 (mean diurnal range), Bio3 (isothermality), Bio8
557 (mean temperature of wettest quarter), Bio12 (annual precipitation), Bio14 (precipitation of driest
558 month), Bio15 (precipitation seasonality) and Bio19 (precipitation of coldest month). Bioclimatic
559 variables were extracted for each occurrence point using “raster” in R (Hijmans, 2021). PCA and
560 multivariate analysis of variance (MANOVA) were conducted to examine whether there was a
561 significant habitat difference among genetic groups. Ecological niche modelling (ENM) was
562 performed using MAXENT version 3.3.1 (Phillips et al., 2006) to predict the geographic
563 distribution of suitable habitats for cultivated mungbean. The ENM analysis was run with a
564 random seed, a convergence threshold of 5,000 and 10-fold cross-validation. As a measure of the
565 habitat overlaps of the four genetic groups, pairwise of Schoener’s D was calculated using
566 ENMTools. The value ranges from 0 (no niche overlap) to 1 (niche complete overlap). In addition,
567 we carried out the same analyses using monthly temperature and precipitation from May, July,
568 and September.

569 **Field evaluation**

570 Among the 52 accessions used for laboratory experiments, phenotyping of 49 accessions was
571 conducted at WorldVeg, Taiwan in 1984 and 2018 and at Crop Sciences Institute, National
572 Agricultural Research Centre, Pakistan in 2015. The traits related to phenology (days to 50%
573 flowering), reproduction (100 seed weight, pod length, pods per plant, 1000 seed weight, seeds
574 yield per plant, and seeds per pod), and plant size (petiole length, plant height, plant height at
575 flowering, plant height at maturity, primary leaf length, primary leaf width, terminal leaflet length,
576 and terminal leaflet width) were included. Trait values were inverse normal transformed.
577 The analysis of variance (ANOVA) was performed to test for inferred genetic groups differences
578 for each trait using R software (version 4.1.0).

579 **Drought phenotyping**

580 A total of 52 accessions with ancestry coefficients $Q \geq 0.7$ from three genetic groups (Southeast
581 Asia, South Asia, and Central Asia) were selected for experiments of seedling-stage drought
582 response. The experiment was laid out in a completely randomized design with three replicates
583 of each accession under two treatments (control/drought). The experiment was conducted in two
584 independent batches, and the whole experiment included 624 plants (52 accessions x 2 treatments
585 x 3 plants per treatment x 2 batches).

586 Mungbean seeds were surface-sterilized with 10% bleach for 10 mins and rinsed with
587 distilled water for three times. Seeds were treated with 70% ethanol for 5 mins and washed three
588 times in distilled water. The sterilized seeds were germinated on wet filter paper in petri dishes
589 for 3 days. The experiment was conducted in a 740FLED-2D plant growth chamber (HiPoint,
590 Taiwan) at a temperature of $25 \pm 1^\circ\text{C}$ and 12 hours of photoperiod (light ratios of red: green: blue
591 3: 1: 1) with light intensity $350 \mu\text{mol m}^{-2}\text{s}^{-1}$ and relative humidity at $60 \pm 5\%$. The seedlings were
592 then transplanted to a hydroponic system with half-strength Hoagland nutrient solution
593 (Phytotechnology Laboratory, USA) and were grown for six days before drought stress started.
594 The nutrient solution was changed on alternate days and the pH of the solution was adjusted to
595 6.0 with 1M KOH or 1M HCl.

596 For drought treatment, seedlings of mungbean were exposed to polyethylene glycol
597 (PEG)-induced drought stress for five days. The solution of PEG6000 with an osmotic potential
598 of -0.6 MPa was prepared by adding PEG6000 (Sigma-Aldrich, Germany) to the nutrient solution
599 according to Michel and Kaufmann (1973), and pH was also adjusted to 6.0. The seedlings grown
600 with the nutrient solution under the same environmental conditions were considered as controls.

601 At the end of the experiment, plants were evaluated for shoot dry weight (SDW) and root
602 dry weight (RDW), measured on digital balance after oven-drying at 70°C for 48 hours. All traits
603 were analysed by mixed-model ANOVA with the treatment (control/drought) and the genetic
604 group as fixed effects. The models included accessions as a random effect nested within genetic
605 groups and a random effect of batches. Tukey's test was conducted to compare genetic groups.
606 All statistics were performed using JMP v3.0.0 (SAS Institute, 2016).

607 ***Q_{ST}-F_{ST} comparisons***

608 For each trait, quantitative trait divergence (Q_{ST}) was calculated separately with respect to each
609 treatment. Our root and shoot weight experiment used a selfed-progeny design, using the self-
610 fertilized seeds from each accession as replicates, as recommended for partially inbred species
611 (Goudet and Buchi, 2006). For the selfed-progeny design of inbred species, (Equation 1) $Q_{ST} =$
612 $V_B / (V_B + V_{Fam})$, where V_B is the among-population variance component and V_{Fam} is the within-
613 population among-family variance component (Goudet and Buchi, 2006). Variance components
614 were estimated using a model with genetic groups, accessions nested within genetic groups, and
615 batches as random factors. To accommodate the possibility that mungbean is not completely
616 selfing, we also applied (Equation 2) $Q_{ST} = (1 + f)V_B / ((1 + f)V_B + 2V_{AW})$ (Goudet and Buchi,
617 2006), where f is the inbreeding coefficient (estimated by VCFtools as 0.8425), V_B is the among-
618 population variance component, and V_{AW} is the additive genetic variance within genetic groups
619 estimated by the kinship matrix using TASSEL software (Bradbury et al., 2007). The results and
620 conclusions are similar to our previous version. The F_{ST} was calculated only using accessions in
621 the phenotyping experiment.

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637 **Data Availability**

638 Sequences generated in this study are available under NCBI BioProject PRJN809503. Accession
639 names, GPS coordinates, and NCBI accession numbers of the Vavilov Institute accessions are
640 available under Supplementary file 1. Plant trait data are available at Dryad
641 <https://doi.org/10.5061/dryad.d7wm37q3h>. Sequences and accession information of the World
642 Vegetable Centre mini-core and the Australian Diversity Panel collections were obtained from
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- 939

940 **Supplementary note.** Text analysis and translation of ancient Chinese texts regarding
941 mungbean

942

943 Qimin Yaoshu (齊民要術, about 544 AD)

944 Qimin Yaoshu, compiled by Sixie Jia (賈思勰), is one of the earliest and most complete
945 agricultural sources in China, detailing agricultural techniques near the lower reaches of Yellow
946 River at that era. This is the earliest record of mungbean in China, demonstrating mungbean has
947 reached northern China at that time and is consistent with our estimates of population divergence
948 time. The popularity of mungbean is demonstrated by it being mentioned multiple times under
949 different contexts, most notably as a green manure:

950 「若糞不可得者，五六月中，概種菘豆，至七月、八月，犁掩殺之。如以糞糞田，
951 則良美與糞不殊，又省功力。」

952 Translation: “Should feces be unavailable, during May and June one could grow mungbean.
953 Until July or August, one could plow mungbean plants into the soil. This is equivalent to using
954 feces to manure the land. This is as good as using feces and saves efforts.”

955 Notice that the months used in ancient China are slightly different from the Gregorian
956 calendar.

957

958 Xiangshan Yelu (湘山野錄, 1068-1077 AD)

959 Xiangshan Yelu was written by a monk, Wen-Ying (文瑩), recording anecdotes during
960 that era. Its records about the Emperor Zhenzong of Song (宋真宗, 968-1022 AD) detailed the
961 phenotypes of Indian mungbean at that time:

962 「真宗深念稼穡，聞占城稻耐旱、西天綠豆子多而粒大，各遣使以珍貨求其種。占
963 城得種二十石，至今在處播之。西天中印土得菘豆種二石，不知今之菘豆是否？」

964 Translation: “Zhenzong of Song deeply concerned about agriculture. He heard Champa rice
965 being drought tolerant and mungbean from India produce numerous and large seeds. Diplomats
966 were sent to exchange the seeds with treasure. Twenty dans of Champa rice were obtained and
967 propagated everywhere. Two dans of mungbean were obtained from India, but it is unclear
968 whether the mungbean today descended from these.”

969 “Dan” (石) is a unit of volume in ancient China and is called “Koku” in Japanese. The
970 exact amount varied with time.

971 The texts provide us with two pieces of important information. First, mungbean from
972 South Asia (likely also includes the SEA genetic groups if accessions near eastern India and
973 Bangladesh were included) at that time had higher yield and larger seeds than native mungbean
974 accessions in northern China, consistent with our results on trait divergence. Second, compared
975 to the clear success of Champa rice in China, it was unclear whether those southern accessions
976 had prospered in northern China, likely suggesting an unsuccessful introduction of southern high-
977 yield and large-seeded accessions to the north.

978

979 Tiangong Kaiwu (天工開物, 1637 AD)

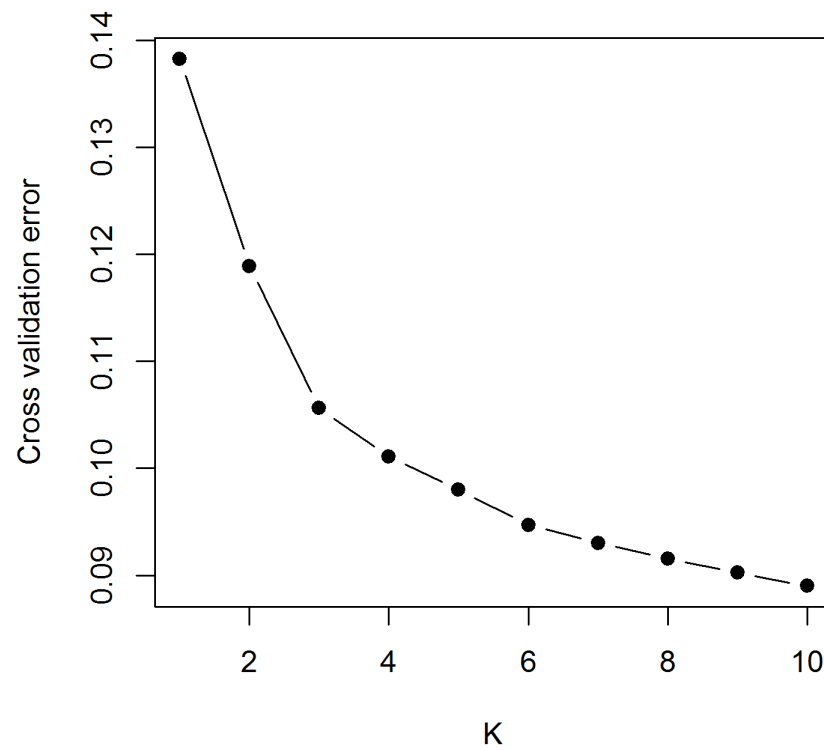
980 Tiangong Kaiwu is a famous Chinese encyclopedia compiled by Song Yingxing (宋應
981 星). While it mostly covers technologies at that time, a section about agricultural practices covers
982 mungbean:

983 「綠豆必小暑方種，未及小暑而種，則其苗蔓延數尺，結莢甚稀。若過期至於處
984 暑，則隨時開花結莢，顆粒亦少。」

985 Translation: “Mungbean must be sown at or after Xiaoshu (Gregorian 7-8 July). Being
986 sown before Xiaoshu, mungbean stems would spread for meters with few pods set. Being sown
987 as late as Chushu (Gregorian 23-24 August), the plants would flower and set pods at any time,
988 also with low yield.”

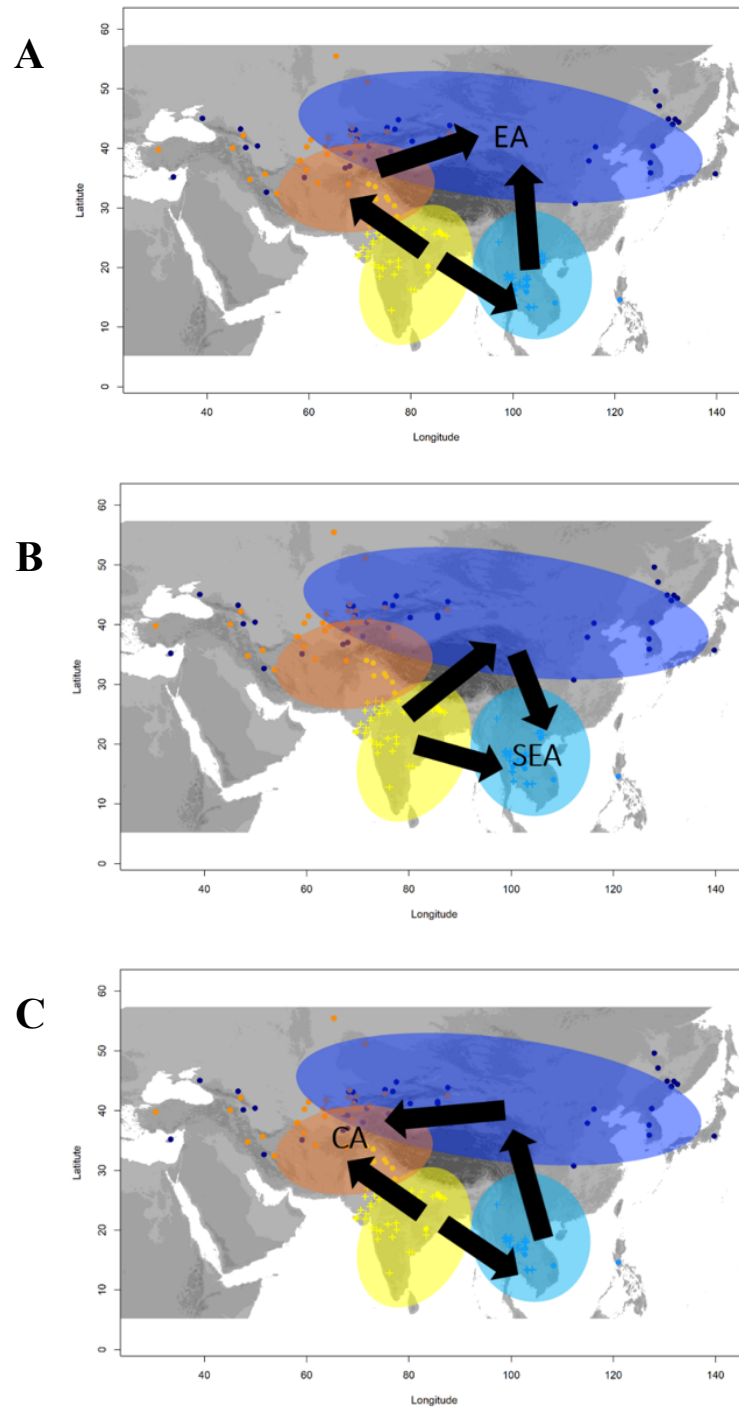
989 As a short-day plant, being sown too early when the days are too long, mungbean would
990 have mostly vegetative growth. Being sown too late when the days are too short, flowering would
991 be induced too quickly before sufficient vegetative development. In addition to our results that
992 short-season accessions were favored in the north due to the requirement for drought escape, this
993 source provides us with another support that mungbean could only be sown in a narrow time
994 window due to daylength requirement. Given the autumn frost damage in the north, not being
995 able to be sown earlier restricts the growing season length in the north, limiting the adoption of
996 southern long-season accessions.

997



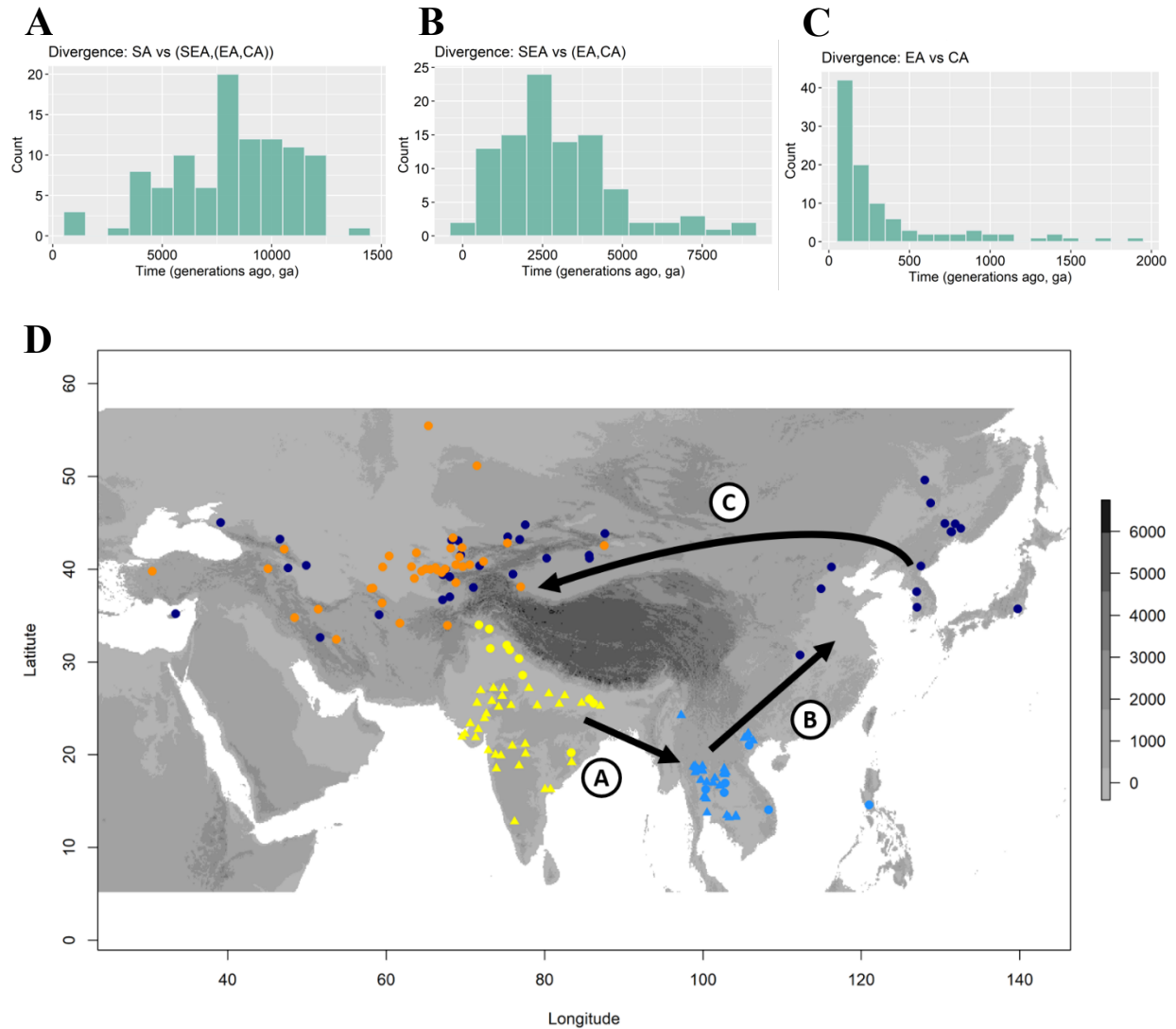
998 **Figure 1-figure supplement 1.** Cross-validation (CV) errors of ADMIXTURE. Means of CV errors
999 were calculated based on K values ranging 1 to 10 with 10 independent runs.

1000

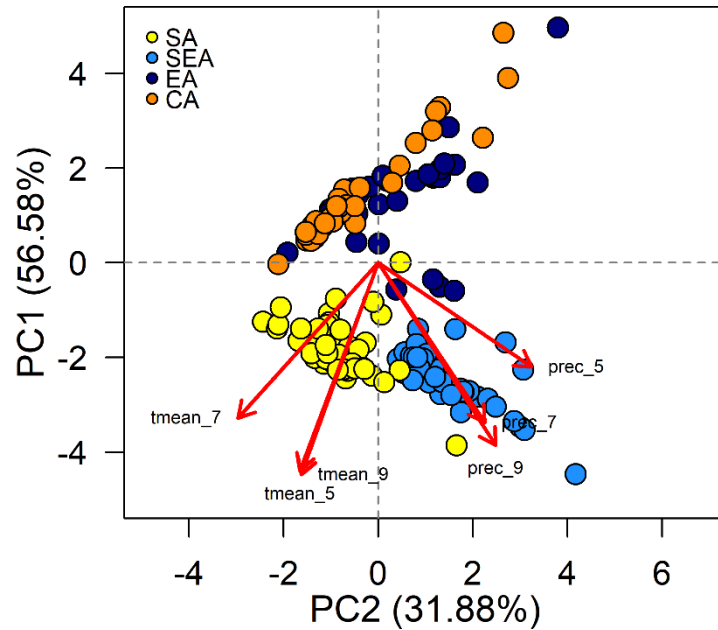


1001 **Figure 2-figure supplement 1.** Schematic representation to investigate presence of admixture in
1002 a target population from two source populations using admixture f_3 -statistics. (A) $f_3(\text{EA}; \text{SEA},$
1003 $\text{CA})$, (B) $f_3(\text{SEA}; \text{SA}, \text{EA})$ and (C) $f_3(\text{CA}; \text{EA}, \text{SA})$. Coloured circles indicate the geographic
1004 area occupied by distinct genetic groups. Arrows indicates the possible direction of expansion
1005 and admixture among populations.

1006

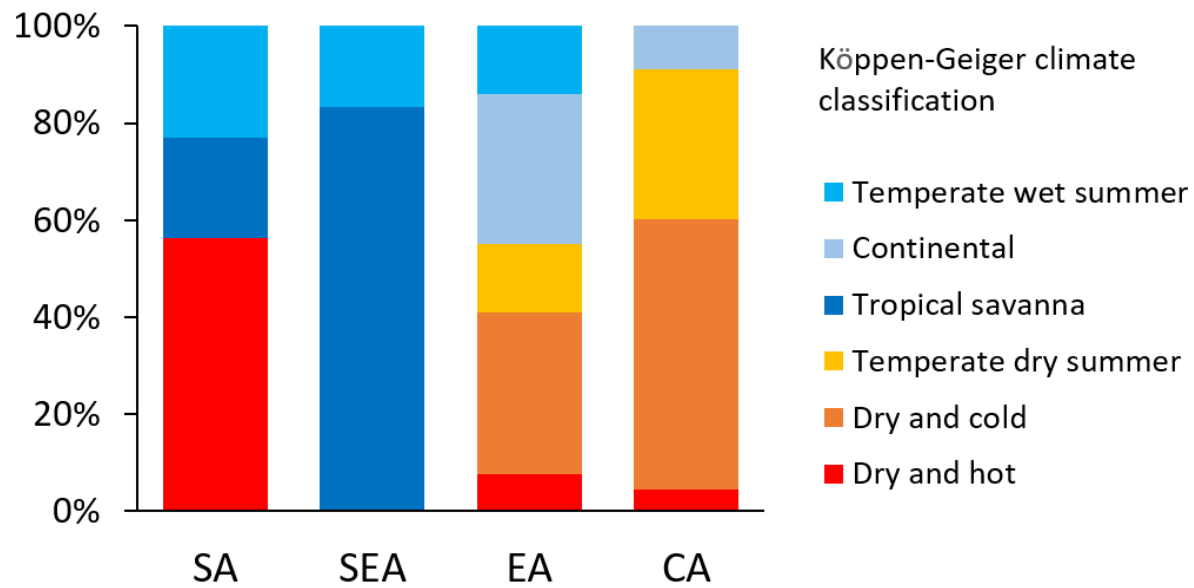


1007 **Figure 2-figure supplement 2.** Estimates of divergence time and inferred mungbean movement
1008 over time across Asia. The histograms of the divergence times represent (A) split time between
1009 SA and (SEA,(EA,CA)), (B) split time between SEA and (EA,CA), and (C) split time between
1010 EA and CA. (D) Geographic distribution of mungbean accessions and proposed mungbean spread
1011 routes. Exact locations for VIR accessions (filled circle) and GBIF records (filled triangle) are
1012 provided. Each accession was coloured the same as an inferred genetic group using
1013 ADMIXTURE in Figure 1. Arrow indicates the possible expansion directions. The map was
1014 shaded as a gray colour representing altitude (meters above sea level).
1015

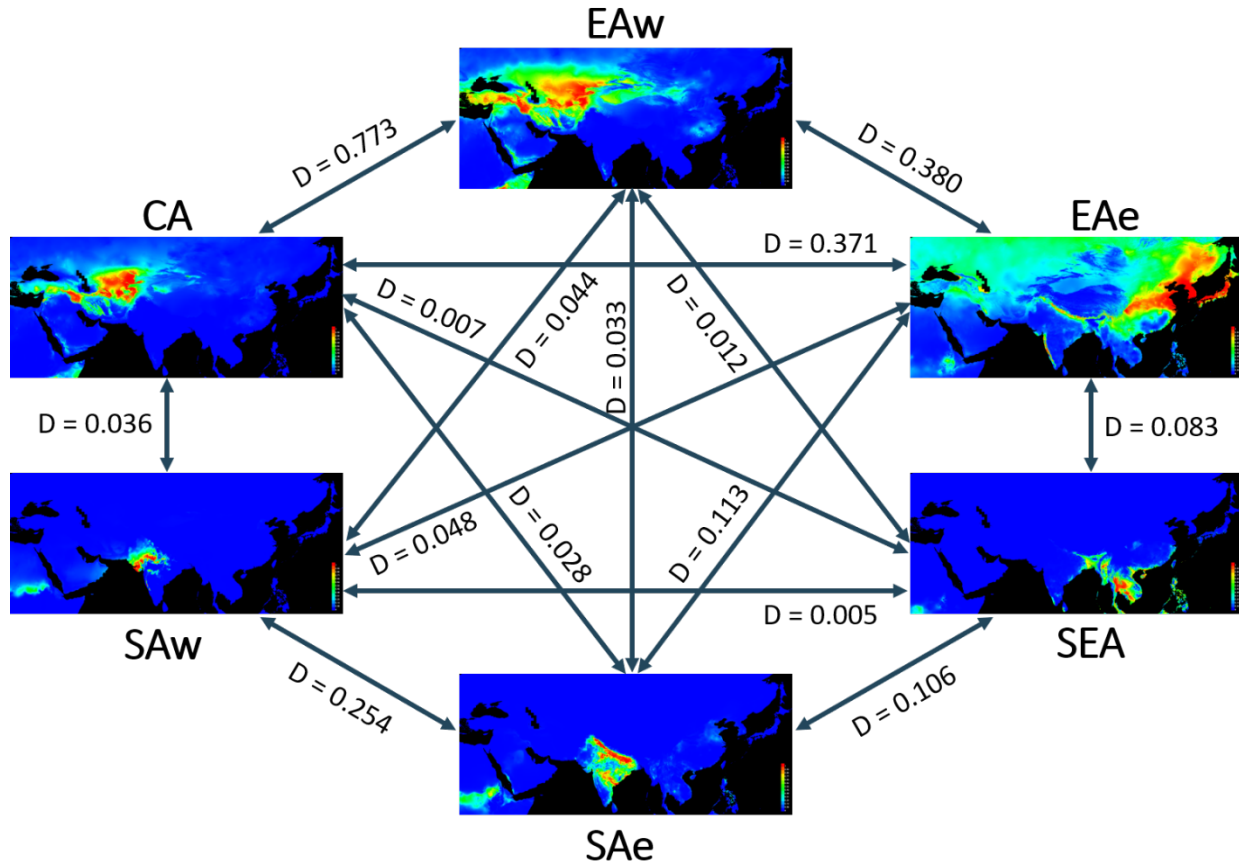


1016 **Figure 3-figure supplement 1.** Principal component analysis (PCA) of the growing season
1017 climatic data including temperature and precipitation of May, July and September. Samples are
1018 coloured according to four inferred genetic groups, as indicated in the legend.

1019

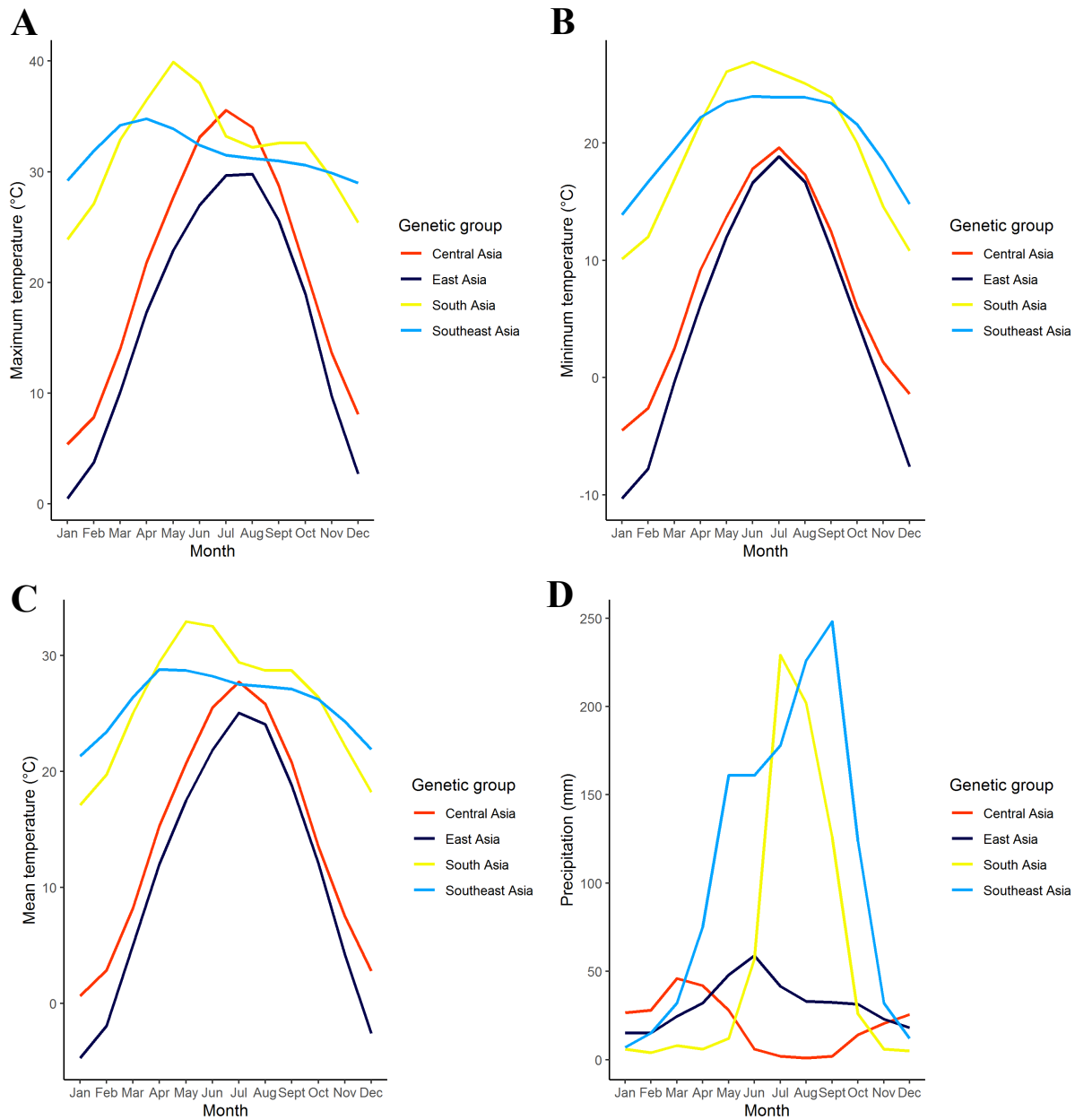


1020 **Figure 3-figure supplement 2.** The distribution of accessions in major climate zones according
1021 to the Köppen climate classification (Köppen, 2011).
1022



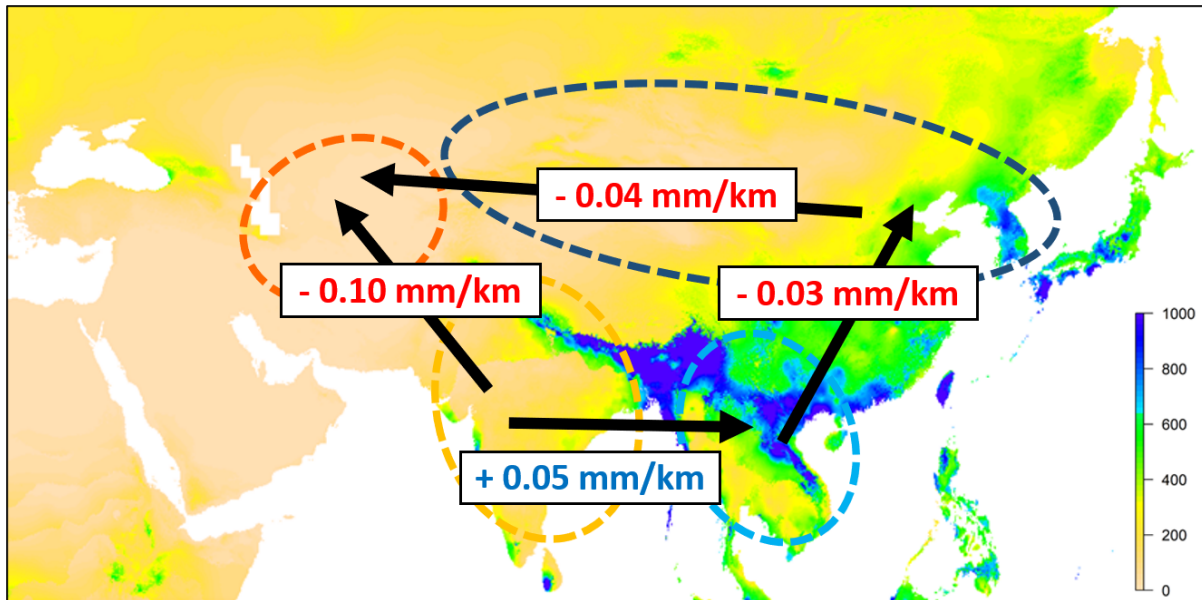
1023 **Figure 3-figure supplement 3.** Predicted distributions of six groups based on monthly
1024 temperature and precipitation (May, July and September) during the summer growing season.
1025 Red colour indicates high suitability and blue indicates low suitability. Values between groups
1026 represent niche overlap measured using Schoener's D. Abbreviations: SAw: South Asia (west),
1027 SAe: South Asia (east); SEA: Southeast Asia; EAe: East Asia (east); EAw: East Asia (west) and
1028 CA: Central Asia.

1029



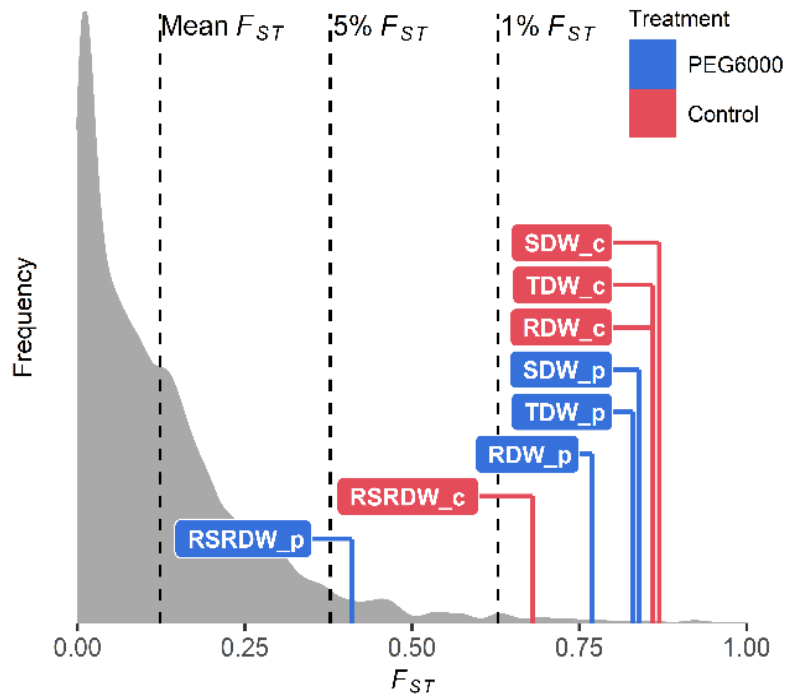
1030 **Figure 3-figure supplement 4.** Monthly temperature and precipitation variations among the four
1031 genetic groups. Monthly (A) maximum temperature, (B) minimum temperature, (C) mean
1032 temperature and (D) precipitation were computed based on median value among all accessions
1033 of a group. Genetic group were coloured the same as in Figure. 1.

1034



1035 **Figure 3-figure supplement 5.** Environmental gradient across Asia. The value on each arrow
1036 indicates a change in mean precipitation for May, July and September (growth season) per
1037 kilometer. The background map is colored according to summer precipitation (Bio18,
1038 precipitation of warmest quarter, in mm).

1039



1040 **Figure 3-figure supplement 6.** Comparison of Q_{ST} - F_{ST} for four drought-related traits under two
1041 environments. F_{ST} values (mean, 5% and 1%) were indicated by black dashed lines. The Q_{ST} for
1042 each trait was colored according to treatment and was calculated as Equation 1 in Materials and
1043 Methods. Abbreviations: RDW: root dry weight; SDW: shoot dry weight; TDW: total dry weight;
1044 RSRDW: root:shoot ratio dry weight; c: control; p: PEG6000.

1045

1046 **Supplementary file 1.** Mungbean accessions from Vavilov Institute (VIR) collection
 1047

Sample name	Country	Latitude	Longitude	NCBI SRA accession	NCBI Biosample accession
1.B	USA	NA	NA	SRR18125483	SAMN26179197
100.A	Cyprus	35.1856	33.3823	SRR18125482	SAMN26179198
101.A	Morocco	33.9716	-6.8498	SRR18125266	SAMN26179199
102.A	Israel	32.4971	35.4973	SRR18125200	SAMN26179200
103.A	Indonesia	-6.5971	106.806	SRR18125367	SAMN26179201
104.B	Russia	44.9003	131.8351	SRR18125356	SAMN26179202
105.A	Chile	-36.8305	-73.1167	SRR18125345	SAMN26179203
106.A	India	22.5587	88.2911	SRR18125298	SAMN26179204
107.A	India	22.5726	88.3639	SRR18125287	SAMN26179205
108.A	India	25.9821	85.6486	SRR18125276	SAMN26179206
109.A	Italy	41.9028	12.4964	SRR18125481	SAMN26179207
10A	China	47.1216	128.7382	SRR18125398	SAMN26179208
110.B	Portugal	38.7223	-9.1393	SRR18125387	SAMN26179209
111.A	Portugal	38.7223	-9.1393	SRR18125376	SAMN26179210
112.A	Russia	45.0347	39.0978	SRR18125329	SAMN26179211
113.A	Tajikistan	40.2675	69.6453	SRR18125318	SAMN26179212
114.B	India	31.5204	74.3587	SRR18125307	SAMN26179213
115.B	India	31.5204	74.3587	SRR18125476	SAMN26179214
116.A	Russia	49.6152	127.9945	SRR18125465	SAMN26179215
116.B	Russia	49.6152	127.9945	SRR18125454	SAMN26179216
117.B	Russia	44.0281	131.3273	SRR18125264	SAMN26179217
118.A	Russia	44.0118	131.3835	SRR18125253	SAMN26179218
119.A	Russia	44.39	132.558	SRR18125242	SAMN26179219
11A	China	47.1216	128.7382	SRR18125231	SAMN26179220
120.A	Argentina	-32.8895	-68.8458	SRR18125434	SAMN26179221
122.A	Japan	31.5969	130.5571	SRR18125423	SAMN26179222
123.B	Kyrgyzstan	55.4649	65.3054	SRR18125412	SAMN26179223
124.B	Ethiopia	12.9545	36.1573	SRR18125223	SAMN26179224
125.A	China	42.5246	87.5396	SRR18125212	SAMN26179225
126.A	China	42.5246	87.5396	SRR18125201	SAMN26179226
127.A	China	42.5246	87.5396	SRR18125199	SAMN26179227
128.A	Democratic Republic of the Congo	1.9293	30.0492	SRR18125198	SAMN26179228
128.B	Democratic Republic of the Congo	1.9293	30.0492	SRR18125197	SAMN26179229
129.A	Korea	37.5665	126.978	SRR18125196	SAMN26179230
12A	China	47.1216	128.7382	SRR18125195	SAMN26179231

13.A	Iran	32.4279	53.688	SRR18125194	SAMN26179232
130.A	Korea	37.5665	126.978	SRR18125371	SAMN26179233
131.A	Korea	37.5665	126.978	SRR18125370	SAMN26179234
132.B	Korea	40.3399	127.5101	SRR18125369	SAMN26179235
133.A	Korea	40.3399	127.5101	SRR18125368	SAMN26179236
134.A	Korea	40.3399	127.5101	SRR18125366	SAMN26179237
135.A	Korea	40.3399	127.5101	SRR18125365	SAMN26179238
136.A	Korea	40.3399	127.5101	SRR18125364	SAMN26179239
137.A	China	42.5246	87.5396	SRR18125363	SAMN26179240
138.A	China	39.4677	75.9938	SRR18125362	SAMN26179241
139.A	Uzbekistan	41.2995	69.2401	SRR18125361	SAMN26179242
14.B	Uzbekistan	39.7681	64.4556	SRR18125360	SAMN26179243
140.B	China	42.5246	87.5396	SRR18125359	SAMN26179244
141.A	China	43.8256	87.6168	SRR18125358	SAMN26179245
142.B	China	42.5246	87.5396	SRR18125357	SAMN26179246
144.A	Ukraine	48.4647	35.0462	SRR18125355	SAMN26179247
145.A	Brazil	-22.9329	-47.0738	SRR18125354	SAMN26179248
146.B	Kazakhstan	43.222	76.8512	SRR18125353	SAMN26179249
147.A	Turkey	39.7646	30.4559	SRR18125352	SAMN26179250
148.B	Turkey	39.7646	30.4559	SRR18125351	SAMN26179251
149.A	Ukraine	48.0386	30.9497	SRR18125350	SAMN26179252
151.A	Tajikistan	38.5598	68.787	SRR18125349	SAMN26179253
154.A	Brazil	-14.235	-51.9253	SRR18125348	SAMN26179254
155.A	United Kingdom	52.3555	-1.1743	SRR18125347	SAMN26179255
156.A	India	25.9821	85.6486	SRR18125346	SAMN26179256
157.A	India	25.9821	85.6486	SRR18125344	SAMN26179257
158.A	India	25.9821	85.6486	SRR18125343	SAMN26179258
159.B	India	25.9821	85.6486	SRR18125342	SAMN26179259
16.A	Uzbekistan	39.7681	64.4556	SRR18125341	SAMN26179260
160.A	Uzbekistan	40.8154	72.2837	SRR18125340	SAMN26179261
161.A	Uzbekistan	40.8154	72.2837	SRR18125339	SAMN26179262
162.A	South Africa	-26.7145	27.097	SRR18125338	SAMN26179263
163.A	South Africa	-26.7145	27.097	SRR18125337	SAMN26179264
164.A	South Africa	-26.7145	27.097	SRR18125336	SAMN26179265
165.B	USA	39.9526	-75.1652	SRR18125299	SAMN26179266
166.A	USA	39.9526	-75.1652	SRR18125297	SAMN26179267
167.B	Kyrgyzstan	42.8224	75.3179	SRR18125296	SAMN26179268
168.A	Kyrgyzstan	42.8224	75.3179	SRR18125295	SAMN26179269
17.B	Uzbekistan	39.7681	64.4556	SRR18125294	SAMN26179270
170.A	Russia	43.2562	46.5893	SRR18125293	SAMN26179271
171.A	Russia	43.2562	46.5893	SRR18125292	SAMN26179272

172.B	Iran	32.4279	53.688	SRR18125291	SAMN26179273
173.A	Iran	32.4279	53.688	SRR18125290	SAMN26179274
174.A	Korea	40.3399	127.5101	SRR18125289	SAMN26179275
175.A	Uzbekistan	40.2504	63.2032	SRR18125288	SAMN26179276
175.B	Uzbekistan	40.2504	63.2032	SRR18125286	SAMN26179277
176.A	Uzbekistan	40.2504	63.2032	SRR18125285	SAMN26179278
177.A	Uzbekistan	40.2504	63.2032	SRR18125284	SAMN26179279
178.A	Uzbekistan	40.2504	63.2032	SRR18125283	SAMN26179280
179.A	Uzbekistan	39.9208	66.4271	SRR18125282	SAMN26179281
18.B	Uzbekistan	39.7681	64.4556	SRR18125281	SAMN26179282
180.A	Uzbekistan	39.9208	66.4271	SRR18125280	SAMN26179283
181.A	Uzbekistan	39.9208	66.4271	SRR18125279	SAMN26179284
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183.A	Uzbekistan	40.3734	71.7978	SRR18125277	SAMN26179286
184.A	China	41.482754	85.626702	SRR18125275	SAMN26179287
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188.B	China	41.1675	80.2634	SRR18125273	SAMN26179289
189.A	China	41.1675	80.2634	SRR18125272	SAMN26179290
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191.A	China	41.175324	85.660861	SRR18125269	SAMN26179293
192.A	India	31.8183	75.2071	SRR18125268	SAMN26179294
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194.A	India	30.3752	76.7821	SRR18125480	SAMN26179297
195.A	India	28.7041	77.1025	SRR18125407	SAMN26179298
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197.A	India	17.6599	75.9064	SRR18125405	SAMN26179300
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203.A	Ethiopia	9.3126	42.1227	SRR18125400	SAMN26179305
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205.A	Indonesia	-6.5971	106.806	SRR18125397	SAMN26179307
205.B	Indonesia	-6.5971	106.806	SRR18125396	SAMN26179308
206.A	Hungary	47.1625	19.5033	SRR18125395	SAMN26179309
207.A	Viet Nam	14.0583	108.2772	SRR18125394	SAMN26179310
21.B	Kazakhstan	51.1605	71.4704	SRR18125393	SAMN26179311
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215.A	Guinea	8.5383	-9.4728	SRR18125390	SAMN26179314
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217.B	Uzbekistan	40.162885	66.227209	SRR18125388	SAMN26179316
218.A	Uzbekistan	40.013465	64.943243	SRR18125386	SAMN26179317
219.A	Turkmenistan	40.243331	59.540314	SRR18125385	SAMN26179318
22.B	Kazakhstan	42.3417	69.5901	SRR18125384	SAMN26179319
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223.A	Uzbekistan	40.483568	70.546311	SRR18125382	SAMN26179321
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225.A	Pakistan	30.3753	69.3451	SRR18125380	SAMN26179323
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229.A	Pakistan	31.4504	73.135	SRR18125375	SAMN26179327
23.A	Kazakhstan	42.3417	69.5901	SRR18125374	SAMN26179328
230.A	Pakistan	31.4504	73.135	SRR18125373	SAMN26179329
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231.A	Tanzania	-6.369	34.8888	SRR18125335	SAMN26179331
232.A	Iran	34.3277	47.0778	SRR18125334	SAMN26179332
233.A	Iran	32.4279	53.688	SRR18125333	SAMN26179333
234.A	India	28.6139	77.209	SRR18125332	SAMN26179334
235.A	Pakistan	33.5651	73.0169	SRR18125331	SAMN26179335
235.B	Pakistan	33.5651	73.0169	SRR18125330	SAMN26179336
236.A	Pakistan	34.0155	71.6888	SRR18125328	SAMN26179337
237.A	Pakistan	28.6001	77.227	SRR18125327	SAMN26179338
238.A	Egypt	26.8206	30.8025	SRR18125326	SAMN26179339
239.A	Kazakhstan	44.7689	77.5573	SRR18125325	SAMN26179340
24.A	Uzbekistan	41.4065	60.3685	SRR18125324	SAMN26179341
240.A	Uzbekistan	39.4065	67.1418	SRR18125323	SAMN26179342
240.B	Uzbekistan	39.4065	67.1418	SRR18125322	SAMN26179343
241.A	Uzbekistan	40.023044	67.433724	SRR18125321	SAMN26179344
242.A	Uzbekistan	39.982851	67.486778	SRR18125320	SAMN26179345
244.A	Uzbekistan	41.773406	63.780613	SRR18125319	SAMN26179346
245.A	Kazakhstan	43.47491	75.335144	SRR18125317	SAMN26179347
246.A	Algeria	35.6971	-0.6308	SRR18125316	SAMN26179348
247.A	Tanzania	-5.0425	32.8197	SRR18125315	SAMN26179349
249.A	Australia	-32.9283	151.7817	SRR18125314	SAMN26179350
25.A	Uzbekistan	41.4065	60.3685	SRR18125313	SAMN26179351
250.B	Russia	NA	NA	SRR18125312	SAMN26179352
251.B	USA	NA	NA	SRR18125311	SAMN26179353
252.A	Afghanistan	34.7602	69.8121	SRR18125310	SAMN26179354

253.A	Afghanistan	34.7602	69.8121	SRR18125309	SAMN26179355
254.A	Yemen	15.5527	48.5164	SRR18125308	SAMN26179356
254.B	Yemen	15.5527	48.5164	SRR18125306	SAMN26179357
255.A	Afghanistan	36.6926	67.118	SRR18125305	SAMN26179358
256.A	Afghanistan	36.6926	67.118	SRR18125304	SAMN26179359
257.B	Afghanistan	36.6926	67.118	SRR18125303	SAMN26179360
258.B	Afghanistan	33.9391	67.71	SRR18125302	SAMN26179361
259.B	Colombia	4.5709	-74.2973	SRR18125301	SAMN26179362
26.B	Uzbekistan	41.4065	60.3685	SRR18125300	SAMN26179363
260.A	Philippines	14.5995	120.9842	SRR18125479	SAMN26179364
261.B	Kenya	-1.2921	36.8219	SRR18125478	SAMN26179365
262.A	Kenya	-1.2921	36.8219	SRR18125477	SAMN26179366
263.A	Kenya	-1.2921	36.8219	SRR18125475	SAMN26179367
264.A	Kenya	-1.2921	36.8219	SRR18125474	SAMN26179368
265.B	Kenya	-1.2921	36.8219	SRR18125473	SAMN26179369
266.A	Kenya	-1.2921	36.8219	SRR18125472	SAMN26179370
267.A	Kenya	-1.2921	36.8219	SRR18125471	SAMN26179371
268.A	Kenya	-1.2921	36.8219	SRR18125470	SAMN26179372
269.B	Kenya	-1.2921	36.8219	SRR18125469	SAMN26179373
27.A	Uzbekistan	39.7681	64.4556	SRR18125468	SAMN26179374
270.A	Kenya	-1.2921	36.8219	SRR18125467	SAMN26179375
271.A	South Korea	35.8987	127.0392	SRR18125466	SAMN26179376
272.A	Australia	NA	NA	SRR18125464	SAMN26179377
273.A	Kenya	-1.2921	36.8219	SRR18125463	SAMN26179378
274.A	Kenya	-1.2921	36.8219	SRR18125462	SAMN26179379
275.A	Kenya	-1.2921	36.8219	SRR18125461	SAMN26179380
276.A	Kenya	-1.2921	36.8219	SRR18125460	SAMN26179381
277.B	Kazakhstan	43.1521	68.2581	SRR18125459	SAMN26179382
278.B	Kazakhstan	43.0631	69.0851	SRR18125458	SAMN26179383
279.A	Kazakhstan	41.5295	69.4133	SRR18125457	SAMN26179384
28.A	Russia	42.1432	47.095	SRR18125456	SAMN26179385
280.B	Kazakhstan	41.5295	69.4133	SRR18125455	SAMN26179386
281.B	South Korea	35.8987	127.0392	SRR18125453	SAMN26179387
282.B	South Korea	35.8987	127.0392	SRR18125452	SAMN26179388
283.B	South Korea	35.8987	127.0392	SRR18125451	SAMN26179389
284.A	Tajikistan	38.0116	71.003	SRR18125450	SAMN26179390
285.A	Tajikistan	37.074793	67.957222	SRR18125449	SAMN26179391
286.A	Tajikistan	37.028926	68.004059	SRR18125448	SAMN26179392
287.A	Tajikistan	39.179338	68.012339	SRR18125447	SAMN26179393
288.A	Philippines	14.5995	120.9842	SRR18125446	SAMN26179394
289.B	Philippines	14.5995	120.9842	SRR18125445	SAMN26179395

290.A	China	40.2374	116.2305	SRR18125444	SAMN26179396
290.B	China	40.2374	116.2305	SRR18125263	SAMN26179397
291.A	China	40.2374	116.2305	SRR18125262	SAMN26179398
292.A	China	30.7378	112.2384	SRR18125261	SAMN26179399
293.A	China	30.7378	112.2384	SRR18125260	SAMN26179400
294.A	China	40.2374	116.2305	SRR18125259	SAMN26179401
295.B	China	37.8957	114.9042	SRR18125258	SAMN26179402
296.A	China	37.8957	114.9042	SRR18125257	SAMN26179403
297.A	China	47.1216	128.7382	SRR18125256	SAMN26179404
298.B	China	47.1216	128.7382	SRR18125255	SAMN26179405
299.A	China	40.2374	116.2305	SRR18125254	SAMN26179406
2A	China	47.1216	128.7382	SRR18125252	SAMN26179407
3.B	China	47.1216	128.7382	SRR18125251	SAMN26179408
30.A	Russia	42.1432	47.095	SRR18125250	SAMN26179409
300.A	China	47.1216	128.7382	SRR18125249	SAMN26179410
31.A	Russia	42.1432	47.095	SRR18125248	SAMN26179411
32.B	Russia	42.1432	47.095	SRR18125247	SAMN26179412
33.A	USA	NA	NA	SRR18125246	SAMN26179413
34.A	USA	NA	NA	SRR18125245	SAMN26179414
34.B	USA	NA	NA	SRR18125244	SAMN26179415
35.A	Russia	NA	NA	SRR18125243	SAMN26179416
36.A	Russia	43.1198	131.8869	SRR18125241	SAMN26179417
37.A	USA	NA	NA	SRR18125240	SAMN26179418
38.B	China	44.9188	130.5244	SRR18125239	SAMN26179419
39.A	Iran	36.3394	59.4698	SRR18125238	SAMN26179420
3A	China	47.1216	128.7382	SRR18125237	SAMN26179421
40.B	Iran	36.3394	59.4698	SRR18125236	SAMN26179422
41.B	Iran	32.4279	53.688	SRR18125235	SAMN26179423
42.A	Iran	32.4279	53.688	SRR18125234	SAMN26179424
43.A	Iran	32.4279	53.688	SRR18125233	SAMN26179425
44.B	Turkmenistan	37.9153	58.0897	SRR18125232	SAMN26179426
45.A	Turkmenistan	37.9153	58.0897	SRR18125230	SAMN26179427
46.A	Turkmenistan	37.9153	58.0897	SRR18125443	SAMN26179428
47.A	Turkmenistan	37.9153	58.0897	SRR18125442	SAMN26179429
48.B	Turkmenistan	37.9153	58.0897	SRR18125441	SAMN26179430
49.B	Turkmenistan	37.9172	58.0907	SRR18125440	SAMN26179431
4A	China	47.1216	128.7382	SRR18125439	SAMN26179432
5.B	China	47.1216	128.7382	SRR18125438	SAMN26179433
50.A	Turkmenistan	37.9172	58.0907	SRR18125437	SAMN26179434
51.B	Turkmenistan	37.9601	58.3261	SRR18125436	SAMN26179435
52.A	Turkmenistan	37.9601	58.3261	SRR18125435	SAMN26179436

53.B	USA	40.1605	-103.2144	SRR18125433	SAMN26179437
54.A	USA	40.1605	-103.2144	SRR18125432	SAMN26179438
55.B	Ukraine	48.3794	31.1656	SRR18125431	SAMN26179439
56.A	Kazakhstan	43.3667	68.4094	SRR18125430	SAMN26179440
57.A	Iran	34.7608	48.3988	SRR18125429	SAMN26179441
58.A	Iran	35.6892	51.389	SRR18125428	SAMN26179442
59.B	Kazakhstan	43.3667	68.4094	SRR18125427	SAMN26179443
6.B	China	47.1216	128.7382	SRR18125426	SAMN26179444
60.A	Kazakhstan	42.2663	68.1431	SRR18125425	SAMN26179445
61.A	Uzbekistan	41.2995	69.2401	SRR18125424	SAMN26179446
62.A	Uzbekistan	41.2995	69.2401	SRR18125422	SAMN26179447
63.A	Uzbekistan	41.2995	69.2401	SRR18125421	SAMN26179448
64.B	Uzbekistan	41.2995	69.2401	SRR18125420	SAMN26179449
65.B	Uzbekistan	41.2995	69.2401	SRR18125419	SAMN26179450
66.B	Uzbekistan	41.2995	69.2401	SRR18125418	SAMN26179451
67.A	Uzbekistan	41.2995	69.2401	SRR18125417	SAMN26179452
68.A	Uzbekistan	41.2995	69.2401	SRR18125416	SAMN26179453
69.A	Uzbekistan	40.4915	68.7811	SRR18125415	SAMN26179454
70.A	Uzbekistan	39.627	66.975	SRR18125414	SAMN26179455
71.A	Uzbekistan	39.627	66.975	SRR18125413	SAMN26179456
72.A	Uzbekistan	39.627	66.975	SRR18125411	SAMN26179457
73.B	Uzbekistan	39.627	66.975	SRR18125410	SAMN26179458
74.A	Uzbekistan	39.627	66.975	SRR18125409	SAMN26179459
75.A	Uzbekistan	39.627	66.975	SRR18125408	SAMN26179460
76.A	Uzbekistan	41.2995	69.2401	SRR18125229	SAMN26179461
77.A	Uzbekistan	39.7681	64.4556	SRR18125228	SAMN26179462
79.B	Uzbekistan	39.7681	64.4556	SRR18125227	SAMN26179463
7A	China	47.1216	128.7382	SRR18125226	SAMN26179464
8.B	China	47.1216	128.7382	SRR18125225	SAMN26179465
81.B	Afghanistan	34.1769	61.7006	SRR18125224	SAMN26179466
82.B	Afghanistan	34.1769	61.7006	SRR18125222	SAMN26179467
83.A	Afghanistan	34.1769	61.7006	SRR18125221	SAMN26179468
84.A	Afghanistan	34.1769	61.7006	SRR18125220	SAMN26179469
85.A	Afghanistan	34.1769	61.7006	SRR18125219	SAMN26179470
86.A	Afghanistan	36.6153	66.9293	SRR18125218	SAMN26179471
88.B	Afghanistan	NA	NA	SRR18125217	SAMN26179472
89.B	Afghanistan	33.9391	67.71	SRR18125216	SAMN26179473
90.B	Uzbekistan	NA	NA	SRR18125215	SAMN26179474
91.A	Japan	35.719	139.7456	SRR18125214	SAMN26179475
92.A	Japan	35.719	139.7456	SRR18125213	SAMN26179476
93.B	Armenia	40.0691	45.0382	SRR18125211	SAMN26179477

94.A	Iran	35.102	59.1042	SRR18125210	SAMN26179478
95.A	Azerbaijan	40.4093	49.8671	SRR18125209	SAMN26179479
96.A	Azerbaijan	40.1431	47.5769	SRR18125208	SAMN26179480
97.A	Turkmenistan	39.0041	63.5688	SRR18125207	SAMN26179481
98.B	Panama	9.3593	-79.8999	SRR18125206	SAMN26179482
99.A	Panama	9.3593	-79.8999	SRR18125205	SAMN26179483
99.B	Panama	9.3593	-79.8999	SRR18125204	SAMN26179484
9A	China	47.1216	128.7382	SRR18125203	SAMN26179485
M7.A	India	31.5204	74.3587	SRR18125202	SAMN26179486

1049 **Supplementary file 2.** Outgroup f_3 statistics among all possible combinations of genetic group
1050 pairs
1051

Outgroup (C)	Source 1 (A)	Source2 (B)	f_3	Standard error	Z-score	Significant
<i>sublobata</i>	CA	EA	0.232	0.005	42.66	Yes
<i>sublobata</i>	CA	SA	0.209	0.005	38.41	Yes
<i>sublobata</i>	CA	SEA	0.213	0.005	39.03	Yes
<i>sublobata</i>	EA	SA	0.209	0.005	38.79	Yes
<i>sublobata</i>	EA	SEA	0.218	0.005	40.75	Yes
<i>sublobata</i>	SA	SEA	0.211	0.005	39.41	Yes

1052 Abbreviations: SA, South Asia; SEA, Southeast Asia; EA, East Asia and CA, Central Asia
1053 (f_3 statistics with Z-score $> |3|$ are considered significant)
1054

1055 **Supplementary file 3.** Admixture f_3 statistics among all possible population trios
1056

Target (C)	Source1 (A)	Source2 (B)	f_3	Standard error	Z-score	Significant
EA	SA	CA	0.005	0.001	4.82	Yes
EA	SEA	CA	-0.001	0.001	-0.51	No
EA	SEA	SA	0.020	0.002	13.48	Yes
SEA	CA	EA	0.030	0.002	14.7	Yes
SEA	SA	CA	0.014	0.002	8.82	Yes
SEA	SA	EA	0.009	0.001	6.86	Yes
SA	CA	EA	0.032	0.002	16.66	Yes
SA	CA	SEA	0.011	0.001	9.02	Yes
SA	EA	SEA	0.017	0.001	12.91	Yes
CA	EA	SA	0.011	0.001	9.55	Yes
CA	EA	SEA	0.016	0.002	10.37	Yes
CA	SEA	SA	0.031	0.002	15.65	Yes

1057 Abbreviations: SA, South Asia; SEA, Southeast Asia; EA, East Asia and CA, Central Asia
1058 (f_3 statistics with Z-score > |3| are considered significant, but only negative f_3 statistics denote
1059 the target population being admixed from source1 and source2.)
1060

1061 **Supplementary file 4.** Mantel tests for isolation by distance of inferred genetic group ($Q \geq 0.5$)
1062

Group	<i>r</i>	<i>P</i>
SA	0.4319	0.008*
SEA	0.3312	0.041*
EA	0.0461	0.052
CA	0.0070	0.435
Southern	0.2934	0.001*
Northern	0.2777	0.001*

1063 Abbreviations: SA, South Asia; SEA, Southeast Asia; EA, East Asia; CA, Central Asia; *r*, Mantel
1064 correlation; significance level * $P < 0.05$
1065

1066 **Supplementary file 5.** Description of bioclimatic variables used in ecological niche modelling
1067

Bioclimatic variable	Variable	Unit
Bio1	Annual mean temperature	°C
Bio2	Mean diurnal range (mean of monthly (max temp - min temp))	°C
Bio3	Isothermality (Bio2/Bio7) ($\times 100$)	°C
Bio4	Temperature seasonality (standard deviation $\times 100$)	°C
Bio5	Max temperature of warmest month	°C
Bio6	Min temperature of coldest month	°C
Bio7	Temperature annual range (Bio5-Bio6)	°C
Bio8	Mean temperature of wettest quarter	°C
Bio9	Mean temperature of driest quarter	°C
Bio10	Mean temperature of warmest quarter	°C
Bio11	Mean temperature of coldest quarter	°C
Bio12	Annual precipitation	mm
Bio13	Precipitation of wettest month	mm
Bio14	Precipitation of driest month	mm
Bio15	Precipitation seasonality (coefficient of variation)	mm
Bio16	Precipitation of wettest quarter	mm
Bio17	Precipitation of driest quarter	mm
Bio18	Precipitation of warmest quarter	mm
Bio19	Precipitation of coldest quarter	mm

1068

1069 **Supplementary file 6.** Pearson's correlation coefficient between pairs of bioclimatic variables (denoted in lower triangle)
 1070

Bioclimatic variable	Bio1	Bio2	Bio3	Bio4	Bio5	Bio6	Bio7	Bio8	Bio9	Bio10	Bio11	Bio12	Bio13	Bio14	Bio15	Bio16	Bio17	Bio18	Bio19
Bio1	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio2	0.087	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio3	0.732	0.321	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio4	-0.814	0.011	-0.876	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio5	0.872	0.258	0.455	-0.449	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio6	0.971	-0.038	0.779	-0.911	0.749	1	-	-	-	-	-	-	-	-	-	-	-	-	-
Bio7	-0.773	0.225	-0.784	0.973	-0.377	-0.896	1	-	-	-	-	-	-	-	-	-	-	-	-
Bio8	0.595	-0.066	0.383	-0.398	0.522	0.530	-0.392	1	-	-	-	-	-	-	-	-	-	-	-
Bio9	0.901	0.122	0.663	-0.759	0.803	0.896	-0.715	0.257	1	-	-	-	-	-	-	-	-	-	-
Bio10	0.915	0.129	0.472	-0.513	0.983	0.808	-0.472	0.590	0.815	1	-	-	-	-	-	-	-	-	-
Bio11	0.977	0.050	0.814	-0.918	0.760	0.994	-0.881	0.544	0.894	0.811	1	-	-	-	-	-	-	-	-
Bio12	0.237	-0.513	0.244	-0.397	-0.030	0.327	-0.477	0.293	0.121	0.073	0.303	1	-	-	-	-	-	-	-
Bio13	0.276	-0.341	0.289	-0.403	0.040	0.332	-0.438	0.342	0.142	0.124	0.330	0.926	1	-	-	-	-	-	-
Bio14	-0.084	-0.620	-0.151	-0.011	-0.205	0.013	-0.154	0.039	-0.120	-0.126	-0.048	0.494	0.233	1	-	-	-	-	-
Bio15	0.258	0.528	0.405	-0.263	0.208	0.191	-0.128	0.302	0.156	0.189	0.262	-0.024	0.213	-0.521	1	-	-	-	-
Bio16	0.271	-0.364	0.283	-0.406	0.028	0.333	-0.446	0.330	0.141	0.115	0.328	0.950	0.992	0.258	0.178	1	-	-	-
Bio17	-0.050	-0.639	-0.110	-0.058	-0.194	0.052	-0.203	0.062	-0.089	-0.110	-0.009	0.555	0.294	0.988	-0.508	0.321	1	-	-
Bio18	-0.031	-0.453	-0.011	-0.119	-0.257	0.027	-0.210	0.245	-0.184	-0.144	0.015	0.805	0.741	0.392	0.016	0.766	0.438	1	-
Bio19	0.177	-0.270	0.266	-0.279	0.048	0.263	-0.335	-0.010	0.218	0.080	0.227	0.460	0.353	0.468	-0.196	0.356	0.490	0.115	1

1071

1072 **Supplementary file 7.** Comparison of bioclimatic variables among the four genetic groups
1073 analysed with multivariate analysis of variance (MANOVA)
1074

Predictor	Test statistic	Df	Observed value	F value	Num. Df	Den. Df	P
Genetic group	Pillai	3	1.790	44.215	24	717	< 2e-16
	Wilks	3	0.024	74.690	24	688	< 2e-16
	Hotelling-Lawley	3	12.716	124.870	24	707	< 2e-16
	Roy	3	10.777	321.960	8	239	< 2e-16

1075 Df = degree of freedom among groups; Num. Df = degrees of freedom of the model; Den. Df =
1076 degree of freedom of residual
1077

1078 **Supplementary file 8.** Summary of analysis of variance (ANOVA) for bioclimatic variables
1079

Bioclimatic variable	Df	Sum square	Mean square	F value	P
Bio1 (Annual temperature)	3	183.688	61.229	235.97	<2.2e-16
Bio2 (Mean diurnal temperature range)	3	66.253	22.085	29.813	<2.2e-16
Bio3 (Isothermality)	3	188.031	62.677	259.34	<2.2e-16
Bio8 (Mean temperature of wettest quarter)	3	177.063	59.021	205.91	<2.2e-16
Bio12 (Annual precipitation)	3	157.890	52.630	144.11	<2.2e-16
Bio14 (Precipitation of driest month)	3	29.859	9.952	11.183	<6.6e-07
Bio15 (Precipitation seasonality)	3	119.810	39.938	76.62	<2.2e-16
Bio19 (Precipitation of coldest quarter)	3	44.212	14.737	17.732	< 1.9e-10

1080 Df = degree of freedom

1081

1082 **Supplementary file 9.** Correlation between eight bioclimatic variables and climatic PC axes 1 to
1083 4
1084

Bioclimatic variable	PC1	PC2	PC3	PC4
Bio1 (Annual temperature)	-0.445	-0.107	0.390	-0.056
Bio2 (Mean diurnal temperature range)	0.193	-0.564	0.062	0.253
Bio3 (Isothermality)	-0.430	-0.082	0.492	-0.184
Bio8 (Mean temperature of wettest quarter)	-0.486	0.042	-0.287	0.049
Bio12 (Annual precipitation)	-0.401	0.366	0.066	0.202
Bio14 (Precipitation of driest month)	0.017	0.582	-0.248	0.259
Bio15 (Precipitation seasonality)	-0.323	-0.365	-0.292	0.667
Bio19 (Precipitation of coldest quarter)	0.279	0.235	0.607	0.587

1085 (Correlation coefficients with absolute values higher than 0.4 are in bold.)
1086

1087 **Supplementary file 10.** Comparison of summer growing season data including temperature and
1088 precipitation of May, July and September among the four genetic groups analysed with
1089 multivariate analysis of variance (MANOVA)
1090

Predictor	Test statistic	Df	Observed value	F value	Num. Df	Den. Df	<i>P</i>
Genetic group	Pillai	3	1.915	70.907	18	723	< 2e-16
	Wilks	3	0.010	156.350	18	676	< 2e-16
	Hotelling-Lawley	3	19.873	262.400	18	713	< 2e-16
	Roy	3	15.400	618.570	6	241	< 2e-16

1091 Df = degree of freedom among groups; Num. Df = degrees of freedom of the model; Den. Df =
1092 degree of freedom of residual
1093

1094 **Supplementary file 11.** ANOVA table for all evaluated field traits (phenology, reproduction and size) as well as drought-related traits
 1095

Trait	Garden	Model r^2	Group F	Group P	SEA ¹	SA ¹	CA ¹	Tukey ²
Phenology:								
Days to 50% flowering	Pakistan 2015	0.2388	7.2144	0.0019*	-0.3702	0.6296	-0.3525	B,A,B
Days to 50% flowering	Taiwan 1984	0.5266	25.5887	<.0001*	0.5723	0.3197	-1.0685	A,A,B
Days to 50% flowering	Taiwan 2018	0.4465	18.5544	<.0001*	0.0974	0.6569	-0.9633	A,A,B
Reproduction:								
100 seed weight, g	Pakistan 2015	0.4722	20.5761	<.0001*	0.9050	-0.6397	-0.2756	A,B,B
Pod length, cm	Pakistan 2015	0.2621	8.1698	0.0009*	0.6810	-0.2605	-0.4896	A,B,B
Pod length, cm	Taiwan 1984	0.7173	58.3615	<.0001*	1.0524	-0.1980	-1.0232	A,B,C
Pods per plant	Pakistan 2015	0.4471	18.5988	<.0001*	-0.6441	0.8455	-0.3062	B,A,B
1000 seed weight, g	Taiwan 1984	0.6248	38.3053	<.0001*	0.9979	-0.8380	-0.1340	A,C,B
Seed yield per plant, g	Pakistan 2015	0.4666	20.1225	<.0001*	-0.6210	0.8752	-0.3712	B,A,B
Seeds per pod	Pakistan 2015	0.1300	3.4372	0.0406*	-0.4875	0.2418	0.2806	A,A,A
Seeds per pod	Taiwan 1984	0.1611	4.4168	0.0176*	0.1413	0.3386	-0.6107	AB,A,B
Plant size:								
Petiole length, cm	Pakistan 2015	0.2943	9.5907	0.0003*	0.5435	0.0878	-0.7798	A,A,B
Plant height, cm	Pakistan 2015	0.0001	0.0024	0.9976	0.0075	0.0055	-0.0158	A,A,A
Plant height at flowering, cm	Taiwan 1984	0.3981	15.2115	<.0001*	0.4811	0.3024	-0.9705	A,A,B
Plant height at maturity, cm	Taiwan 1984	0.5472	27.8000	<.0001*	0.3480	0.5605	-1.1362	A,A,B
Primary leaf length, cm	Taiwan 1984	0.5454	27.5930	<.0001*	0.9813	-0.4212	-0.6253	A,B,B
Primary leaf width, cm	Taiwan 1984	0.6053	35.2773	<.0001*	1.0244	-0.6010	-0.4313	A,B,B
Terminal leaflet length, cm	Pakistan 2015	0.2186	6.4340	0.0034*	0.3062	0.2643	-0.7167	A,A,B
Terminal leaflet width, cm	Pakistan 2015	0.1680	4.6458	0.0145*	0.4361	0.0387	-0.5734	A,AB,B
Drought (PEG6000):								

Shoot dry weight (SDW), mg	NTU 2021	0.5998	36.7246	<.0001*	1.0508	-0.5607	-0.5483	A,B,B
Root dry weight (RDW), mg	NTU 2021	0.5964	36.2048	<.0001*	1.0299	-0.6933	-0.3336	A,B,B
Total dry weight (TDW), mg	NTU 2021	0.5934	35.7555	<.0001*	1.0448	-0.5883	-0.5028	A,B,B
Root:Shoot ratio dry weight (RSRDW)	NTU 2021	0.3261	11.8577	<.0001*	-0.2316	-0.4342	0.9112	B,B,A
Drought (Control):								
Shoot dry weight (SDW), mg	NTU 2021	0.5779	33.5453	<.0001*	1.0275	-0.4709	-0.6484	A,B,B
Root dry weight (RDW), mg	NTU 2021	0.5205	26.5962	<.0001*	0.9716	-0.6077	-0.3811	A,B,B
Total dry weight (TDW), mg	NTU 2021	0.5638	31.6722	<.0001*	1.0195	-0.5374	-0.5430	A,B,B
Root:Shoot ratio dry weight (RSRDW)	NTU 2021	0.1753	5.2083	0.0089*	-0.3036	-0.2004	0.6773	B,B,A

1096 *Significant at $P < 0.05$; 1: least-square means of each group after inverse normal transformation of raw data; 2: levels not connected
1097 by same letter are significantly different.
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1099 **Supplementary file 12.** Mean of eight bioclimatic variables of the genetic groups

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Bioclimatic variable	Northeast Asia (N = 37) Mean ± SD	Northwest Asia (N = 45) Mean ± SD	Southeast Asia (N = 45) Mean ± SD	South Asia (N = 49) Mean ± SD	Central Asia (N = 72) Mean ± SD
Bio1	62.49 ± 55.39	117.58 ± 54.30	256.47 ± 19.29	255.57 ± 14.80	128.72 ± 41.10
Bio2	115.22 ± 15.44	124.38 ± 20.14	102.02 ± 16.11	127.24 ± 16.12	130.08 ± 13.89
Bio3	25.14 ± 2.04	31.02 ± 4.47	51.58 ± 7.19	42.8 ± 4.58	32.4 ± 3.52
Bio8	207.86 ± 31.97	107.89 ± 67.90	271.47 ± 11.68	283.92 ± 23.02	92.04 ± 42.23
Bio12	821.59 ± 299.98	301.24 ± 201.00	1477.69 ± 380.18	750.39 ± 329.19	285.67 ± 145.24
Bio14	12.14 ± 11.72	5.18 ± 8.99	6.33 ± 4.34	2.47 ± 3.44	2.51 ± 4.98
Bio15	95.35 ± 24.52	65.4 ± 20.35	84.78 ± 7.25	124.47 ± 23.48	70.92 ± 12.92
Bio19	42.51 ± 38.23	88.33 ± 67.16	48.02 ± 24.91	34 ± 33.99	92.15 ± 45.19

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