

# A chickpea genetic variation map based on the sequencing of 3,366 genomes

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Zero hunger and good health could be realized by 2030 through effective conservation, characterization and utilization of germplasm resources<sup>1</sup>. So far, few chickpea (*Cicer arietinum*) germplasm accessions have been characterized at the genome sequence level<sup>2</sup>. Here we present a detailed map of variation in 3,171 cultivated and 195 wild accessions to provide publicly available resources for chickpea genomics research and breeding. We constructed a chickpea pan-genome to describe genomic diversity across cultivated chickpea and its wild progenitor accessions. A divergence tree using genes present in around 80% of individuals in one species allowed us to estimate the divergence of *Cicer* over the last 21 million years. Our analysis found chromosomal segments and genes that show signatures of selection during domestication, migration and improvement. The chromosomal locations of deleterious mutations responsible for limited genetic diversity and decreased fitness were identified in elite germplasm. We identified superior haplotypes for improvement-related traits in landraces that can be introgressed into elite breeding lines through haplotype-based breeding, and found targets for purging deleterious alleles through genomics-assisted breeding and/or gene editing. Finally, we propose three crop breeding strategies based on genomic prediction to enhance crop productivity for 16 traits while avoiding the erosion of genetic diversity through optimal contribution selection (OCS)-based pre-breeding. The predicted performance for 100-seed weight, an important yield-related trait, increased by up to 23% and 12% with OCS- and haplotype-based genomic approaches, respectively.

Pulses are an important crop commodity providing protein for human health. Worldwide pulse productivity has been stagnant for the last five decades, contributing to low per-capita availability of these foods and high levels of malnutrition in developing countries<sup>3</sup>. Chickpea (*Cicer arietinum* L.) production ranks third among pulses, and chickpea is cultivated in more than 50 countries, especially in South Asia and sub-Saharan Africa. As it is an important source of protein, dietary fibre and micronutrients, chickpea is key to nutritional security. More than 80,000 chickpea germplasm accessions are being conserved in 30 genebanks across the world<sup>4</sup>, but only a few have been used for chickpea improvement<sup>2</sup>.

Germplasm sequencing efforts in some crop plants have provided insights into the global distribution of genetic variation<sup>5</sup>; how this diversity has been shaped by the genetic bottlenecks associated with domestication<sup>6</sup> and by the effects of selective breeding<sup>7</sup>; and, finally, how we can link this genetic variation to phenotypic diversity<sup>2</sup> for breeding applications. Haplotype maps developed using whole-genome sequencing (WGS) data have helped to determine the percentage of the constrained genome and detect deleterious mutations that can be purged for accelerated breeding<sup>8,9</sup>. Furthermore, sequencing and genotyping of a germplasm collection allows better conservation and management in genebanks<sup>5,10</sup>.

A list of affiliations appears at the end of the paper.

On the basis of WGS of 3,366 chickpea germplasm accessions, we report here a rich map of the genetic variation in chickpea. We provide a chickpea pan-genome and offer insights into species divergence, the migration of the cultigen (*C. arietinum*), rare allele burden and fitness loss in chickpea. We propose three genomic breeding approaches—haplotype-based breeding, genomic prediction and OCS—for developing tailor-made high-yielding and climate-resilient chickpea varieties.

We sequenced 3,366 chickpea germplasm lines, including 3,171 cultivated and 195 wild accessions at an average coverage of around 12× (Methods, Extended Data Fig. 1, Supplementary Data 1 Tables 1, 2). Alignment of WGS data to the CDC Frontier reference genome<sup>11</sup> identified 3.94 million and 19.57 million single-nucleotide polymorphisms (SNPs) in 3,171 cultivated and 195 wild accessions, respectively (Extended Data Table 1, Supplementary Data 1 Tables 3–7, Supplementary Notes). This SNP dataset was used to assess linkage disequilibrium (LD) decay (Supplementary Data 2 Tables 1, 2, Extended Data Fig. 2, Supplementary Notes) and identify private and population-enriched SNPs (Supplementary Data 3 Tables 1–4, Supplementary Notes). These private and population-enriched SNPs suggest rapid adaptation and can enhance the genetic foundation in the elite gene pool.

## Pan-genome

We developed a chickpea pan-genome (592.58 Mb) using an iterative mapping and assembly approach by combining the CDC Frontier reference genome, an additional 2.93 Mb from a desi genome (ICC 4958)<sup>12</sup>, 3.70 Mb from a *Cicer reticulatum* genome<sup>13</sup> and 53.66 Mb from de-novo-assembled sequences from cultivated (48.38 Mb; 3,171) and *C. reticulatum* (5.28 Mb; 28) accessions (Supplementary Data 4 Table 1). Although similar pan-genome studies have been conducted in other crops, including rice<sup>5,14</sup>, soybean<sup>15</sup> and *Brassica oleracea*<sup>16</sup>, our pan-genome comprises more than 3,000 individuals.

A total of 29,870 genes (1,601 additional gene models) were identified, of which 1,582 were to our knowledge novel compared to previously reported genes<sup>11</sup>. Gene ontology (GO) annotations identified genes that encode response to oxidative stress, response to stimulus, heat shock protein, cellular response to acidic pH and response to cold (Supplementary Data 4 Tables 2, 3), suggesting a possible role in adaptation. The modelling analysis curve eventually reaching saturation suggested that the pan-genome is closed, in concurrence with other plant pan-genomes<sup>14,16</sup> (Fig. 1a). N50, a widely used metric to assess the quality of an assembly, is the length of the shortest contig for which larger and equal size contigs cover 50% of the total assembly. The N50 values for sequences from de-novo-assembled cultivated and *C. reticulatum* accessions, *C. reticulatum* and the desi genome were 2.61 kb, 1.30 kb, 1.78 kb and 1.76 kb, respectively, whereas the average gene length was 4.72 kb, 1.09 kb, 1.09 kb and 0.98 kb (Supplementary Data 4 Table 1). This pan-genome was further used to assess the effect of presence–absence variations on protein-coding genes (Supplementary Data 4 Table 4, Supplementary Notes).

Cultivated (2,258) and *C. reticulatum* (22) accessions with a coverage of greater than 10× were analysed to discover structural variations, including insertions (139,483), deletions (47,882), inversions (61,171), intra-chromosomal translocations (417) and inter-chromosomal translocations (2,410) in cultivated and 287,854 insertions, 67,351 deletions, 58,070 inversions, 446 intra-chromosomal translocations and 2,066 inter-chromosomal translocations among *C. reticulatum* accessions as compared to the CDC Frontier genome<sup>11</sup> (Fig. 1b, Extended Data Table 1, Supplementary Data 5 Table 1, Supplementary Notes). More structural variations in the *C. reticulatum* accessions were expected because of their high divergence from cultivated chickpea. We further identified 793 gene-gain copy number variants (CNVs) and 209 gene-loss CNVs in cultivated accessions, and 643 gene-gain and 247 gene-loss CNVs in *C. reticulatum* accessions (Supplementary Data 5 Tables 2, 3).

## Species divergence and migration

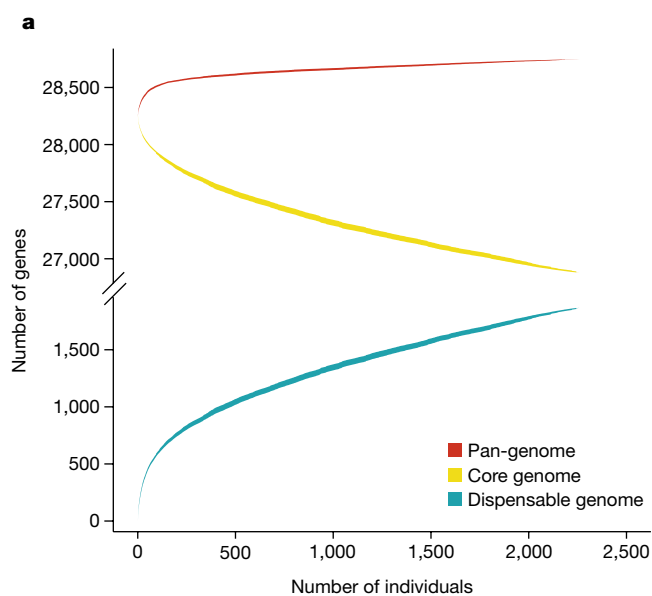
To understand speciation and estimate species divergence time in the eight *Cicer* species analysed here, single-copy genes identified using ‘fabales’ genes from the BUSCO<sup>17</sup> database were used to carry out homologue-based gene annotation in preliminary genome assemblies, the CDC Frontier<sup>11</sup> and *Medicago truncatula*<sup>18</sup>. Using these single-copy genes, *Cicer cuneatum* was estimated to have diverged from other *Cicer* species around 21.4 (19.6–22.8) million years ago (Ma) (Extended Data Fig. 3a, Supplementary Notes), about the time that Arabia collided with Asia, and a time when ‘Rand Flora’ taxa like *Cicer* may have migrated from Africa into Southwest Asian habitats<sup>19</sup>. *C. reticulatum* and *Cicer echinospermum* were estimated to have diverged around 15.3 (14.0 to 16.2) Ma, which is higher than previous estimates and might be influenced by: (i) wild accessions conserved at the International Crops Research Institute for the Semi-Arid Tropics (ICRISAT) representing only some populations of these species, when recent work has shown that only some *C. echinospermum* populations are cross-compatible with *C. arietinum*; and (ii) introgression from *C. echinospermum* into cultivated chickpea, which is widespread in Australian and North American breeding lines, and is also likely to have occurred in International Center for Agricultural Research in the Dry Areas (ICARDA) lines.

Phylogenetic analysis grouped all 195 wild accessions into 6 clusters (Clusters I–VI) (Extended Data Fig. 3b, Supplementary Notes). Cluster IVa included all *C. reticulatum* and one *C. echinospermum* (ICC 20192; green colour), whereas cluster IVb included all *C. echinospermum* and one *C. reticulatum* (ICC 73071; golden-yellow colour). Similarly, one *Cicer pinnatifidum* (ICC 20168; red colour) was grouped with the *Cicer bijugum* accessions in cluster II, and one *C. bijugum* (ICC 20167; blue colour) was grouped with *C. pinnatifidum* accessions in cluster I. These are two cross-compatible species. Spontaneous hybridization might have occurred in nature. In terms of post-species divergence, a homologue (*Ca\_25684*) of *SHATTERPROOF2* (also known as Agamous-like MADS-box protein (*AGL5*)), which is responsible for seed dispersal, was analysed for haplotypic variation (Supplementary Notes). We found an association of the ‘C’ allele with low or minimal shattering in cultivated species, as seen at the low shattering allele (‘C’) on chromosome 5 at position 1,022,962 of the orthologue in common bean<sup>20</sup>.

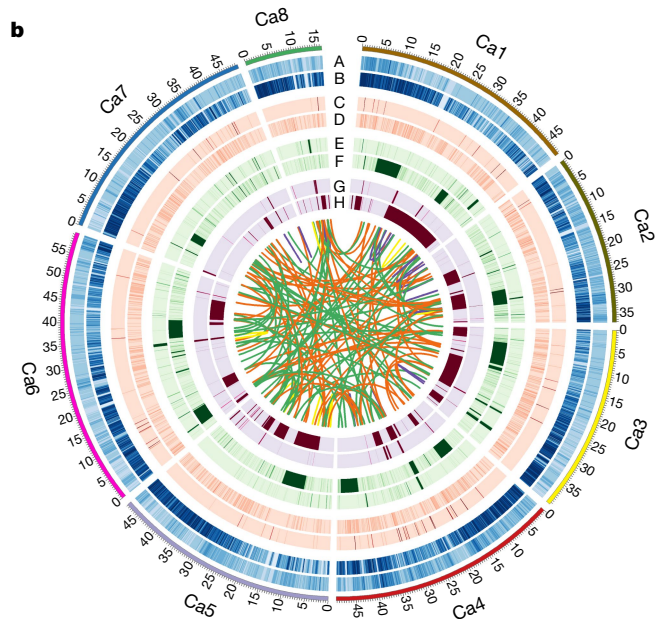
The neighbour-joining tree grouped most South Asian accessions with no distinct clustering for other geographic origins (Extended Data Fig. 4). Our principal component analysis (PCA) of accessions suggests two paths of diffusion or migration of chickpea from the centre of origin in the Fertile Crescent: one path indicates diffusion to South Asia and East Africa, and the other suggests diffusion to the Mediterranean region (probably through Turkey) as well as to the Black Sea and Central Asia (up to Afghanistan) (Fig. 2a–f, Extended Data Fig. 5). This diffusion translated into a pattern of nucleotide diversity ( $\pi$ ), among accessions from Central Asia ( $4.74 \times 10^{-4}$ ) and South Asia ( $3.62 \times 10^{-4}$ ) (Supplementary Data 6 Table 1), which is consistent with earlier reports<sup>2</sup>. Pairwise fixation index ( $F_{ST}$ ) estimations further supported these findings (Supplementary Data 6 Table 2, Supplementary Notes).

## Domestication and breeding bottlenecks

Our analysis indicates that chickpea experienced a strong bottleneck beginning around 10,000 years ago. The population size reaching its minimum around 1,000 years ago, followed by a very strong expansion of the population within the last 400 years (Extended Data Fig. 6), suggest a strong recent expansion of chickpea agriculture. One consequence of this bottleneck is shown by the higher  $\pi$  in *C. reticulatum* ( $2.20 \times 10^{-3}$ ) relative to cultivated accessions (Extended Data Table 1, Supplementary Data 6 Table 1).



**Fig. 1 | Global chickpea genetic variations. a**, The chickpea pan-genome. Modelling analysis of the pan-genome and core genome shows an increase and decrease in the number of genes with each added genotype, indicating that the pan-genome is a closed pan-genome. The thickness of the curves represents the 99% confidence interval. **b**, Circos diagram illustrating the variation density among chickpea lines. Overall, higher numbers of variations were observed among wild accessions. Tracks indicate SNP density among cultivated (A) and



wild (B), insertion density among cultivated (C) and *C. reticulatum* (D), deletion density among cultivated (E) and *C. reticulatum* (F), and inversion density among cultivated (G) and *C. reticulatum* (H). Links represent inter- and intra-chromosomal translocations. Yellow (cultivated) and purple (*C. reticulatum*) denote intra-chromosomal translocations, whereas orange (cultivated) and green (*C. reticulatum*) represent inter-chromosomal translocations.

Genetic relationship analysis between cultivated and wild chickpea showed that one cultivated accession (ICC 16369) from East Africa was grouped with wild chickpea (Extended Data Fig. 7). This same genotype also showed the presence of the 'T' allele, specific to wild species in *SHATTERPROOF2*, suggesting that ICC 16369 has been mislabelled as belonging to the cultivated chickpea (Supplementary Data 7).

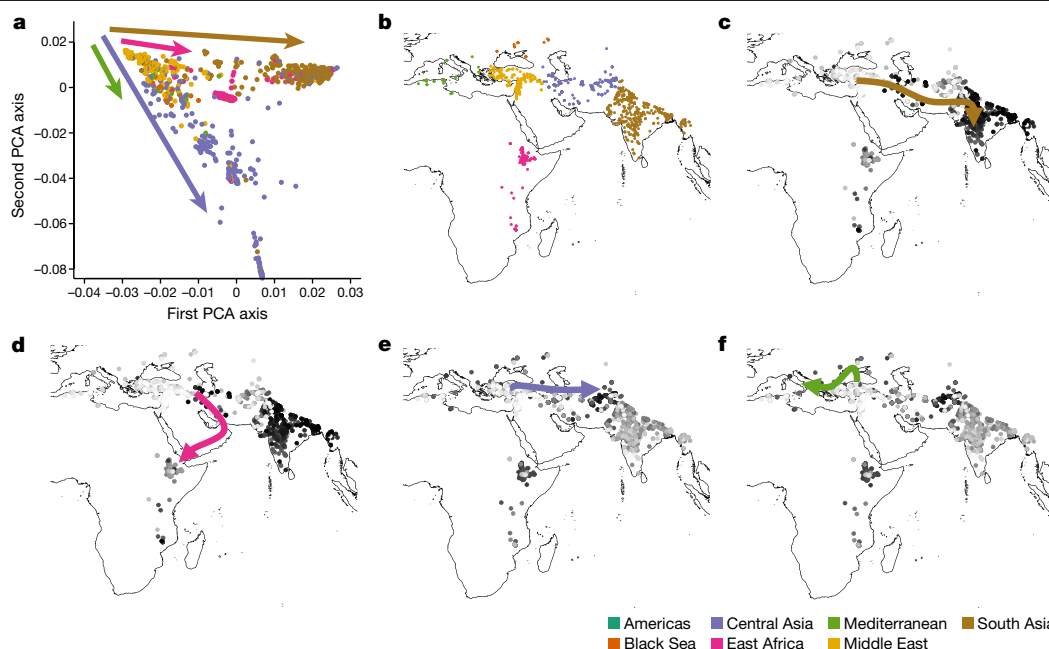
To detect selection sweeps, we pinpointed 18 fragments in cultivated chickpea using the composite likelihood ratio (CLR) (Extended Data Fig. 6, Supplementary Data 6 Tables 8, 9). Combined analysis with reduction of diversity (ROD),  $F_{ST}$  and Tajima's  $D$  identified genomic regions for *C. reticulatum* (immediate wild species progenitor) versus landraces (2,899; 42,148 kb), landraces versus breeding lines (191; 4,360 kb) and breeding lines versus cultivars (14; 404 kb) that might have undergone selection during domestication and breeding (Supplementary Data 6 Tables 3–6, Supplementary Notes, <https://doi.org/10.6084/m9.figshare.15015327>). We identified 35 regions (222 kb) common between *C. reticulatum* versus landraces and landraces versus breeding lines, and similarly one region (4 kb) between landraces versus breeding lines and breeding lines versus cultivars. Furthermore, we identified a total of 37 unique potential genes in these 36 regions that may have a role in the adaptation of chickpea during migration to different environments by regulating flowering time and plant growth (Supplementary Data 6 Table 7). For example, *FLP2* (flower development and vegetative to reproductive phase transition of meristem), *LRP1* (root growth), *PIPSKL1* (signalling pathways for survival and T cell metabolism) and *MYB12* (flavonoid biosynthesis) are some key genes we pinpointed that are critical for plant growth, metabolic pathways and adaptation in changing environments.

We used genomic evolutionary rate profiling (GERP) analysis to identify 29 Mb (8.36%) genomic regions as evolutionarily constrained (GERP score of greater than 0), indicating purifying selection (Extended Data Fig. 8a). Using constrained genome, sorting intolerant from tolerant<sup>21</sup> (SIFT) score (less than 0.05) and GERP (greater than 2), 10,616

non-synonymous SNPs were identified as candidate deleterious mutations (Extended Data Fig. 8b). Using the derived allele frequency (DAF) spectrum, we selected 37 non-synonymous deleterious mutations (SIFT < 0.05; GERP > 2; DAF > 0.8) in 36 genes (Supplementary Data 8 Tables 1–4), as fixed that have not been purged through traditional breeding. Detailed analysis indicated a higher (17.88%,  $P = 0.01772$ ) abundance of deleterious alleles in the wild progenitor (*C. reticulatum*) than in cultivated accessions (Extended Data Fig. 8c). Furthermore, the mutation burden for genomic regions under selection suggested that the number of deleterious mutations in landraces was approximately twofold that in breeding lines (206.91%;  $P = 2.195676 \times 10^{-60}$ ) (Extended Data Fig. 8d). To increase the fitness of cultivated chickpea, these deleterious alleles are potential targets for genomics-assisted breeding and genome editing.

### Superior haplotypes for key traits

We used 3.94 million SNPs and phenotyping data for 16 traits on 2,980 cultivated genotypes to identify 205 SNPs associated with 11 traits (Methods, Supplementary Data 9 Table 1, Supplementary Notes). Of the 205 associated SNPs, 152 were present in 79 unique genes with potential roles in controlling seed size and development. Analysis of these genes across cultivated genotypes identified 350 haplotypes (Supplementary Data 9 Tables 2–4, Supplementary Notes). Using 19.10 million haplo-pheno combinations, we identified 24 consistent and stable superior haplotypes for 20 genes (Supplementary Data 9 Tables 5–7, Extended Data Fig. 9a). This analysis revealed that the majority of breeding lines (80%) lacked superior haplotypes that are present in the landraces. We validated superior haplotypes by using historical data on 129 chickpea varieties released between 1948 and 2012 (Extended Data Fig. 9b, c). Finally, we identified 56 lines as potential donors for introducing superior haplotypes in breeding (Supplementary Data 9 Tables 8–10).



**Fig. 2 | Insights into chickpea migration.** **a–f**, The PCA based on geographic origin suggests two paths of diffusion (**a**, **b**). The first path illustrates a diffusion to South Asia (**c**) and East Africa in parallel (**d**). The second path suggests a diffusion to Central Asia (**e**) together with the Mediterranean region (**f**).

## Enriching the genetic base

We combined OCS<sup>22</sup> with a mate allocation method that takes into account genetic gain and genetic diversity as a guide for potential future chickpea pre-breeding programmes or ‘evolving gene banks’<sup>22,23</sup> (Supplementary Notes). With a price bonus for earliness and for large seeds, we chose 274 (9.4%) unique genotypes for 325 matings from the 2,898 available genotypes, divided among desi (190), kabuli (120) and intermediate (15), using MateSel<sup>24</sup> (Supplementary Data 10 Table 1).

The frequency distribution of predicted progeny index (mean of nine environments) values was bimodal. Higher predicted progeny index values were observed in kabuli as compared with desi. However, marked improvements were predicted in desi and kabuli, from candidate parents to predicted progenies (Extended Data Fig. 10a, b). The frequency distribution of predicted progeny genomic estimated breeding value (GEBV) for yield per plant (YPP) in desi (13.79 g) exceeded kabuli (12.65 g) and a higher response to selection was observed for desi (0.6 g; 4.3%) than for kabuli (0.4 g; 3.5%) (Extended Data Fig. 10c, d). For 100-seed weight (100SW), the mean 100SW of predicted progeny in kabuli (30.6 g) was almost twice that of desi (16.9 g), and the response to selection was three times higher for kabuli (5.7 g; 23%) than for desi (2.0 g; 13%) (Fig. 3a, Extended Data Fig. 10e, f). Kabuli progeny, with a later flowering time, did not respond to selection for earliness (–1.0 day) as rapidly as desi progeny (–3.3 days) (Extended Data Fig. 10g, h). These predicted responses to selection in the next cycle occurred with a relatively small increase in predicted progeny inbreeding in the desi (0.03) and intermediate (0.02), but a large increase in the kabuli (0.17) (Extended Data Fig. 10i, Supplementary Data 10 Table 2, Supplementary Notes).

## Breeding population improvement

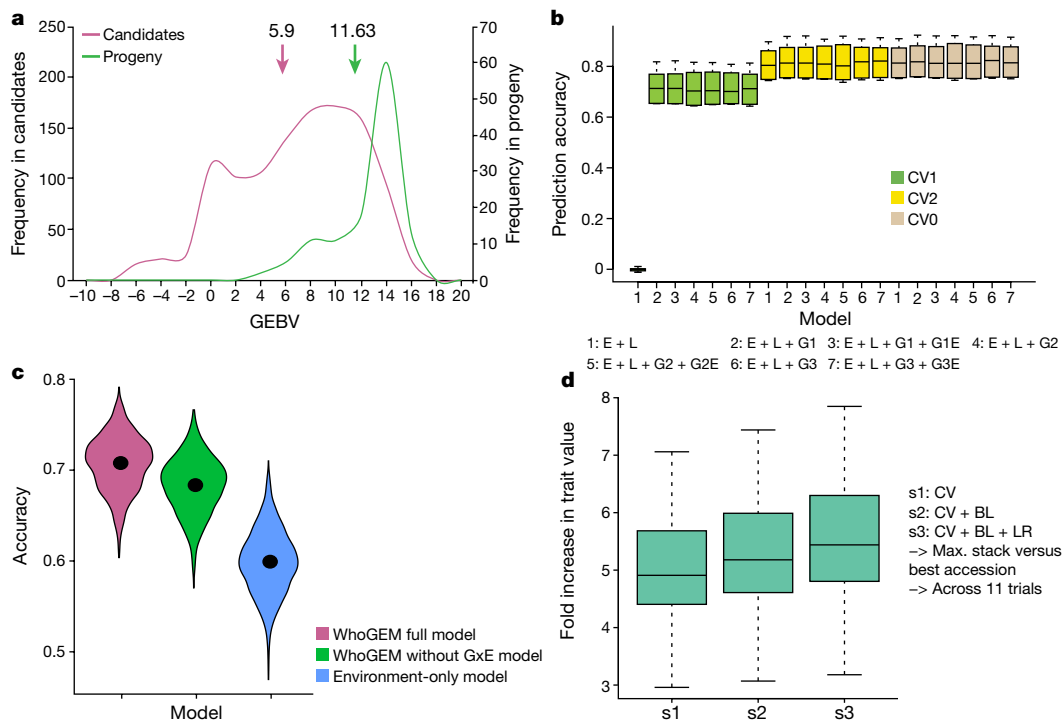
We used different subsets of SNPs and phenotyping data on 16 traits across 12 combinations of year and location, following 3 genomic prediction approaches: (i) interaction of marker and environment covariates ( $G \times E$ )<sup>25</sup>; (ii) implementation of the WhoGEM approach<sup>26</sup>; and (iii) a haplotype-based approach for estimating local GEBVs<sup>27</sup>.

In the first approach, 3 genomic relationship matrices with 223,119 (G1), 531,457 (G2) and 754,576 (G3) SNPs, and phenotyping data for 9 traits on 2,980 genotypes, were used to understand the variability explained within the groups and environments (Supplementary Data 10 Table 3). Overall, the environment (E) + genotype (L) + marker effects (G3) model for cross-validation scheme 0 (CV0; see ‘Prediction using the interaction of genomic and environmental covariates’ in Methods) produced the highest average correlation (0.719) for 100SW, and the E + L model returned the lowest value (0.031) for basal secondary branch (Supplementary Data 10 Table 4). For 100SW, genomic prediction accuracy varied from 0.611 (E + L + G3 + G3E) to 0.719 (E + L + G3) for CV1 and CV0, respectively (Fig. 3b).

In the second approach, we used WhoGEM with 276,956 LD-pruned SNPs and phenotyping data for 9 traits on 1,318 genotypes (with GPS data). Prediction accuracies of the full model ranged from 0.25 to 0.91 (Supplementary Data 10 Table 5). Although the highest prediction accuracy was obtained for plot yield (0.914), this method was still efficient in predicting 100SW, with an accuracy of 0.599 (environment-only model) to 0.707 (WhoGEM full model) (Fig. 3c). Evidence for interactions between admixture components and the environment was presented for phenology, plant production and plant architecture traits (Extended Data Fig. 11a–m). The use of admixture components integrates the effects of demography (that is, gene flow and genetic drift) and artificial or natural selection to explain phenotypic variation with reasonable accuracy. This shows considerable potential to detect the accumulation of favourable admixture components from the wider genepool.

In the third approach, 124,833 selected SNPs were used to construct LD blocks, called haplotypes. These SNPs and phenotyping data for 100SW and YPP for 2,980 genotypes were used to estimate local GEBVs for the haplotypes. The local GEBV analysis revealed substantial genetic potential in each subgroup for trait improvement (Extended Data Fig. 12). When comparing the best accessions with the highest GEBVs to the in silico genotypes that combined all haplotypes with the highest trait effect across the whole genome, the predicted performance increased by more than threefold for YPP and by more than fivefold for 100SW (Fig. 3d). Our results indicate that capturing novel alleles from landraces through a haplotype-based prediction approach could improve YPP or 100SW by 6–12% (Fig. 3d).





**Fig. 3 | An example of the use of four genomic breeding strategies for improving 100SW. a**, Mean GEBV and total genetic values predict a 23% increase in one generation for 100SW in kabuli candidates. **b**, Genomic-enabled predictions using Bayesian generalized linear regression (BGLR) on three cross-validation schemes provided the highest mean prediction accuracy with scheme CV0 ( $n = 2,980$  cultivated accessions). **c**, A general linear model using the WhoGEM prediction machine provided the highest prediction accuracies for the WhoGEM full model ( $n = 1,500$ ; 300 replicates of a fivefold

cross-validation). In each violin plot, the black dot represents the mean. GxE, genotype and environment interaction. **d**, Haplotype-based local GEBVs that are suggested to provide a fivefold improvement in performance over the best different groups (cultivars (CV,  $n = 152$ ), breeding lines (BL,  $n = 396$ ) and landraces (LR,  $n = 2,439$ )). Each of the box plots shows the upper and lower whisker (indicated by dashed lines), the 25% and 75% quartiles and the median (as a solid line).

## Discussion

Our study reports global polymorphisms in chickpea by sequencing 3,366 germplasm accessions (3,171 cultivated and 195 wild). This analysis brings greater resolution to our understanding of the within-species diversity of *C. arietinum*. The chickpea pan-genome (592.58 Mb) developed from cultivated draft genomes<sup>11,12</sup> and the *C. reticulatum* genome<sup>13</sup>, together with WGS data on cultivated and *C. reticulatum* accessions, provided insights into gene content variation across cultivated chickpea and its wild progenitor.

Although some studies based on chloroplast DNA<sup>28</sup> and nuclear ribosomal DNA<sup>29</sup> have been conducted to investigate the evolution and domestication of *Cicer* species in the past, their resolution was limited. Here, by using WGS data for a large number of individuals, we estimated the divergence time between chickpea and its closest progenitor species. Our study also provides opportunities to rectify misclassifications of accessions to the correct species and to determine whether chickpea seeds preserved in archaeological sites were wild or cultivated.

We identified selective sweeps and candidate genes under domestication and breeding that were responsible for reducing genetic diversity in the cultivated gene pool. Most importantly, our study analysed genetic loads in *Cicer* species. Although selection and recombination have successfully purged many deleterious alleles, the current collection of breeding lines and cultivars still contains substantial genetic loads that affect crop fitness. Here, we have identified deleterious alleles for purging through genome-informed breeding and/or gene editing.

We identified numerous superior haplotypes for improvement-related traits in landraces, and used the concept of superior haplotypes by comparing the yield of the released varieties carrying superior versus

regular haplotypes for yield-related traits<sup>30</sup>. Furthermore, we estimated prediction accuracies for agronomic traits using three genomic prediction approaches and provided a case study for 100SW, demonstrating that genomic prediction approaches have great potential for enhancing crop productivity. We suggest using haplotype mining and genomic prediction approaches in chickpea and other crops to provide climate resilience and improved nutrition to meet future worldwide demand.

## Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-021-04066-1>.

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## Methods

### Germplasm sequencing and variant calling

We performed WGS of 2,967 *Cicer* accessions from a global composite collection<sup>4</sup> using the HiSeq2500 at the Center of Excellence in Genomics and Systems Biology, ICRISAT. By including sequence data of 399 lines from an earlier study<sup>2</sup>, we analysed 3,366 accessions (3,171 cultivated and 195 wild species accessions) altogether (Supplementary Notes).

We aligned sequencing data from the 3,366 chickpea accessions to the reference genome of CDC Frontier<sup>11</sup>, using BWA-MEM<sup>31</sup> v.0.7.15. SNP calling was performed using GATK<sup>32</sup> v.3.7 as per GATK best practices for SNP calling, thus creating the base SNP set. We defined two other SNP sets: (i) Set-A: only SNPs with <30% missing call, and biallelic calls, and (ii) Set-B: SNPs with less than 30% missing calls, biallelic calls, and LD-pruned using PLINK<sup>33</sup> v.1.90 (“--indep-pairphase 50 10 0.2” parameter). Set-B SNPs were only used to depict the population genetic structure.

### Private and population-enriched SNPs

To determine the private and population-specific SNPs, the frequency of alleles within a given population was determined using VariantsToTable<sup>34</sup> of GATK v3.8.1. We defined ‘private alleles’ as those present in at least four accessions within a population and absent in other populations, and ‘population-enriched alleles’ as those present in a given population ( $\geq 20\%$ ) and less frequent in other populations<sup>5</sup> ( $\leq 2\%$ ).

### LD decay, diversity and $F_{ST}$

LD decay was determined using the software PopLDdecay<sup>35</sup> v.3.29 with the parameter “-MaxDist 1000”. Nucleotide diversity ( $\pi$ ) was calculated from a 100-kb sliding window with a 10-kb step using VCFtools<sup>36</sup> v.0.1.13. The average of all valid windows was considered the population genetic diversity. The fixation index ( $F_{ST}$ ) was calculated from 100-kb non-overlapping windows using VCFtools. The global weighted  $F_{ST}$  was used to measure the differentiation of populations.

### Construction of a pan-genome

The chickpea draft genome of CDC Frontier<sup>11</sup> (a kabuli variety; considered as the foundation genome) together with ICC 4958<sup>12,37</sup> (a desi genome sequence), a *C. reticulatum* genome<sup>13</sup>, and de-novo-assembled sequences from 3,171 cultivated and 28 *C. reticulatum* accessions were used to guide the assembly of the chickpea pan-genome using a conservative approach<sup>38</sup>. Following the alignment of reads from each accession to the reference, unmapped and dangling mapped read pairs were extracted using SAMTools<sup>39</sup> v.1.2 based on the FLAG field. The extracted reads were de-novo-assembled using MEGAHIT<sup>40</sup> v.1.2.9 with default parameters. To identify possible redundancies among assembled contigs that were already present in the foundation genome, the assembled contigs were aligned to the foundation genome using NUCmer<sup>41</sup> v.4.0.0beta2 with the parameters “-l 20 -c 65” and the alignments with length  $\geq 500$  bp and identity of greater than 80% were extracted to be added into the intermediate pan-genome. The processes were performed one by one: ICC 4958, de-novo-assembled sequences from 3,171 cultivated accessions, the *C. reticulatum* genome, and de-novo-assembled sequences from 28 *C. reticulatum* accessions. Further, to identify redundancy among the ‘novel’ sequences, all-versus-all alignment was performed using CD-HIT<sup>42</sup> v.4.81. The same process was performed for the next iteration until no sequence was left. Finally, we removed the potential containments from vectors, bacteria, viruses, animals, fungi and organelle sequences using BLASTN<sup>43</sup> v.2.2.31 to the corresponding NT databases and obtained the final pan-genome. As a result, the CDC Frontier genome<sup>11</sup> and novel assembled sequences were combined to construct the chickpea pan-genome.

### Structural and copy number variations

A total of 2,258 cultivated and 22 *C. reticulatum* accessions (with sequence depth of greater than  $10\times$ ) were used to identify structural variations

against the reference genome of CDC Frontier<sup>11</sup>, such as large insertions, deletions, inversions, and intra- and inter-chromosomal translocations. The insertions, deletions and inversions were identified using a dual calling strategy through BreakDancer<sup>44</sup> v.1.1.2 and Pindel<sup>45</sup> v.0.2.5b9. First, BreakDancer was used to detect structural variations with parameter “-q 20 -y 20 -r 1”. Secondly, the output of BreakDancer was used as an input for Pindel using the parameter “-x 4 -breakdancer” to increase the sensitivity and specificity. To merge the results from BreakDancer and Pindel, two structural variants with a distance between the two break-points of less than 100 bp were considered the same structural variation and merged. Owing to the inability of Pindel to detect intra- and inter-chromosomal translocations, only BreakDancer was used for their analysis. Furthermore, a structural variation was considered if it was present in at least 5% of the individuals in a given population.

For CNVs, we first generated a GC-content profile using gccount (<http://bioinfo-out.curie.fr/projects/freec/src/gccount.tar.gz>) with parameter “window = 1000 step = 1000” to normalize non-uniform read coverage of genomic position. Then, Control-FREEC<sup>46</sup> v.11.0 was used to detect CNVs in 1-kb non-overlapping windows (bins) with parameter “ploidy = 2 window = 1000 step = 1000 mateOrientation=FR” for each high-depth individual (sequencing depth > 10X). Next, the sample-level copy numbers were combined to produce a matrix of copy numbers for each bin at the cohort level. To further reduce false positives, we filtered out the bins with a CNV rate of less than 1%. The affected genes were identified by the presence of overlapping regions with CNVs.

### Divergence and phylogenetic relationship

For divergence time estimation, 195 wild species accessions were assembled individually using MEGAHIT<sup>40</sup> v.1.2.9 with default parameters. Then, the ‘fabales’ genes were downloaded from the BUSCO<sup>17</sup> database (odb10), which contains 5,366 single-copy orthologues to predict the genes for 195 wild species accessions, CDC Frontier genome<sup>11</sup> and *M. truncatula* genome<sup>18</sup> (as outgroup) using GeneWise<sup>47</sup> v.2.4.1 with the parameters “-both -sum -genesf”. On the basis of the gene annotations of 195 wild species accessions, only one sample with the longest average coding sequence (CDS) length was chosen for each wild species. The CDS sequences of single-copy genes in seven wild species, CDC Frontier and *M. truncatula* were extracted. For each single-copy family, multiple sequence alignment was performed using MUSCLE<sup>48</sup> v.3.8.31 with default parameters and poorly aligned and divergent regions were eliminated using Gblocks<sup>49</sup> v.0.91b with the parameter “-t=c”. The aligned matrix from each single-copy family was combined to construct the super aligned matrix. The maximum likelihood tree was constructed using RAxML<sup>50</sup> v.8.2.12 with parameters “-f a -x 12345 -p 12345 -m GTR-CATX”. Finally, divergence time was estimated by MCMCTree<sup>51</sup> v.4.4 with three time-calibration points (0.007–0.013 Ma for *C. reticulatum*–*C. arietinum*, 12.2–17.4 Ma for *C. arietinum*–*C. pinnatifidum*, and 30.0–54.0 Ma for *C. arietinum*–*M. truncatula*) from the literature<sup>52–54</sup>.

To assess the relatedness among 195 wild accessions and 3,171 cultivated lines, the genetic distance matrix based on identity by state (IBS) was calculated through PLINK v1.90 with the parameter “--distance 1-ibs” using LD-pruned SNPs (--indep-pairwise 50 10 0.2) present on pseudomolecules. On the basis of the distance matrix, neighbour-joining phylogenetic trees were then constructed using ‘neighbor’ in PHYLIP<sup>55</sup> v.3.6.

APCA was undertaken to study the relatedness and clustering among cultivated chickpea accessions. The top 20 principal components (PCs) of the variance-standardized relationship matrix were estimated using EIGENSOFT<sup>56</sup> v.7.2.0 with default parameters on LD-pruned SNPs present on pseudomolecules. PCA results were plotted using the R package ‘rworldmap’ (ref.<sup>57</sup>).

### Diversity and genetic bottleneck

To characterize variation among populations, population differentiation statistics ( $F_{ST}$ ) were calculated in a 10-kb/2-kb sliding window using VCFtools v.0.1.13. A range of pairwise  $F_{ST}$  was calculated in the same

combinations as for the ROD calculations. Tajima's  $D$  was calculated using VCFtools ("--TajimaD 100000") in 100-kb non-overlapping windows. A window was considered a selection window in the upper 90% of the population's empirical distribution for ROD and  $F_{ST}$  statistics, along with a negative Tajima's  $D$  value (less than  $-2$ ). Genes located on the selection windows were identified, and functional enrichment of the KEGG pathway (v.87.0) and GO term for these candidate genes was conducted using the Fisher's exact test with false discovery rate correction using EnrichmentPipeline<sup>58</sup> (<https://sourceforge.net/projects/enrichmentpipeline/>).

For determining population size histories and split times, the SMC++ programme<sup>59</sup> v.1.13.1 was used. Individuals with more than 20% missing data were filtered out. We built 20 random datasets of 150 genotypes. For each of the 20 datasets, SMC++ was used with a generation time of one year and a mutation rate of  $6.5 \times 10^{-9}$  (ref. <sup>60</sup>). To avoid potential bias in the estimates owing to the long run of homozygosity, we filtered out homozygous regions longer than 5 kb in the 150 samples. For each of the 20 estimations, we used 5 different combinations of distinguished lineages, as suggested previously<sup>59</sup>. We then calculated the median of the 20 independent estimates for each time point.

SweeD (v.3.3.1) analysis was performed as previously<sup>61</sup> on chromosomes Ca1 to Ca8. To keep calculation time and resource into reasonable burdens while staying conservative in pointing genomic regions as being likely to be under positive selection, 2 random sub-samples of 251 landraces, proportional to 2,439 landraces for each geographical region, were considered. The analysis computes in each sub-sample a CLR for each SNP along the genome. We used a grid value of 10,000 for each chromosome, corresponding roughly to computing a CLR ratio every 9 kb. We considered the highest 1% CLR values for each sample and kept them as candidate SNPs for positive selection of the positions detected in both samples. Owing to linkage disequilibrium, a high CLR value detected on an SNP can result from selection acting on a nearby gene. Therefore, we computed a list of intervals that are likely to be targeted by selection from the list of SNPs detected under selection, without pointing to particular SNPs but including all SNPs within 10 kb of each other.

Effect of nucleotide variations on protein function was predicted with SIFT 4G<sup>21</sup> v.2.0.0. Putative deleterious mutations were identified with a SIFT score of less than 0.05. The *Medicago* genome was used as an outgroup to identify the derived alleles in the chickpea genome. Mutation burden was computed by counting the number of derived deleterious alleles present in constrained regions of the genome in each genotype as described before<sup>8</sup>.

### Genome-wide association analysis

Genome-wide association study (GWAS) analysis was performed using 3.94 million genome-wide SNPs and phenotypic data generated on 16 traits for 2 seasons and 6 locations. Only biallelic SNPs in cultivated genotypes were used in the GWAS analysis. Furthermore, the filtration was done with a minor allele frequency (MAF) cut-off of 0.05, missing rate cut-off of 0.8 and heterozygosity rate of 0.1. Marker trait association (MTA) analysis was then performed using a mixed linear model with the filtered HapMap file and phenotyping data. The first three PCs were used to control the population structure. The Manhattan plots and QQ plots were generated from the GWAS results. A  $P$  value of  $3.16 \times 10^{-7}$  was used to consider the MTA as significant.

### Identification of superior haplotypes

For haplotype analysis, we retained a SNP set for 3,171 cultivated chickpea lines according to the following criteria: (i) MAF > 0.001; and (ii) proportion of missing calls per SNP < 30%. The haplotypes present within trait-associated genes were examined and only homozygous calls were considered for haplotype analysis. The identified haplotypes were visualized in Flapjack<sup>62</sup> v.1.19.09.04.

For the haplo-phenotype analysis, haplotypes carrying only one genotype were removed from the analysis. The accessions were categorized on the

basis of haplotype groups, and together with phenotypic data, superior haplotypes were identified<sup>63</sup>. Haplotype-wise means for 100SW, days to flowering (DF) and YPP were compared to define superior haplotypes. Duncan's multiple range test was used for statistical significance.

### OCS approach

We used GEBV from the genomic prediction section for key production traits (YPP, 100SW, DF and days to maturity (DM)) to generate a genomic relationship matrix based on 754,576 SNPs. We used the breeding program implementation platform MateSel v.6.3 (<http://matesel.une.edu.au>) to generate an optimized mating design within desi, kabuli and intermediate types. The relative emphasis on the mean index versus co-ancestry was set by choosing the target degrees on the response surface<sup>24</sup>. We chose a target of 60 degrees to minimize the increase in population co-ancestry (maximize population genetic diversity) while achieving an acceptable rate of genetic gain. As this study aimed to maintain a diverse pre-breeding pool while making economic improvements, we followed the conservative approach for 'evolving gene banks' (ref. <sup>23</sup>).

We generated unique economic indices for desi and kabuli chickpea, which were calculated on a US\$ per ha basis and included yield (average GEBV for YPP over 9 sites) with a bonus price for large seeds (when average GEBV for 100SW over 9 sites exceeded the average for kabuli of +5.9 g) and earliness (average GEBV for DF and DM over 9 sites < 0 days). The base price for chickpea was assumed to be US\$400 per tonne, and YPP was converted to an equivalent grain yield value per hectare by assuming that the mean YPP of 18 g per plant is equivalent to 1.8 tonnes per hectare. The index was also adjusted for a price bonus for large seeds and earliness as follows. The starting values for GEBV for 100SW are low in desi candidates (mean  $-4.0$  g) and high in kabuli candidates (mean +5.9 g). Hence, the starting value for a price bonus for 100SW begins at GEBV + 5.9 g, and there is no bonus below this value. The price bonus per gram (GEBV 100SW > 5.9 g) is US\$35 per gram, which is added to the base price. Similarly, a bonus was provided in price per tonne for GEBV earliness (average of GEBV DF and GEBV DM). The average GEBV earliness in the desi group was  $-1.6$  days, and in the kabuli group was +2.4 days. The starting value for a price bonus for earliness begins at average GEBV 0 days; there is a bonus for negative values of US\$10 per day added to the base price and no bonus for positive values.

### Genomic prediction analyses

**Prediction using the interaction of genomic and environmental covariates.** As described previously<sup>25</sup>, three models, a basic model (E + L) with main effects of environments (E) and lines (L), a model (E + L + G) including the main effects of markers, and a genomic by environment interaction model (E + L + G + GE) were used. Three different SNP datasets (G1, cultivated accessions; G2, wild accessions; and G3, G1 + G2) were used as a genomic matrix (G), post-conventional quality controls on missing values (<20%) and MAF (>0.05). Phenotyping data for nine traits across 12 different year  $\times$  location combinations were used. The Pearson's correlation coefficient between observed phenotype and predicted genomic breeding value was used to estimate the accuracy of genomic prediction. Three different random cross-validation (CV) schemes, CV1 (evaluate the prediction accuracy of models when a certain percentage of lines are not observed in any environment), CV2 (estimates the prediction accuracy of models when some lines are evaluated in some environments but not in others) and CV0 (predicts an unobserved environment using the remaining environments as a training set) were used. CV1 and CV2 with fivefold cross-validation were implemented to generate the training and testing sets, and the prediction accuracy was assessed for each testing set. The permutation of the five subsets led to five possible training and validation datasets. This procedure was repeated 20 times, and 100 runs were performed for each trait-environment combination on each population. The



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same partition was used for the analysis of all the GS models. For CVO, each environment was predicted using the remaining environments. For fitting the GS models, the R package Bayesian Generalized Linear Regression (BGLR)<sup>64</sup> v.1.0.7 was used.

**Prediction using the WhoGEM method.** For WhoGEM analysis, 1,318 accessions with the validated geographical location were selected and used as a reference dataset. The SNP dataset was filtered for missing (>0.1) and MAF (<0.01) and used for a detailed search with ADMIXTURE<sup>65</sup> v.1.3.0 between  $K = 19$  and  $K = 30$  to identify the most likely number of admixture components. To confirm the admixture value, another method, DAPC (discriminant analysis of principal components), was used. The optimal number of admixture components in the WhoGEM method was obtained by comparing the predicted and recorded locations (ProvenancePredictor algorithm<sup>26</sup>) and fixed to  $K = 23$ .

A general linear model explored the relationships between the phenotypes and admixture components, and land types. A forward-backward algorithm was used to reduce the set of predictors to the most significant ones. The model is fitted on the whole dataset, and the significant factors are identified and conserved. A negative control (a model without any genetics (called environment-only)) is also fitted to the data. The models were fitted on the whole dataset, and the significant factors were identified and conserved.

A test of WhoGEM significance is given by a likelihood ratio test comparing the WhoGEM-based model and the environment-only-based model. The performances of the three models (full WhoGEM-based model, additive and environment-only model) are then evaluated using 100–300 replicates of a fivefold cross-validation scheme.

## Prediction using a haplotype-based approach

The SNP set was filtered, first by excluding all markers with more than two called alleles, missing (>10%) and MAF (<5%). A subset of 124,833 (20%) of 2.4 million high-quality SNPs were randomly selected to reduce the computational load in further analyses. Those SNPs were used to construct LD blocks and estimate local GEBVs for haplotypes of those LD blocks. Details on the method used to calculate local GEBVs for haplotypes of LD blocks are described in a previous report<sup>27</sup>.

We also ran a ridge-regression best linear unbiased prediction (BLUP) model in the R-package rrBLUP (ref.<sup>66</sup>) v.4.6.0 to predict marker effects for seven agronomic traits, then summed up the predicted allelic effects of each observed haplotype for all genome-wide LD blocks. Finally, we estimated variances among local GEBVs for haplotypes within each LD block to highlight regions in the genome showing molecular variation linked to observed phenotypic variation for the agronomic traits measured in the field trials.

## Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

## Data availability

The data that support the findings of this study have been deposited in the NCBI under accession code BioProject: PRJNA657888. The chickpea pan-genome assembly and annotations developed in this study are available at <https://doi.org/10.6084/m9.figshare.16592819>. The variant calls for each accession and phenotype data are available to download at <https://cegresources.icrisat.org/cicerseq>. Manhattan and QQ-plots for GWAS analysis are available at <https://doi.org/10.6084/m9.figshare.15015309> and <https://doi.org/10.6084/m9.figshare.15015315>, respectively. Source data are provided with this paper.

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**Competing interests** The authors declare no competing interests.

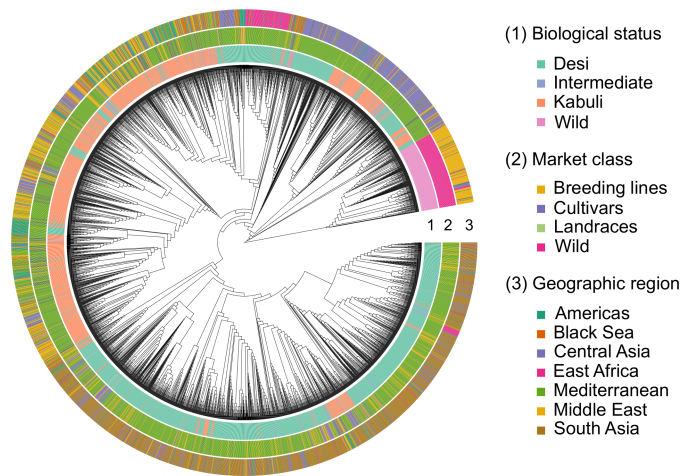
#### Additional information

**Supplementary information** The online version contains supplementary material available at <https://doi.org/10.1038/s41586-021-04066-1>.

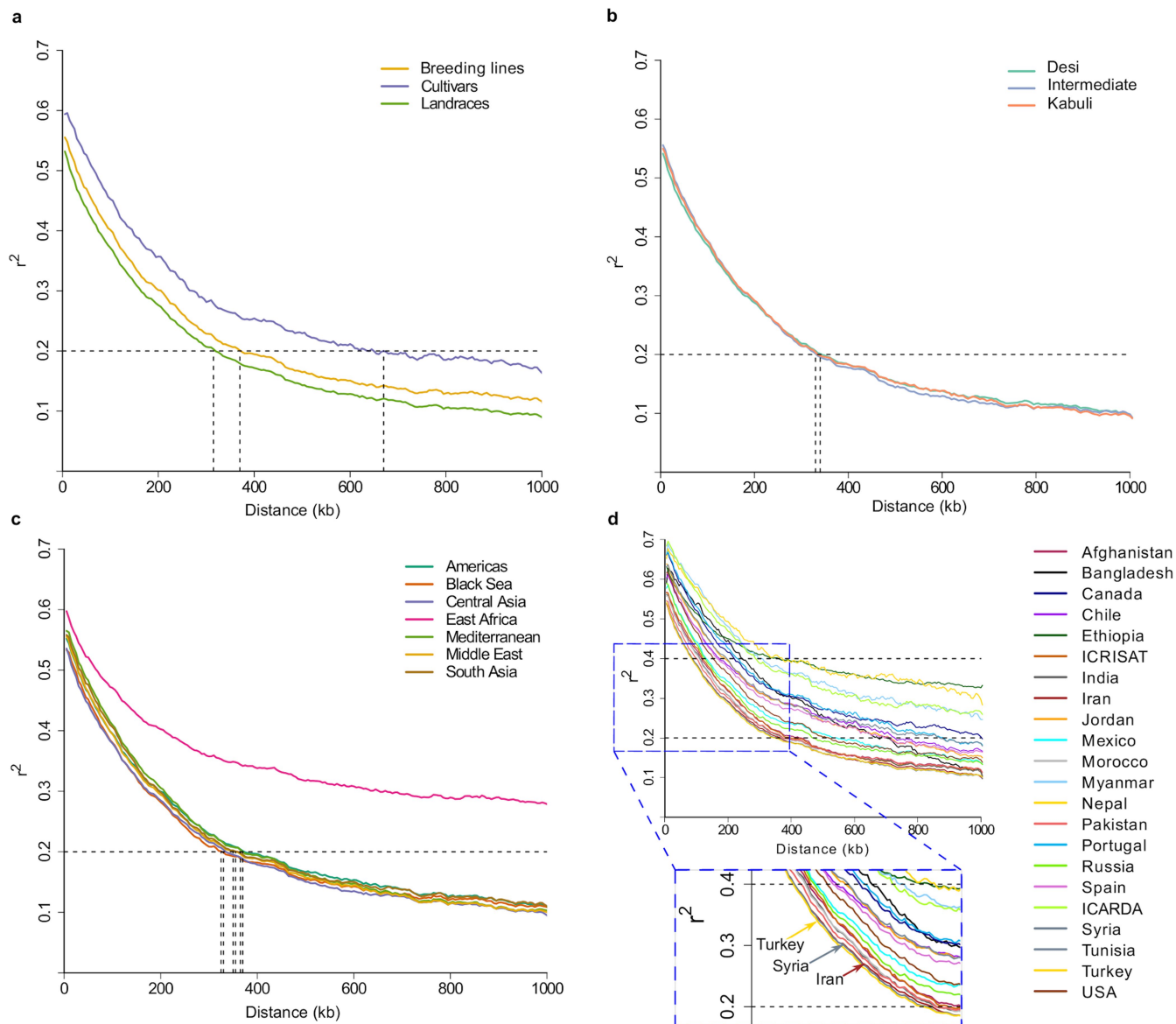
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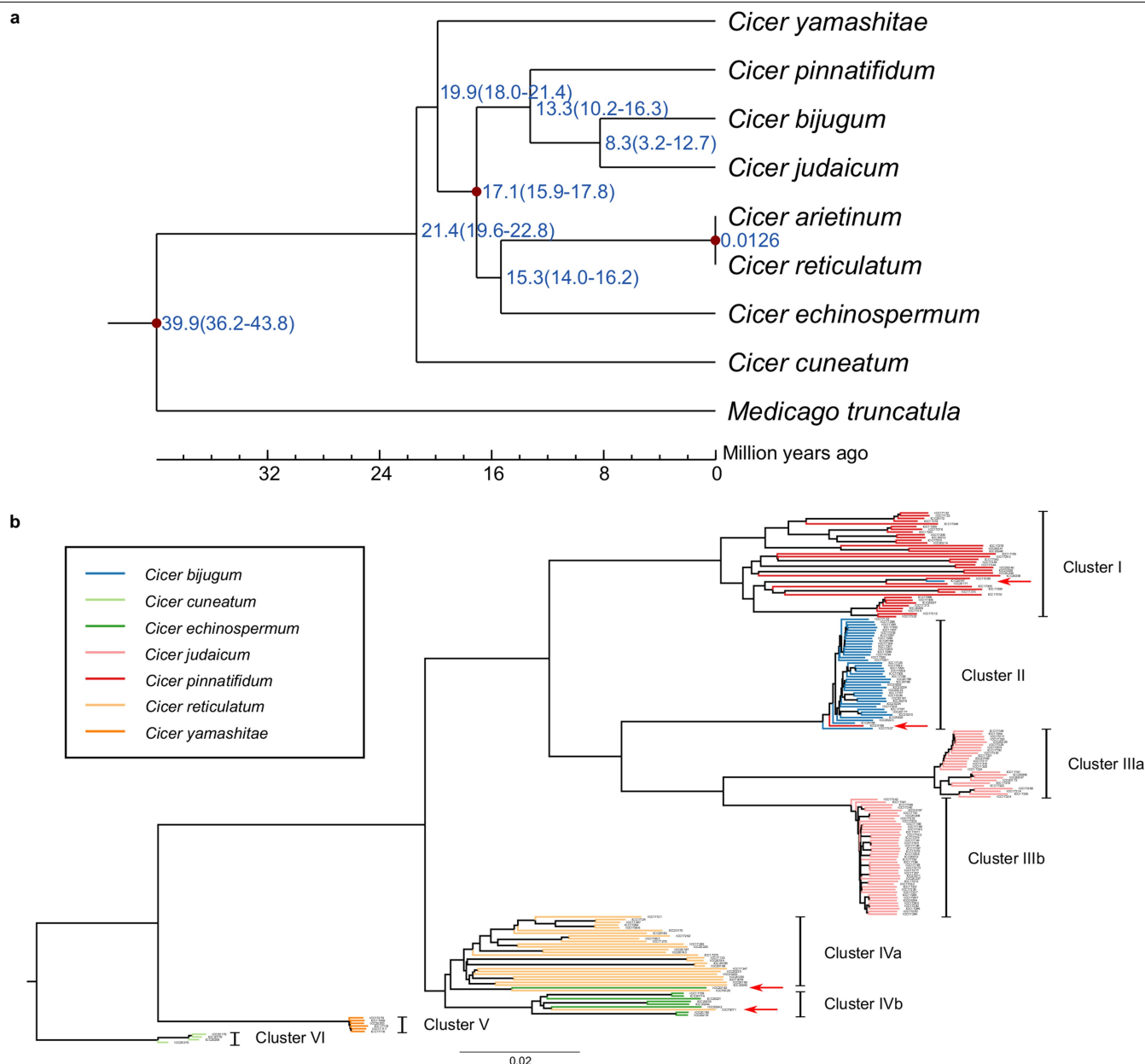
**Extended Data Fig. 1 | Phylogeny-based clustering.** Phylogenetic tree represents clustering of individuals, represented through respective tracks (from inside to outside), Track 1: Biological status; Track 2: Market class; and Track 3: Geographical regions. A clear outgroup of wild accessions is observed.



**Extended Data Fig. 2 | Linkage disequilibrium decay observed among cultivated chickpea genotypes.** (a) Rapid LD decay was observed in landraces (315 kb) based biological status followed by breeding lines (370 kb) and cultivars (670 kb). (b) Similar LD decay rate was observed among population based on market class, namely desi (340 kb), intermediate (330 kb) and kabuli (330 kb). (c) Among seven geographic populations, genotypes from Black Sea (352 kb) had lowest rate of LD decay followed by Central Asia (330 kb), Middle

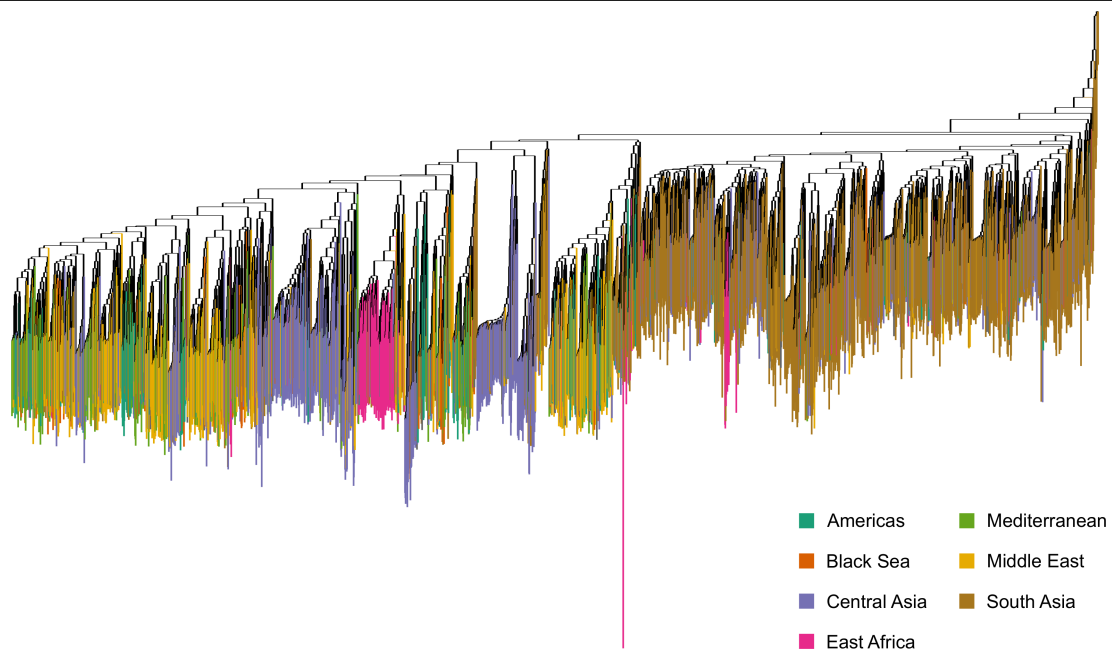
East (350 kb), South Asia (355 kb), Mediterranean (365 kb) and Americas (370 kb). The population East Africa had much slower LD decay compared to other population based geographic regions. (d) Cultivated accessions from countries Turkey (306.51 kb), Syria (316.22 kb) and Iran (320.61 kb) had more rapid decline of LD decay compared to cultivated accessions from other countries, indicating more recombination events and haplotype diversity/number.



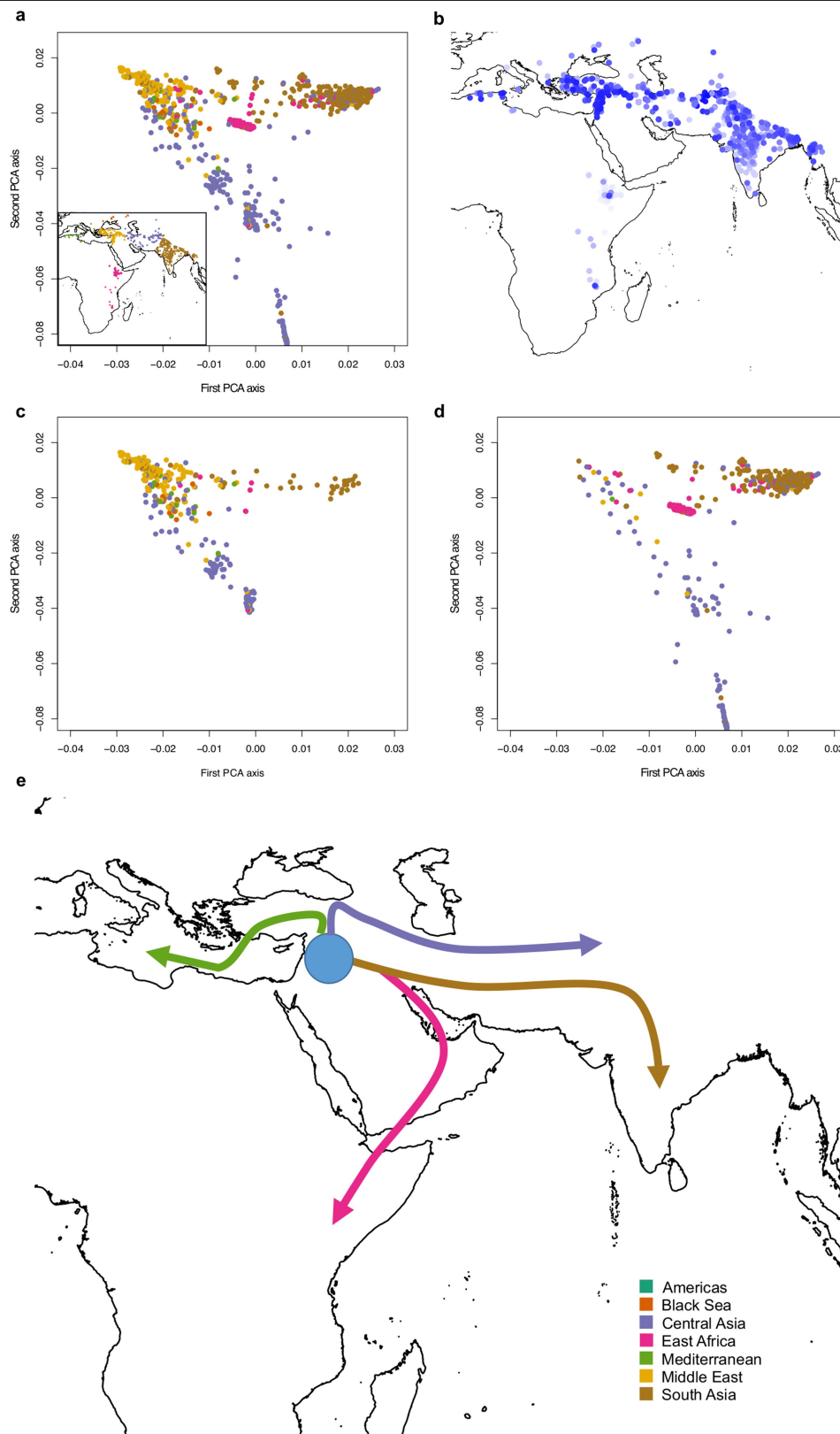


**Extended Data Fig. 3 | *Cicer* species evolution.** (a) Speciation and divergence time for eight species in the genus *Cicer*. The maximum likelihood phylogenetic tree showed clear out-grouping of *C. cuneatum* from the other *Cicer* species and *C. reticulatum* being nearest to the cultivated chickpea species (*C. arietinum*). Three time-calibration points (0.007–0.013 Ma for *C. reticulatum*–*C. arietinum*, 12.2–17.4 Ma for *C. arietinum*–*C. pinnatifidum*, and 30.0–54.0 Ma for *C. arietinum*–*M. truncatula*) were used for estimating divergence time. The nearest wild species (*C. reticulatum* and *C. echinospermum*) related to the cultivated *C. arietinum* were estimated to be diverged from other *Cicer* species around ~15.3 (14.0–16.2) Ma. (b) Genetic diversity among wild species accessions. Phylogenetic tree constructed based on SNPs grouped 195 wild species accessions into six clusters. A clear grouping for accessions of *C. judaicum*,

*C. yamashitae* and *C. cuneatum* was observed in Cluster III, Cluster V and Cluster VI, respectively. However, ICC 20168 (one *C. pinnatifidum* accession; red colour) grouped along with *C. bijugum* accessions in Cluster II; similarly, ICC20167 (one *C. bijugum* accession; blue colour) grouped along with *C. pinnatifidum* accessions in Cluster I. Cluster III and Cluster IV were divided into two sub-clusters each, in which both sub-clusters of Cluster III possessed all accessions of *C. judaicum*. In Cluster IVa we observed grouping of all *C. reticulatum* accessions except one *C. echinospermum* accession (ICC 20192; green colour); similarly, in Cluster IVb one accession of *C. reticulatum* (ICC 73071; golden-yellow colour) grouped along with *C. echinospermum* accessions.

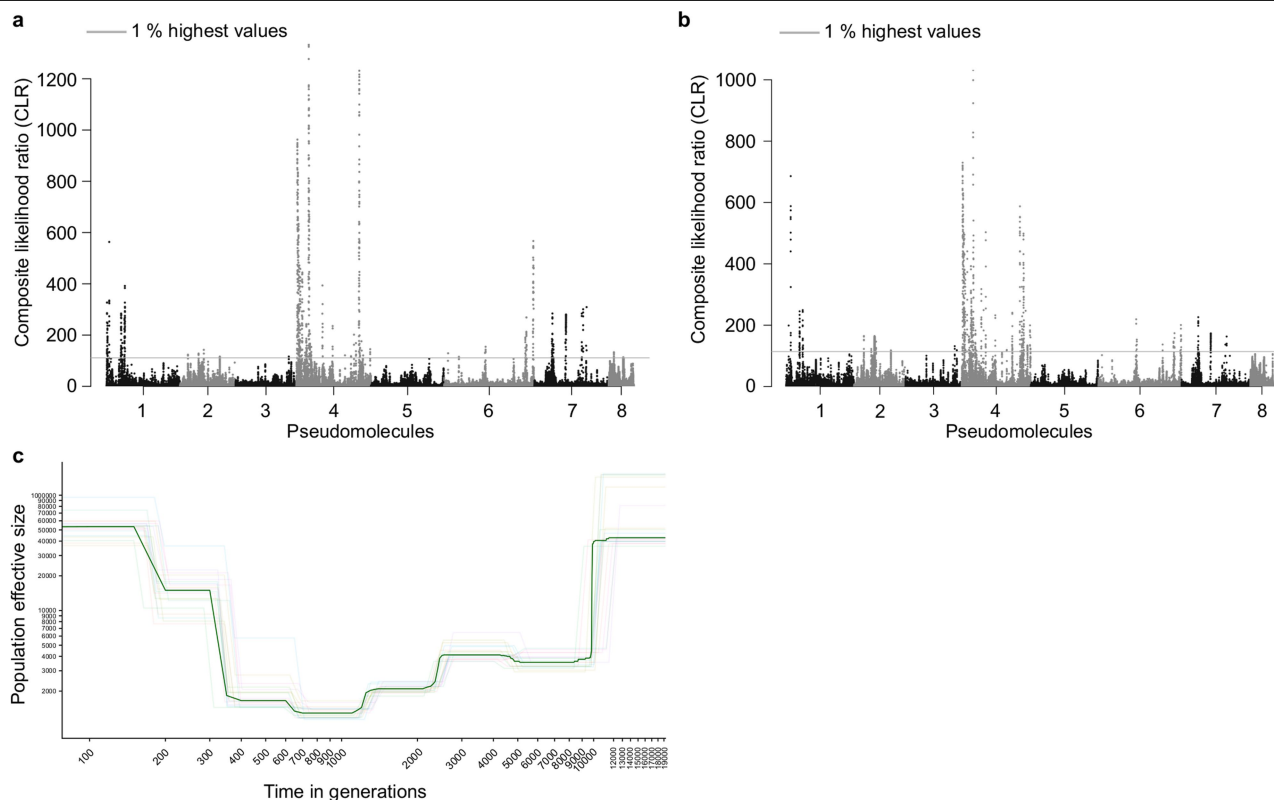


**Extended Data Fig. 4 | Phylogenetic tree based on  $F_{ST}$ .** Accessions from Mediterranean region, Middle East, Americas and Black Sea regions were clustered together, and South Asia as a separate cluster.



**Extended Data Fig. 5 | Relationship route of chickpea diffusion and seed morphology.** (a) PCA analysis for landraces. (b) Distance to the most extreme cultivated sample (closest to wild relatives) were plotted on the map. (c, d) For landraces with large seed morphology (kabuli; c) and small seed morphology (desi; d) indicated that small seed was mainly found in East-Asia, South-East

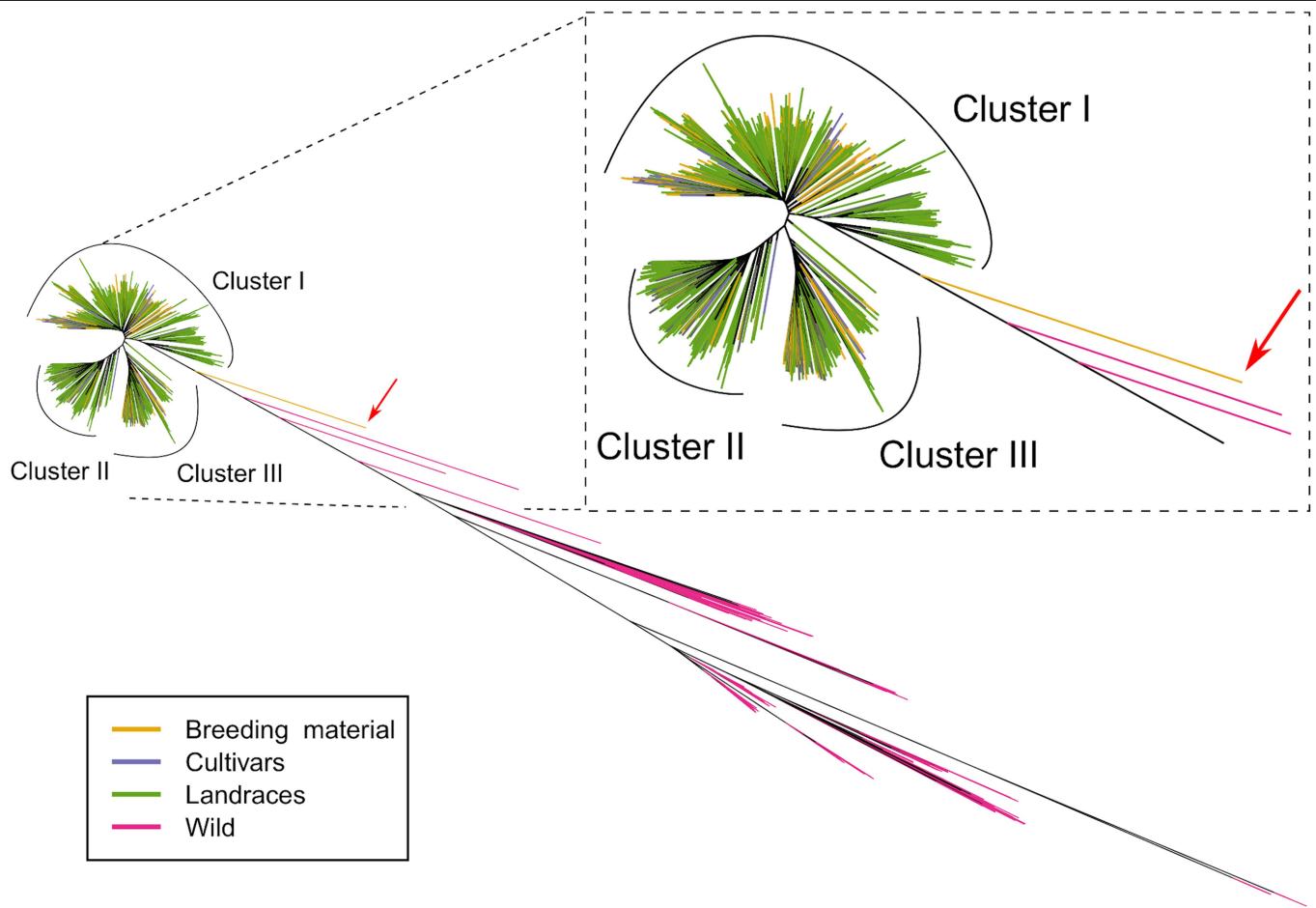
Asia and Africa. These suggest large and small seeds were selected independently during chickpea diffusion of agriculture. (e) PCA results summarised a Central Asian diffusion alongside a Mediterranean diffusion, and a South Asian diffusion associated with the diffusion to East Africa.



**Extended Data Fig. 6 | Composite likelihood ratio values along the chickpea genome and inference of past evolution of effective size.** The composite likelihood ratio for chromosomes 1 to 8 on the x axis is computed for two random subsets of 251 individuals: subset 1 (a) and subset 2 (b). Horizontal grey line shows the threshold above which the highest 1% CLR values are found. (c) Using sequentially markovian coalescent as implemented in SMC++

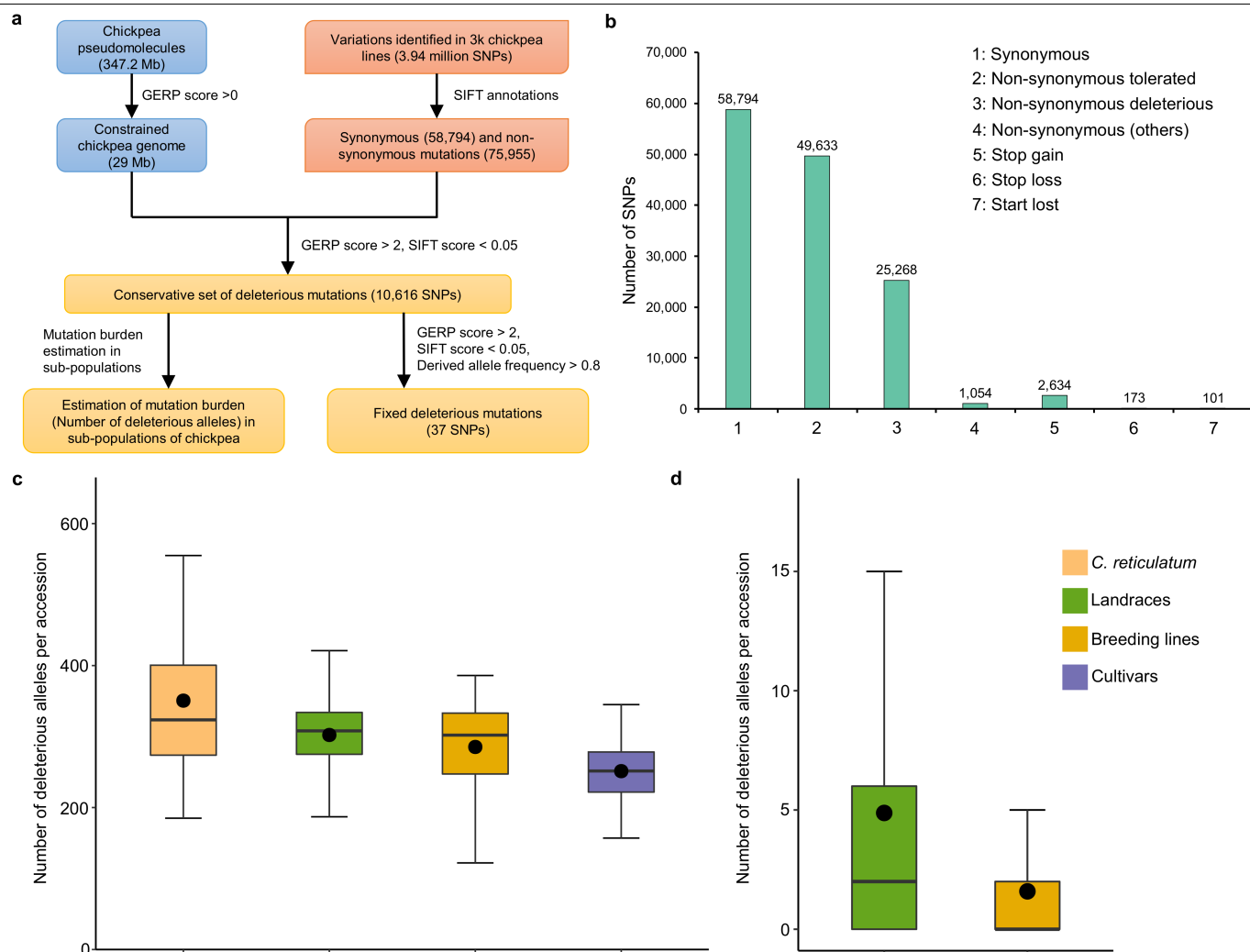
(Terhorst et al. 2017), we reconstructed the past history of effective size for 20 sets of 150 randomly chosen cultivated genotypes (thin lines). We computed at each time point the median of the estimated histories and plotted it (bold lines). Focus was made for the plotting on timeframe 100 – 20,000 generations ago. Both x and y axes are log-scaled.





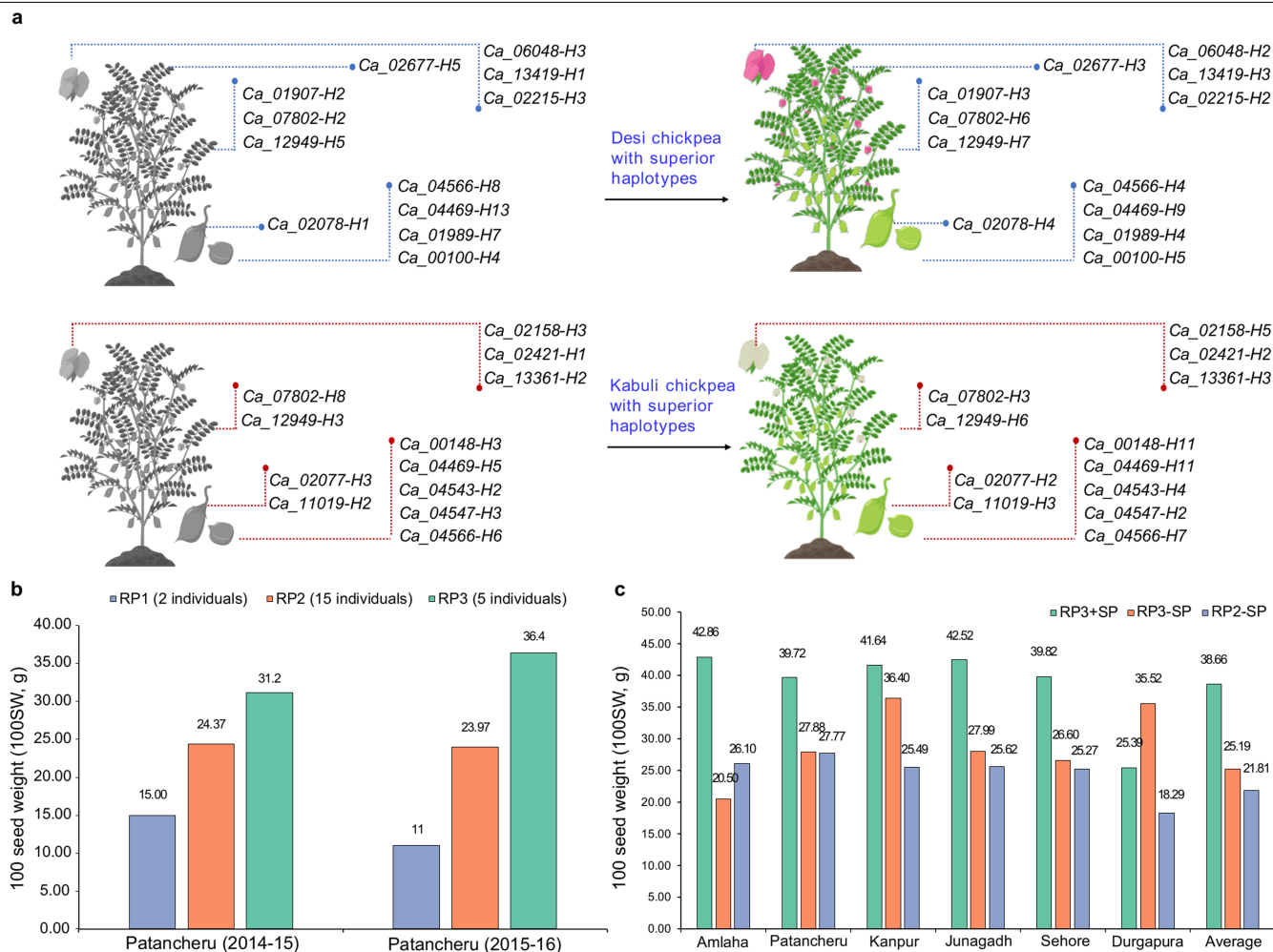
**Extended Data Fig. 7 | Neighbour-joining trees constructed using SNPs present on the pseudomolecules indicates a clear out-grouping of wild species accessions from cultivated accessions.** The cultivated accessions

formed three distinct clusters. One landrace from East Africa (ICC 16369) (red arrow) grouped together with wild species accessions.



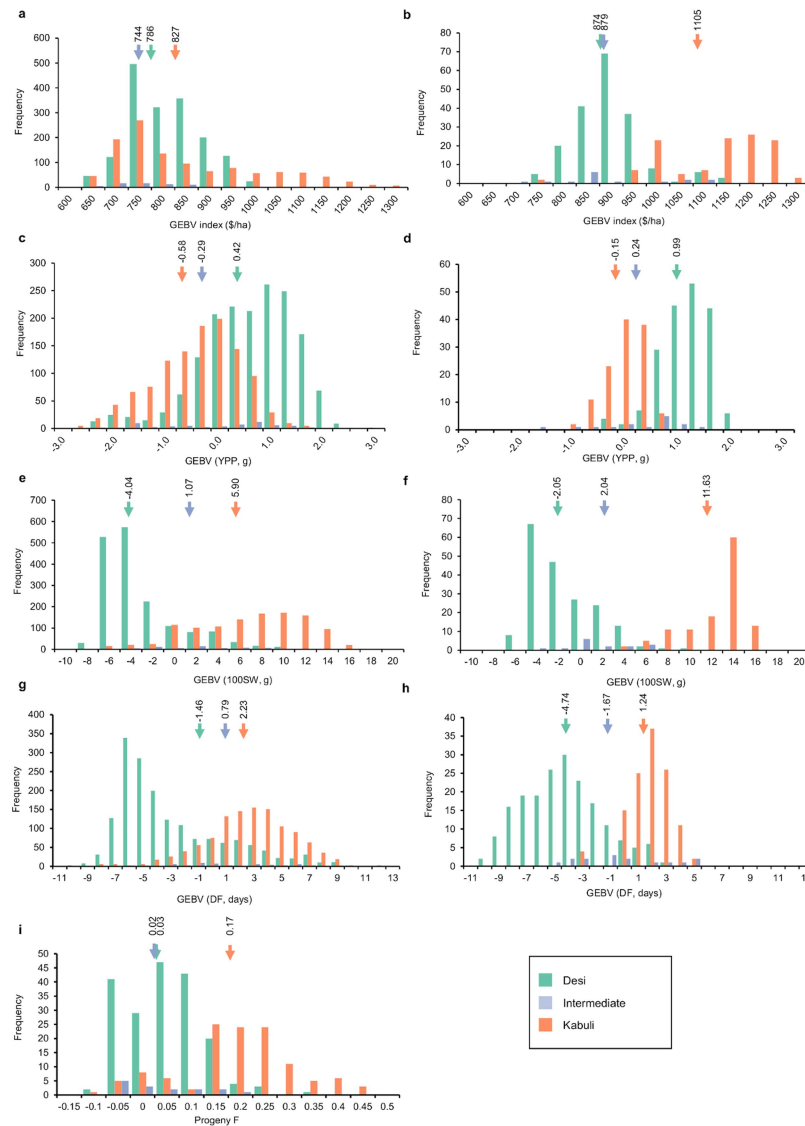
**Extended Data Fig. 8 | Genetic loads in chickpea.** a) A snapshot of steps and parameters used to estimate the mutation burden and fixed deleterious alleles. b) Variant annotation using SIFT revealed higher non-synonymous mutations, of which non-synonymous deleterious variants were used to identify deleterious mutations. c) Mutation burden analysis indicated a 17.88% decrease (two-tailed Welch's t-test;  $t = 2.525$ ,  $df = 27$ ,  $p = 0.01772$ ,  $CI = 95\%$ ) in mutation burden in cultivated ( $n = 2987$ ) as compared to progenitor (*C. reticulatum*;  $n = 28$ ). d)

Mutation burden for genomic regions under selection showed that landraces ( $n = 2439$ ) contained 206.91% higher (two-tailed Welch's t-test,  $t = -17.087$ ,  $df = 1645$ ,  $p = 2.195676 \times 10^{-60}$ ,  $CI = 95\%$ ) deleterious mutations than breeding lines ( $n = 396$ ). The black solid dots in box plots represent mean values for the respective population. Each of the box plots shows the upper and lower whisker, the 25% and 75% quartiles, the median (as solid line) and the mean (black dot).



**Extended Data Fig. 9 | Towards developing tailored chickpea with superior haplotypes for yield and related traits.** (a) Representative desi and kabuli chickpea plant (on left) carrying inferior haplotype combination for key traits including 100 seed weight (100SW), days to maturity (DM), plant height (PLHT), pods per plant (PPP), and plot yield (PY). Target desi and kabuli chickpea plant (on right) carrying superior haplotype for 100SW, DM, PLHT, PPP and PY. New breeding lines can be developed by introgressing the superior haplotype combination through haplotype-based breeding. (b) Comparison of average performance among RP1 vs RP2 vs RP3 varieties for 100SW at Patancheru location. An increase in 100SW between the varieties of RP1 vs RP3 was observed, whereas no differences were observed in the case of RP1 vs RP2 and RP2 vs RP3 varieties (datasets of ICRISAT 2014-15 and 2015-16). RP1 indicates

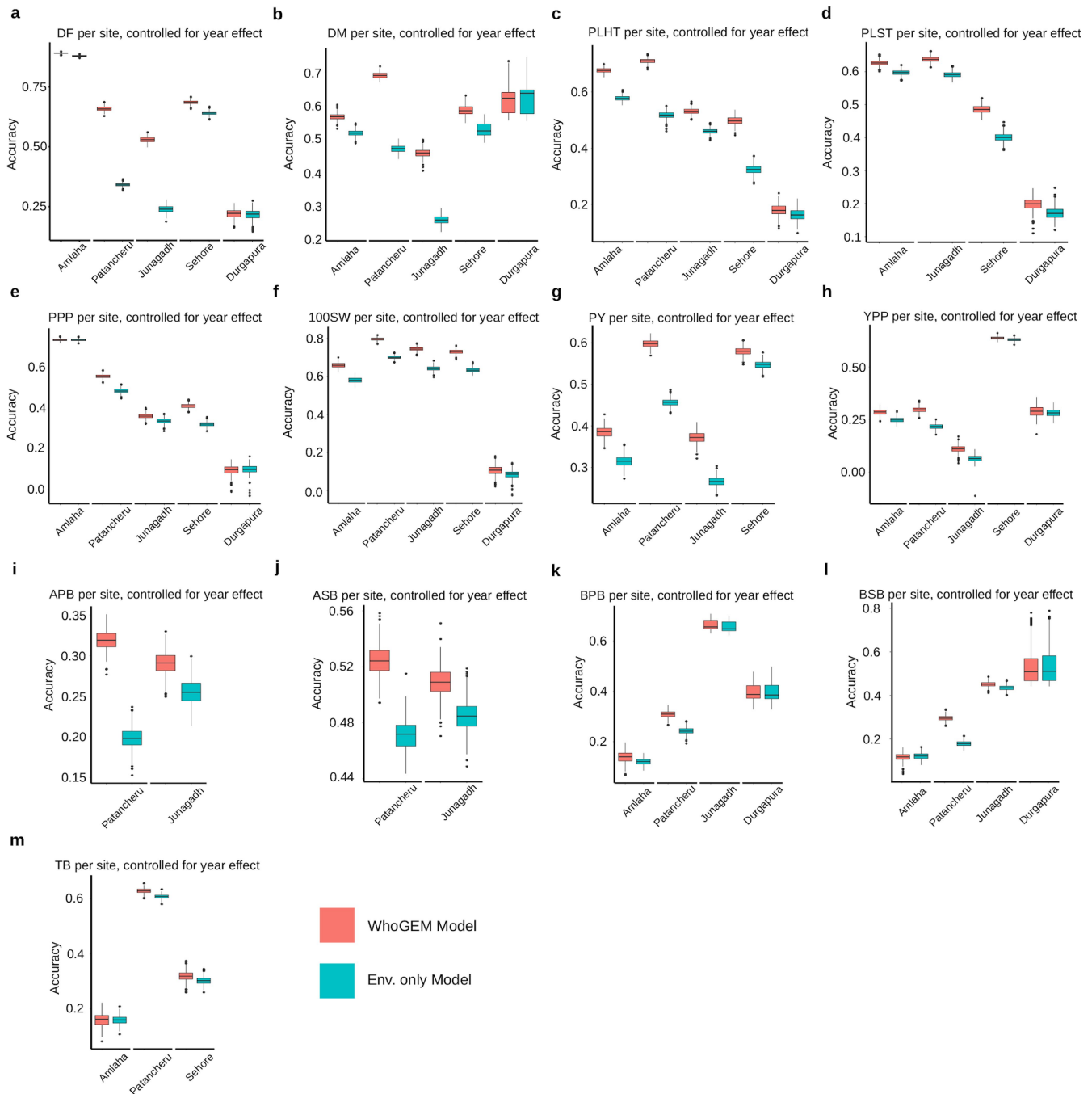
chickpea varieties released before 1993, RP2 indicates chickpea varieties released between 1993-2002 and RP3 indicates chickpea varieties released after 2002. (c) Comparison of RP2 and RP3 varieties for 100SW (with and without superior haplotypes) for six locations. A difference between lines carrying the superior haplotypes (RP3+SP) for 100SW was observed in comparison to those which did not (RP3-SP and RP2-SP) except for the Durgapura location. However, marked differences were also observed between the RP3-SP and RP2-SP lines, except for Patancheru and Amlaha locations. RP3+SP indicates RP3 varieties with superior haplotypes, RP3-SP indicates RP3 varieties without 100SW superior haplotype and RP-SP indicates RP2 varieties without 100SW superior haplotype. RP2 indicates chickpea varieties released between 1993-2002 and RP3 indicates chickpea varieties released after 2002.



**Extended Data Fig. 10 | Response to OCS based on mating allocation for candidate parents and predicted cycle 1 progeny family means in economic index.** Index increased from parents to cycle 1 progeny in the kabuli group by US\$274/ha, and in the desi group by US\$94/ha, and reflects the high value of large seeds in the kabuli group. Arrows indicate the population mean GEBVs for desi (green), kabuli (orange) and intermediate (blue) groups. (a) Response to selection for candidate parents. (b) Response to selection for predicted cycle 1 progeny family. (c, d) Response to OCS among desi, kabuli and Intermediate accessions is shown for genomic estimated breeding values (GEBVs) for yield per plant (YPP, g). YPP increased by 0.6 g (4.5%) above the average candidate parent YPP (13.2 g) in the desi group, and by 0.4 g (3.3%) above the average candidate parent yield (12.2 g) in the kabuli group. Arrows indicate the population mean GEBVs for desi (green), kabuli (orange) and intermediate (blue) groups. (e) candidate parents. (f) predicted cycle 1 progeny family. (g, h) Response to OCS for GEBVs for 100 seed weight (100SW, g). 100SW increased by 2.0 g

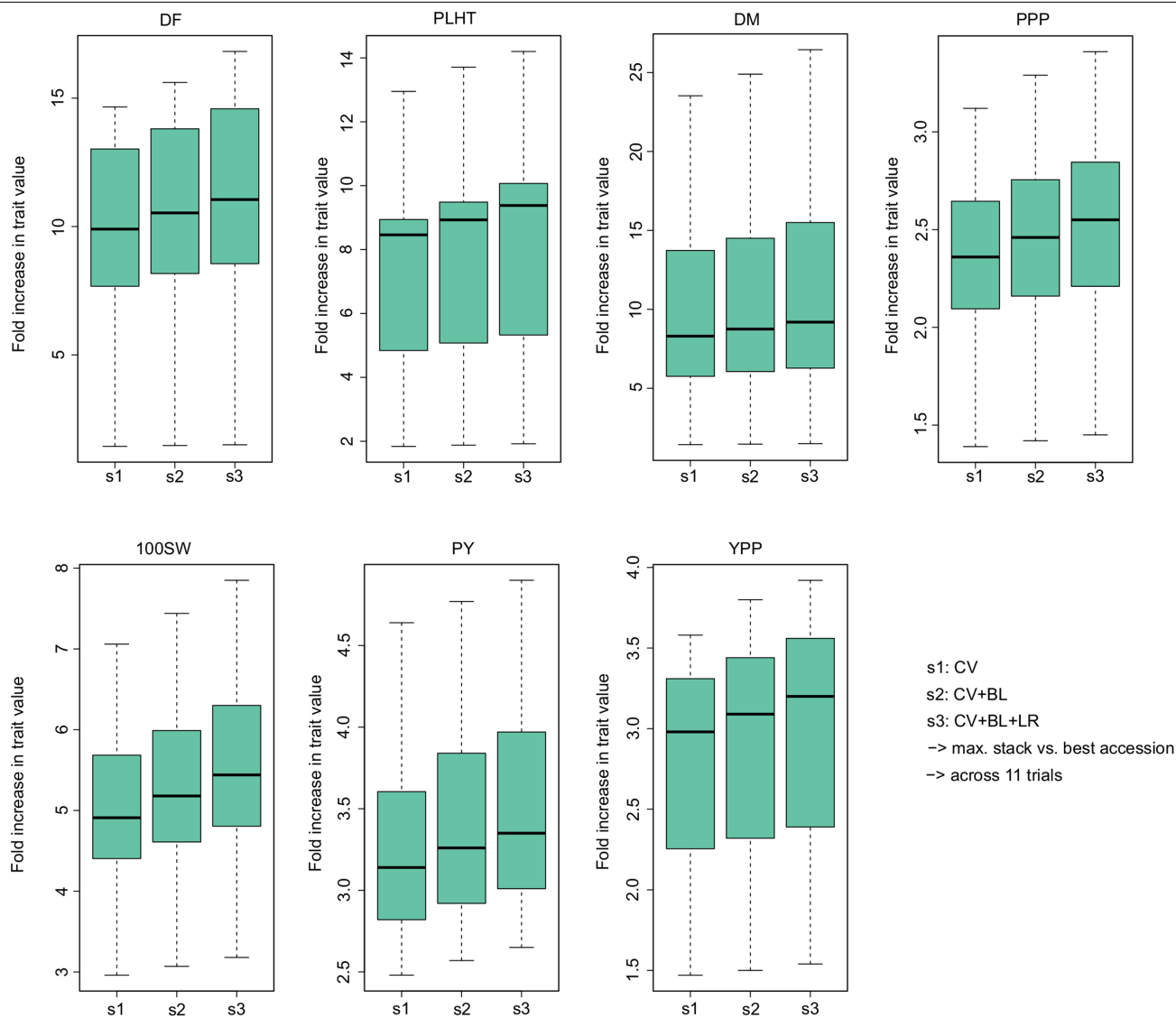
(12.7%) above the average candidate parent 100SW (15.0 g) in the desi group, and by 5.7 g (22.9%) above the average candidate parent 100SW (24.9 g) in the kabuli group. Arrows indicate the population mean GEBVs for desi (green), kabuli (orange) and intermediate (blue) groups. (e) candidate parents. (f) predicted cycle 1 progeny family. (g, h) Response to OCS for GEBVs for days to flower (DF). DF decreased by 3.3 d (–4.7%) below the average candidate parent DF (68.6 d) in the desi group, and by 1.0 d (–1.4%) below the average candidate parent DF (72.3 d) in the kabuli group. Arrows indicate the population mean GEBVs for desi (green), kabuli (orange) and intermediate (blue) groups. (g) candidate parents. (h) predicted cycle 1 progeny family. (i) Predicted average inbreeding (F) in cycle 1 progeny in among desi, kabuli and intermediate accessions. Progeny inbreeding increased by 0.170 in the kabuli group, by 0.025 in the desi group, and by 0.015 in intermediate group. Arrows indicate the population mean GEBVs for desi (green), kabuli (orange) and intermediate (blue) groups.





**Extended Data Fig. 11 | WhoGEM prediction accuracies for different traits in different sites.** A general linear model was used for predicting performance in selected (with a geolocation) 1,318 cultivated chickpea accessions. At each site, 200 replicates of a fivefold cross-validation scheme are applied to estimate the accuracies of WhoGEM model (phenotype as a function of admixture components and market class) compared to environment-only model i.e. a model without genetic effects. Tests of WhoGEM significance are given by likelihood ratio tests between the WhoGEM-based models and the environment-only-based model. Phenology traits: (a) days to flowering (DF),

(b) days to maturity (DM), (c) plant height (PLHT) and (d) plant stand (PLST); Production traits: (e) pods per plant (PPP), (f) 100 seed weight (100SW), (g) plot yield (PY) and (h) yield per plant (YPP) and Plant architecture traits: (i) apical primary branch (APB), (j) apical secondary branch (ASB), (k) basal primary branch (BPB), (l) basal secondary branch (BSB), and (m) tertiary branch (TB). Each of the box plots shows the upper and lower whisker, the 25% and 75% quartiles and the median (as solid line) of the fold change ( $n = 1,318$  cultivated accessions)..



**Extended Data Fig. 12 | Assessment of trait improvement potential by stacking the superior haplotypes for target traits.** The genotypes were classified into three different groups (cultivars (CV,  $n = 152$ ); breeding lines (BL,  $n = 396$ ) and landraces (LR,  $n = 2,439$ )) and these genotypes were grouped in three subgroups s1 (CV), s2 (CV+BL) and s3 (CV+BL+LR). Local GEBVs for haplotypes were calculated by firstly grouping SNP markers based on their

pairwise linkage disequilibrium, and then summing up allele effects for each haplotype of each block. The best possible genotype for each trait was generated in silico by adding up the best haplotypes across the whole genome. This in silico genotype was then compared to the accession with the highest GEBV. Each of the box plots shows the upper and lower whisker (indicated by dashed lines), the 25% and 75% quartiles and the median (as solid line).

Extended Data Table 1 | Summary of genome diversity features

Germplasm	SNP			Structural variations					LD decay (kb)	Nucleotide diversity (π)
	Total SNP	Private SNP	Population enriched SNP	INS	DEL	INV	ITX	CTX		
Cultivated (3171)	3,941,492	--	--	139,483	47,882	61,171	417	2,410	--	--
Market class										
Desi	3,383,484	185,645	1,223	139,481	47,832	61,149	415	2,409	340	4.13 × 10 <sup>-4</sup>
Intermediate	1,654,675	198	15	125,843	40,448	52,359	381	2,068	330	4.73 × 10 <sup>-4</sup>
Kabuli	2,563,386	60,120	1,026	139,430	47,722	61,083	416	2,404	330	4.55 × 10 <sup>-4</sup>
Biological status										
Cultivars	1,467,163	87	156	106,355	35,115	39,828	291	1,743	670	3.09 × 10 <sup>-4</sup>
Breeding lines	2,417,773	1,790	17	138,633	47,055	60,230	409	2,381	370	4.63 × 10 <sup>-4</sup>
Landraces	3,415,287	311,581	112	139,483	47,882	61,171	417	2,410	315	4.75 × 10 <sup>-4</sup>
Geographic regions										
Americas	1,785,351	1,870	23	129,681	43,488	55,256	388	2,238	370	4.53 × 10 <sup>-4</sup>
Black Sea region	1,558,472	58	469	118,806	40,406	48,517	361	1,985	325	4.72 × 10 <sup>-4</sup>
Central Asia	2,444,447	22,821	6,911	139,418	47,839	61,107	416	2,406	330	4.74 × 10 <sup>-4</sup>
East Africa	2,049,973	8,469	11,356	124,723	42,448	52,510	375	2,170	--	3.69 × 10 <sup>-4</sup>
Mediterranean region	1,881,691	347	10	133,869	45,577	57,563	398	2,300	365	4.36 × 10 <sup>-4</sup>
Middle East	2,173,446	3,881	36	138,710	47,376	60,526	414	2,396	350	4.52 × 10 <sup>-4</sup>
South Asia	2,670,568	28,856	71	139,431	47,710	61,037	414	2,407	355	3.62 × 10 <sup>-4</sup>
Wild Species										
Wild (195)	19,574,878									5.47 × 10 <sup>-3</sup>
<i>Cicer bijugum</i>	5,928,544	308,193	811,415	--	--	--	--	--	--	4.91 × 10 <sup>-4</sup>
<i>Cicer cuneatum</i>	3,445,598	888,326	1,498,916	--	--	--	--	--	--	3.43 × 10 <sup>-4</sup>
<i>Cicer echinospermum</i>	4,055,280	353,170	926,927	--	--	--	--	--	--	1.85 × 10 <sup>-3</sup>
<i>Cicer judaicum</i>	6,406,872	1,469,225	1,364,687	--	--	--	--	--	--	1.49 × 10 <sup>-3</sup>
<i>Cicer pinnatifidum</i>	7,630,762	859,830	1,000,204	--	--	--	--	--	--	1.40 × 10 <sup>-3</sup>
<i>Cicer reticulatum</i>	6,419,606	1,000,994	758,515	287,854	67,351	58,070	446	2,066	--	2.20 × 10 <sup>-3</sup>
<i>Cicer yamashitae</i>	3,939,064	1,215,444	1,492,985	--	--	--	--	--	75	3.16 × 10 <sup>-4</sup>

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### Software and code

Policy information about [availability of computer code](#)

Data collection	Sequencing data was generated on Illumina HiSeq2500.
Data analysis	Admixture (v1.3.0), AUGUSTUS (v3.1), BCFTools (v1.4), BGLR (V1.0.7), BLAST (v2.2.31), BreakDancer (v1.1.2), BUSCO (odb10), BWA (v0.7.15), CD-HIT (v4.81), Control-FREEC (v11.0), EIGENSOFT (v7.2.0), EnrichmentPipeline ( <a href="https://sourceforge.net/projects/enrichmentpipeline">https://sourceforge.net/projects/enrichmentpipeline</a> ), EVM (v1.1.1), Flapjack (v1.19.09.04), GAPIT3 (v20191108), GATK (v3.7), GATK (v3.8.1), Gblocks (v0.91b), gccount ( <a href="http://bioinfo-out.curie.fr/projects/freec/src/gccount.tar.gz">http://bioinfo-out.curie.fr/projects/freec/src/gccount.tar.gz</a> ), GeneWise (v2.4.1), GERP++ (May 22 2011), KEGG (v87.0), LASTZ (v1.4.00), Matesel (v6.3), MCMCTree (v4.4), Megahit (v1.2.9), MUSCLE (v3.8.31), NUCmer (v4.0.0beta2), PHYLIP (v3.6), Pindel (v0.2.5b9), PLINK (v1.90), PopLDdecay (v3.29), RAxML (v8.2.12), rrBLUP (v4.6.0), SAMTools (v1.2), SelectionTools (v19.4), SIFT 4G (v2.0.0), SMC++ (v1.13.1), SweeD (v3.3.1), SWISS-PROT (release-2018_07), VCFtools (v0.1.13).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

### Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

The data that support the findings of this study has been deposited in the NCBI under accession code BioProject: PRJNA657888. The chickpea pan-genome assembly



and annotations developed in this study are available at doi: 10.6084/m9.figshare.16592819. The variant calls for each accession and phenotype data are available to download at <https://cegresources.icrisat.org/cicerseq>. Manhattan and QQ-Plots for GWAS analysis are available at doi:10.6084/m9.figshare.15015309 and doi:10.6084/m9.figshare.15015315, respectively. BUSCO (odb10) and SWISS-PROT (release-2018\_07).

## Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☒ Life sciences ☐ Behavioural & social sciences ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://nature.com/documents/nr-reporting-summary-flat.pdf)

## Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	We sequenced 3000 accessions from global chickpea composite collection, which was developed by ICRISAT genebank in collaboration with ICARDA to define the genetic structure and represent the maximum diversity for the isolation of allelic variants of candidate gene associated with beneficial traits. The composite collection is a useful resource for detecting new sources of genetic variation and allelic variants of candidate gene(s) associated with beneficial traits, identifying diverse lines for use in functional and comparative genomics, in mapping and cloning gene(s), and in applied breeding (Upadhyaya et al. 2005, Plant Genetic Resources).
Data exclusions	Genotyping data was filtered using various well established criteria including % of missing, minor allele frequency and others. Similarly low quality phenotyping data from 3 site/year combination was filtered out. These exclusion have been defined for each analysis in the Methods section.
Replication	The composite collection, along with very promising checks (Annigeri, G130, ICCV10, JG11, KAK2 & L550) lines, were evaluated in an augmented block design. The experiment was conducted at Six locations Patancheru, Amlaha, Junagadh, Kanpur, Durgapura and Sehore during the post-rainy season of 2014-15 and 2015-16 years. For sequencing data, no replication was attempted.
Randomization	Analysis of the phenotyping data was performed by considering block as random and entry as fixed effects using the restricted maximum likelihood estimation procedure. Different populations in the analysis were defined based on passport information for germplasm accessions. For instance, based on seed type all cultivated (3171) accessions were divided into three populations/groups namely desi, kabuli and intermediate. Similarly, we also grouped accessions based on biological status (wild, landraces, breeding lines and cultivars) and their country of origin.
Blinding	No blinding. All data were processed equally.

## Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

### Materials & experimental systems

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input checked="" type="checkbox"/>	<input type="checkbox"/> Animals and other organisms
<input checked="" type="checkbox"/>	<input type="checkbox"/> Human research participants
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern

### Methods

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging