

# The pattern of genetic variability in a core collection of 2,021 cowpea accessions

Christopher J. Fiscus ,<sup>1,6</sup> Ira A. Herniter ,<sup>1,7</sup> Marimagne Tchamba,<sup>2</sup> Rajneesh Paliwal ,<sup>2</sup> María Muñoz-Amatriaín ,<sup>3</sup> Philip A. Roberts,<sup>4</sup> Michael Abberton ,<sup>2</sup> Oluwafemi Alaba ,<sup>1</sup> Timothy J. Close ,<sup>1,5</sup> Olaniyi Oyatomi ,<sup>2,\*</sup> Daniel Koenig ,<sup>1,5,\*</sup>

<sup>1</sup>Department of Botany and Plant Sciences, University of California, Riverside, Riverside, CA 92521, USA

<sup>2</sup>International Institute of Tropical Agriculture (IITA), Ibadan 200001, Nigeria

<sup>3</sup>Departamento de Biología Molecular, Universidad de León, León 24071, Spain

<sup>4</sup>Department of Nematology, University of California, Riverside, Riverside, CA 92521, USA

<sup>5</sup>Institute for Integrative Genome Biology, University of California, Riverside, Riverside, CA 92521, USA

<sup>6</sup>Present address: Department of Ecology and Evolutionary Biology, University of California, Irvine, Irvine, CA 92697, USA

<sup>7</sup>Present address: Department of Plant Biology, Rutgers University, New Brunswick, NJ 08901, USA

\*Corresponding author: Department of Botany & Plant Sciences, 2142 Batchelor Hall, University of California, Riverside, 900 University Avenue Riverside, California 92521 USA. Email: dkoenig@ucr.edu; \*Corresponding author: IITA Headquarters, PMB 5320, Oyo Road, Ibadan 200001, Oyo State, Nigeria. Email: o.oyatomi@cgiar.org

Cowpea is a highly drought-adapted leguminous crop with great promise for improving agricultural sustainability and food security. Here, we report analyses derived from array-based genotyping of 2,021 accessions constituting a core subset of the world's largest cowpea collection, held at the International Institute of Tropical Agriculture (IITA) in Ibadan, Nigeria. We used this dataset to examine genetic variation and population structure in worldwide cowpea. We confirm that the primary pattern of population structure is two geographically defined subpopulations originating in West and East Africa, respectively, and that population structure is associated with shifts in phenotypic distribution. Furthermore, we establish the cowpea core collection as a resource for genome-wide association studies by mapping the genetic basis of several phenotypes, with a focus on seed coat pigmentation patterning and color. We anticipate that the genotyped IITA Cowpea Core Collection will serve as a powerful tool for mapping complex traits, facilitating the acceleration of breeding programs to enhance the resilience of this crop in the face of rapid global climate change.

**Keywords:** cowpea; population structure; GWAS; seed pigmentation

## Introduction

Enhancing the sustainability of agriculture is essential to feeding the world population, especially in the context of climate change. Among crops, legumes are unique in their ability to improve soil fertility through symbiotic association with rhizobia, which reside within root nodules to facilitate nitrogen fixation (Nap and Bisseling 1990). This quality has made legumes important components of crop rotation and intercropping, both of which help to increase productivity (Reckling et al. 2016). In addition, grain legumes are notable for the high protein content of their grains (termed "pulses"), and their amino acid compositions complement those of cereals, making them exceptionally nutritious for both human and animal diets. Productivity gains in grain legumes over the past 50 years have been attributed primarily to expanded planting area rather than to advances in crop improvement as seen in other types of crops, such as cereals (Foyer et al. 2016). As such, the development of genetic resources linking genotype to phenotype is crucial to improve grain legumes and to advance crop research and breeding.

Cowpea [*Vigna unguiculata* (L.) Walp.] is a leguminous crop of global importance as a source of both food and fodder. The crop is produced around the world with the majority of contemporary

production in West Africa (7.6 of 8.9 million tons in 2021) (Food and Agriculture Organization of the United Nations 2022), where it is grown mainly for its protein-rich dry grain. However, other parts of the plant are edible including the leaves, stems, and immature pods (Boukar et al. 2019; Weng et al. 2019), which, in addition to culinary uses, are used in traditional medicine (Abebe and Alemayehu 2022). Cowpea haulms, stems, and leaves also constitute a nutritious fodder for livestock (Tarawali et al. 1997). In addition to its ability to grow in nutrient-poor soil, cowpea is also remarkable for having exceptional resilience to drought and high temperatures (Hall 2004), rendering it an ideal candidate for bolstering food system sustainability and for mitigating the repercussions of global climate change.

Linguistic and archaeological evidence suggest that cowpea was domesticated at least 5,000 years ago in Sub-Saharan Africa (Herniter et al. 2020), presumably from *V. unguiculata* ssp. *dekindtiana* var. *spontanea* (Pasquet and Padulosi 2012). The domestication region, however, remains a subject of debate with proposed domestication centers in West Africa (Vaillancourt and Weeden 1992; Padulosi and Ng 1997; Coulibaly et al. 2002; Ba et al. 2004), East Africa (Xiong et al. 2018), or in both regions (Huynh et al. 2013; Xiong et al. 2016; Herniter et al. 2020). Previous efforts to

study genetic variation in contemporary cowpea germplasm collections have indicated a strong pattern of genetic differentiation between germplasm from West and East Africa (Huynh et al. 2013; Xiong et al. 2016; Muñoz-Amatriaín et al. 2017; Fatokun et al. 2018; Herniter et al. 2020; Muñoz-Amatriaín et al. 2021), with West Africa as the region of maximum diversity of domesticated cowpea (Fatokun et al. 2018; Muñoz-Amatriaín et al. 2021).

Although genetic resources for cowpea are underdeveloped compared to cereal crops such as maize, rice, and wheat, cowpea benefits from several available sequenced genomes (Lonardi et al. 2019; Liang et al. 2023) as well as a high quality genotyping array (Muñoz-Amatriaín et al. 2017). Several quantitative trait locus (QTL) mapping populations have been developed to link genotype to phenotype in this species and have been useful in identifying QTL for diverse traits such as flowering time (Andargie et al. 2013; Huynh et al. 2018; Lo et al. 2018; Olatoye et al. 2019), disease resistance (Huynh et al. 2016; Santos et al. 2018), seed coat color and patterning (Herniter et al. 2018; Herniter et al. 2019), perenniability (Lo et al. 2020), and several domestication-related traits such as seed shattering and yield (Lo et al. 2018). Additionally, several genome-wide association studies (GWAS) have been conducted to discover single nucleotide polymorphisms (SNPs) associated with flowering time (Muñoz-Amatriaín et al. 2021; Paudel et al. 2021), disease resistance (Dong et al. 2022; Kpoviessi et al. 2022), seed size (Lo et al. 2019), seed protein content (Chen et al. 2023), drought response (Ravelombola et al. 2021; Wu et al. 2021), pod shattering (Lo et al. 2021), and salt tolerance (Ravelombola et al. 2022). Nevertheless, the majority of QTL mapping studies in cowpea have been limited in their ability to fine-map associated loci due to constrained sample sizes or low molecular marker densities.

Here, we report the genetic characterization of a core collection of cowpea held at the International Institute of Tropical Agriculture (IITA), a CGIAR research center in Ibadan, Nigeria. The IITA Cowpea Collection represents the most comprehensive collection of worldwide cowpea germplasm, currently consisting of 16,460 accessions collected from 91 countries (March 2022) (IITA 2023). The bulk of the collection (~12,000 lines) consists of locally adapted landraces (hereafter traditional cultivars). The core subset of the collection was selected to represent the diversity of geographic origin and agrobotanical traits present among accessions in the full collection (Mahalakshmi et al. 2007) and has been subsequently refined to include 2,082 accessions. To study genetic variation in the Core Collection, we made use of previously generated SNP data (Close et al. 2023) consisting of 2,021 accessions genotyped at nearly 50,000 polymorphic sites using the Illumina Cowpea iSelect Consortium Array (Muñoz-Amatriaín et al. 2017). We then leveraged this dataset to characterize the pattern of population structure in global cowpea and demonstrated the utility of this resource for GWAS by dissecting the genetic basis of variation in seed pigmentation.

## Materials and methods

### Accession collection sites

Collection sites for each accession were provided by the IITA (IITA 2023). World regions are described according to the United Nations M49 standard (United Nations Statistics Division).

### Genotyping and variant filtering

Accessions were genotyped using the Illumina Cowpea iSelect Consortium Array which assays nearly 50,000 SNPs (Muñoz-Amatriaín et al. 2017) as described in Close et al. (2023). All alleles reported in this work belong to the Watson strand.

SNPs with greater than 95% of missing calls across samples were filtered first, followed by filtering of samples with greater than 50% missing calls or more than 20% heterozygous calls across sites. Near identical lines were identified by calculating pairwise Hamming distance between samples with plink 1.9 (Chang et al. 2015), applying hierarchical clustering to the distance matrix, and defining clusters of accessions using a static tree cut at height of 500. One sample from each cluster was randomly selected to be retained for subsequent analysis.

From the filtered SNP set, a set of SNPs in approximate linkage equilibrium was produced by selecting SNPs with minor allele frequency greater than or equal to 0.01 and no more than 5% missing calls. The selected SNPs were then pruned pairwise in 50 SNP windows with a step size of 10 and an  $R^2$  threshold of 0.20. All SNP filtering and pruning was done using plink 1.9 (Chang et al. 2015). The pruned set, composed of 4,245 SNPs, was used for all population structure analyses.

### Variant effect prediction

The predicted effect of variants were determined using Ensembl Variant Effect Predictor v. 95.0 (McLaren et al. 2016) using the *V. unguiculata* v.1.0 reference genome assembly and v.1.2 of the genome annotation (Lonardi et al. 2019).

### Genetic diversity statistics

Allele frequencies and pairwise genetic distances were calculated using plink v.1.9 (Chang et al. 2015). Expected heterozygosity was calculated according to the following formula:  $\frac{1}{N} \sum_{n=1}^N (1 - p^2 + q^2)$ , where N is the total number of sites and p and q are the frequencies of the major and minor frequency alleles at each site, respectively.

An iterative sampling method was used to calculate the minimum number of accessions to represent 90 and 95% of the genetic diversity in the IITA Cowpea Core Collection. The method consisted of sampling an increasing number of accessions without replacement and computing the number of segregating sites and expected heterozygosity per sampled cohort. The procedure was conducted 1,000 times to approximate the minimum number of accessions, with accession sampling order randomized in each iteration.

### Population structure

Population structure was analyzed using ADMIXTURE 1.3.0 (Alexander et al. 2009; Alexander and Lange 2011) for  $K = 1$  to 10 with 10 replicates per K. The most likely K value was determined using the Evanno method (Evanno et al. 2005). ADMIXTURE runs were postprocessed and visualized using the pophelper 2.3.1 R package (Francis 2017). Samples were assigned to the group contributing 70% or more of the sample's estimated ancestry fraction. All other samples were considered to be unassigned.

Principal component analysis (PCA) on allele frequencies was done using the adegenet v. 2.1.5 (Jombart 2008) R package. The mean allele frequency at a locus was substituted for missing genotype calls.

### Phenotypic analyses

The IITA Cowpea Characterization and Cowpea Evaluation phenotypic datasets were downloaded from the IITA website (IITA 2023) and consisted of 15 quantitative and 23 qualitative traits. The data were collected as part of the IITA cowpea breeding program across multiple field trials spanning many years and thus represent reference values used for breeding. For the analysis, seed color pigmentation traits were defined according to

the majority color of the seed and similar traits were combined as described in [Supplementary Table 5](#). “Small eye” is synonymous with the “Eye 1” pattern and “Large eye” is synonymous with the Eye 2 pattern described by [Hermite et al. \(2019\)](#). “Self-colored” is described herein as the synonymous “full coat” pattern ([Singh 2020](#)). Qualitative traits were coded into binary “case/control” variables using the fastDummies 1.6.3 R package ([Kaplan 2020](#)) or considered to be discrete numerical variables ([Supplementary Table 6](#)). Differentiation in phenotype distribution between population structure clusters was tested using Mann–Whitney–Wilcoxon tests for quantitative phenotypes and Fisher’s exact test for qualitative phenotypes in R ([R Core Team 2023](#)). Two-sided tests were used for all analyses and a Bonferroni adjusted threshold was used to determine significance at  $\alpha = 0.05$ .

### Linkage disequilibrium

Pairwise linkage disequilibrium (LD) between markers was calculated with plink 1.9 ([Chang et al. 2015](#)). Since calculation of pairwise LD for all markers in a dataset can be computationally prohibitive, LD was only calculated between markers that were within 10 Mbp of each other. Genome-wide and chromosome-specific background levels of LD were determined by first calculating the mean  $R^2$  value in 500 bp distance bins and then taking the median value across all bins.

### Genome-wide association mapping

Genome-wide association mapping was done by fitting univariate linear mixed-effects models with GEMMA 0.98.4 ([Zhou and Stephens 2012](#)). The models included a centered relatedness matrix to correct for spurious associations due to population structure. Only sites with minor allele frequency (MAF) greater than 0.01 and missing in fewer than 5% of samples were tested in this analysis. Continuous quantitative phenotypes were log-transformed prior to analysis. Qualitative phenotypes were encoded in binary as 1 or 0 corresponding to presence or absence of the trait. Bonferroni correction was used to correct P-values for multiple testing and significance was assessed at  $\alpha = 0.05$ . Linked significant SNPs ( $R^2 > 0.20$ ) within 1 Mbp were considered to define a single peak and the SNP with the lowest P-value per peak was assigned as the “tag SNP.” Candidate genes were identified by examining the overlaps between significant peaks and loci described in the *Vigna unguiculata* v.1.2 genome annotation ([Lonardi et al. 2019](#)).

### Gene expression data

The *Vigna unguiculata* Gene Expression Atlas (VuGEA) data ([Yao et al. 2016](#)) were retrieved from the Legume Information System ([Berendzen et al. 2021](#)) at <https://data.legumeinfo.org>.

## Results

### Genetic diversity of the IITA cowpea core collection

Genomic DNA was extracted from 2,021 accessions from the IITA Cowpea Core Collection and scored at nearly 50,000 polymorphic sites using the Illumina Cowpea iSelect Consortium Array ([Close et al. 2023](#)). Following removal of sites and samples with missing data and excess heterozygosity, the dataset comprised genotypes from 1,991 accessions collected from 86 countries across 6 continents ([Fig. 1a](#)). All major cowpea producing regions were represented in the dataset with 1,547 accessions from Africa, 207 from Asia, 110 from North America, 68 from Latin America, 43 from Europe, and 14 from Oceania. The accessions mostly

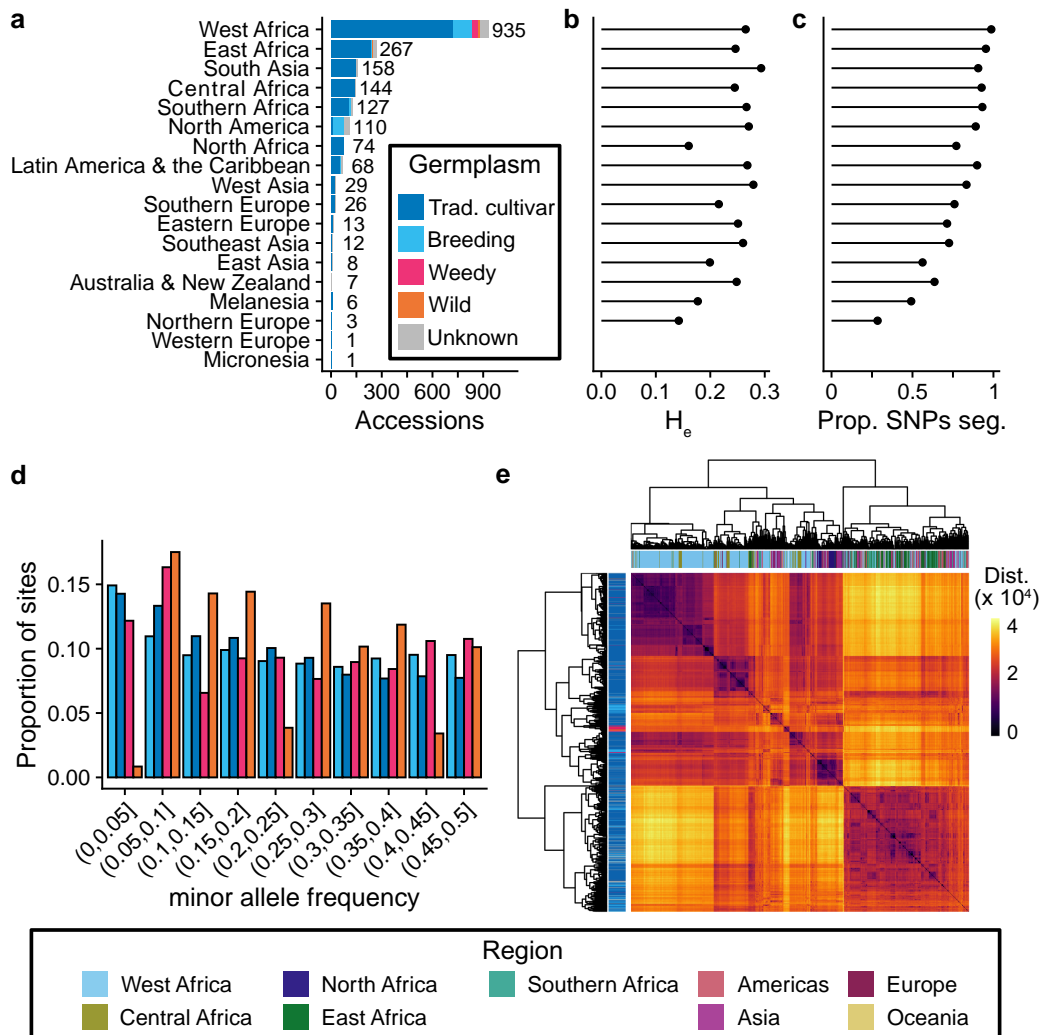
consisted of 1,590 traditional cultivars [including 3 accessions of yardlong bean (*V. unguiculata* ssp. *sesquipedalis*)], with an additional 206 breeding lines mostly from Nigeria and the United States of America. Also included were 57 uncultivated accessions, presumed to be wild or weedy.

Pairwise genetic distances calculated between accessions across filtered sites revealed many nearly identical accessions in the Core Collection ([Supplementary Fig. 1](#)). In total, we identified 130 clusters (comprising 400 accessions) of nearly identical accessions by applying hierarchical clustering to the genetic distance matrix ([Supplementary Table 1](#), static tree cut threshold of 500 corresponded to differences at  $\sim 0.5\%$  or fewer sites). The vast majority of clusters consisted of either two or three accessions (79 and 25 clusters, respectively), while 26 clusters consisted of 4 or more accessions (max cluster size 15). Most clusters (72/130) contained accessions collected from different countries with 40 clusters consisting of accessions from countries on different continents, 19 clusters consisting of accessions collected from adjacent countries within Africa, 12 clusters consisting of accessions from nonadjacent countries within Africa, and 1 cluster with accessions from adjacent countries in South America. The geographic distribution of accessions within clusters suggests that accessions were dispersed between continents by human activity and then resampled during the construction of the Core Collection. Of the 130 clusters, 86 consisted of accessions of the same germplasm status (e.g. traditional cultivars only), 20 clusters contained both traditional cultivars and breeding lines, 12 clusters had traditional cultivars and accessions of unknown status, 3 clusters contained breeding material and accessions of unknown status, and 2 clusters contained wild or weedy accessions and traditional cultivars. We randomly selected one accession from each of the 130 clusters for subsequent analysis.

The final dataset consisted of 1,722 accessions genotyped for 48,004 biallelic SNP markers. The included accessions reflect the geographic diversity of the IITA Cowpea Collection, with lines originating from 85 countries and comprising  $\sim 83\%$  (1,722/2,081) of accessions from the IITA Cowpea Core Collection ([Mahalakshmi et al. 2007](#)) and 86% (324/376) of accessions from the IITA Cowpea Mini-Core Collection ([Fatokun et al. 2018](#)). The analyzed samples encompassed 1,394 traditional cultivars, 171 breeding and research lines, 32 weedy accessions, 17 wild accessions, and 110 lines with unknown germplasm status. While the traditional cultivars constitute a global collection, the breeding and research material was predominately collected from Africa and North America (98 and 51 accessions, respectively). The presumed wild accessions were exclusively collected from Africa, while the presumed weedy accessions originated solely from West Africa.

Crop dispersal and improvement can lead to genome-wide depletion of genetic diversity. We sought to determine how these processes impacted cowpea by estimating average expected heterozygosity ( $H_e$ ) within the major geographic regions represented in our dataset.  $H_e$  ranged from 0.14 in Northern Europe to 0.29 in South Asia among regions that contributed 3 or more accessions ([Fig. 1b](#), [Supplementary Table 2](#)). Within Africa,  $H_e$  was similar for all regions ( $H_e = 0.25\text{--}0.27$ ) except for North Africa, which had a substantial reduction in diversity ( $H_e = 0.15$ ). Outside of Africa,  $H_e$  varied considerably, with higher values exceeding those observed in Africa in South Asia, West Asia, North America, and Latin America. In contrast, all regions of Europe and Oceania had reduced  $H_e$  compared to Sub-Saharan Africa.

Although  $H_e$  provides a summary of genetic variation within a population, the measure is influenced by the number of segregating alleles and their frequencies. The number of segregating sites per geographic region ranged from 13,660 in Northern Europe to



**Fig. 1.** Genetic diversity of the IITA Core Cowpea Collection. a) Region of origin and germplasm status for genotyped accessions. Labels beside each bar indicate the number of accessions per region. b) Average expected heterozygosity per region of origin calculated using all sites. c) Proportion of SNPs segregating per region. d) Folded allele frequency spectrum by germplasm status. Germplasm status labels are shared with (a). e) Heatmap of pairwise genetic distances. Row labels indicate germplasm status and column labels indicate region of origin.

47,361 in West Africa (Fig. 1c, Supplementary Table 2). The number of segregating sites per region was greatest in Sub-Saharan Africa, with a loss of over 10% of segregating sites observed in all other regions except for South Asia. The proportion of segregating sites was especially low in Oceania and East Asia, though these regions were not well sampled in the IITA core.

Taken together, our analyses are consistent with the hypothesis that West Africa is the center of domesticated cowpea genetic diversity with the largest number of segregating sites observed in this region. We observed a reduction of genetic diversity in all regions out of Africa except South Asia, which is consistent with founder effects upon dispersal. The elevation of  $H_e$  in South Asia is largely due to the majority of alleles segregating at intermediate frequencies (median MAF among segregating alleles = 0.25). The reduction of both  $H_e$  and the proportion of segregating sites in North Africa suggests strong founder effect with low genetic diversity among the founding lines. We also found evidence that cowpea improvement has reduced genetic diversity, with breeding lines having fewer segregating sites (46,332 vs 47,660) and an enrichment for both rare ( $MAF \leq 0.05$ ) and higher intermediate frequency ( $MAF > 0.30$ ) alleles compared to traditional cultivars (Fig. 1d).

To support cowpea breeding and improvement, we calculated “micro-population” subsets of accessions constituting 95 and 90% of the genetic diversity present in the IITA Cowpea Core Collection. The group representing 95% of genetic diversity included 36 accessions, while the group representing 90% of genetic diversity consisted of 19 accessions (Supplementary Table 1). Despite the Core Collection being composed of accessions that reflect the geographic and phenotypic diversity of the broader IITA Cowpea Collection, our findings indicate that the vast majority of common genetic diversity within the core subset can be represented with a minimal number of lines. However, as the genotyping approach used here focuses on alleles segregating at high frequencies in cowpea, it is important to note that the subsets are missing much of the rare variation present in the Core Collection.

## Patterns of population structure

After removal of nearly identical samples, we observed a multi-modal distribution of pairwise genetic distances between accessions suggesting that strong population structure might exist in the Core Collection. Hierarchical clustering of accessions by

genetic distance linked population structure to geographic origin and germplasm status (Fig. 1e). The primary genetic pattern in the dataset consisted of two large clusters: one cluster composed predominately of accessions from Eastern and Southern Africa and a second cluster encompassing accessions mostly from Northern, Central, and Western Africa. Germplasm collected from outside Africa is interspersed within the two main clusters, consistent with the aforementioned pattern of migration out of Africa likely facilitated by human activity. We also observed that many wild and weedy accessions clustered together within the larger cluster. It remains uncertain whether this observation is a result of ascertainment bias of our genotyping approach or if accessions designated as wild or weedy are actually feral accessions that have escaped cultivation.

To examine population structure in more detail, we next estimated the ancestries of each accession using ADMIXTURE (Alexander et al. 2009; Alexander and Lange 2011) (Fig. 2a). Evanno's  $\Delta K$  method (Evanno et al. 2005) suggested an optimum of two subpopulation clusters (Supplementary Fig. 2). We assigned accessions to the respective cluster that contributed 70% or more ancestry, resulting in 614 accessions in cluster 1, 680 accessions in cluster 2, and 428 accessions unassigned to a cluster (Supplementary Table 1). Accessions were nonrandomly distributed between clusters for 13 of the 17 geographic regions (Chi-squared tests,  $P < 0.05$ , Supplementary Table 3). Cluster 1 contains the majority of accessions from East Africa (90%), Latin America and the Caribbean (75%), North America (71%), Southern Africa (72%), and Australia and New Zealand (67%). In contrast, cluster 2 consists of the majority of accessions from North Africa (92%), West Africa (59%), and Central Africa (52%). The majority of lines from East Asia (86%), Melanesia (83%), and Southeast Asia (82%) were unassigned to a cluster. Germplasm from Eastern Europe, Micronesia, Southern Europe, and West Asia did not have significant enrichment for belonging to a particular cluster. Given the strong partitioning of accessions by geographic subregion within Africa, we hereafter refer to cluster 1 as the ESA (Eastern and Southern Africa) cluster and cluster 2 as the WNCA (West, North, and Central Africa) cluster. The two clusters have significant genetic differentiation ( $F_{st} = 0.292$ ) and the ESA cluster has a slightly higher level of genetic diversity compared to the WNCA cluster ( $H_e = 0.25$  for ESA, 0.22 for WNCA, 0.28 for unassigned).

To explore the possibility of population substructure in West Africa, we considered additional subpopulations. Considering  $K = 3$  subpopulations reassigned 124 accessions from the WNCA cluster and 82 previously unassigned accessions to the new cluster 3. Additionally, 78 and 146 accessions from the ESA and WNCA clusters, respectively, were unassigned at  $K = 3$ . Cluster 3 consisted of nearly all accessions from North Africa in proximity to the Nile River Delta (57/62) and the majority from Southern Europe (20/25), East Asia (5/7), and Eastern Europe (6/10) (Fig. 2a). We also observed that the ancestry previously attributed to the WNCA cluster in accessions found outside of Africa was reassigned to cluster 3. Cluster 3 accessions were nearly exclusively collected in arid climates (Köppen climate classification: BWh/BWk), while other clusters are found to be mostly confined to wet tropical climates (Supplementary Fig. 3), hinting at the possibility that cluster 3 ancestry includes alleles adapted to drier environments.

Modeling population structure with an additional subpopulation ( $K = 4$ ) grouped 148 accessions previously unassigned to a cluster when considering fewer subpopulations to cluster 4 (Fig. 2a and d). Cluster 4 consisted of 133 accessions from West

Africa, 7 accessions from either Central or East Africa, and 8 accessions collected in other world regions. The West African cluster 4 accessions were not associated with a particular landform such as a river or mountain range and, like the majority of accessions in the WNCA cluster, originated from tropical savanna climates (Supplementary Fig. 3). This group of accessions likely represents early-flowering traditional varieties that were cultivated alongside later maturing lines as a precaution against drought (Mortimore et al. 1997).

A PCA of allele frequencies was largely consistent with the population structure modeling. The first 10 PCs explain 27% of the variance in the dataset, with nearly half of this variance (12.8%) explained by the first 2 PCs alone (Supplementary Fig. 4). Overall, the distribution of individuals along the first 2 PCs was continuous, suggesting frequent admixture between population groups (Fig. 2b-d). PC 1 separates the ESA cluster from the WNCA cluster (Fig. 2b-d), while PC 2 separates cluster 3 from the other clusters (Fig. 2c). Cluster 4 is not resolved along the first 2 PCs and is instead resolved on PC 3 (Fig. 2d, Supplementary Fig. 5).

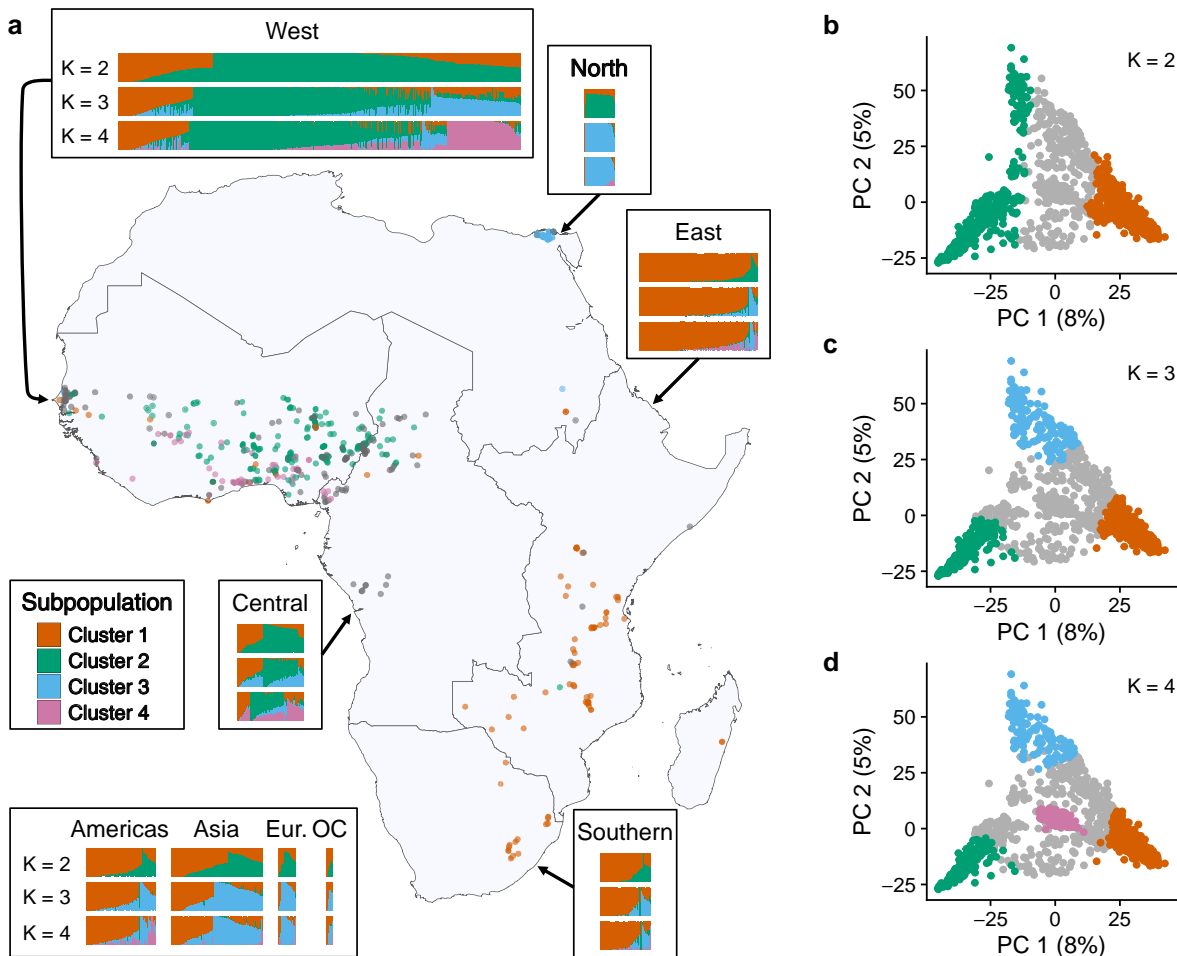
ESA ancestry was also overrepresented in the modern breeding material present in the IITA core though a large proportion of accessions remained unassigned (Supplementary Table 4), perhaps the result of controlled crosses between accessions from the two clusters. Consistently, increasing  $K$  did not reduce the number of unassigned accessions, pointing to true admixture rather than cryptic population structure in the sample. In addition, we found limited evidence for contribution from clusters 3 and 4 in the breeding germplasm. Two of the three yardlong bean accessions were assigned to the ESA cluster while the third accession remained unassigned to a subpopulation cluster.

Overall, our population structure analyses confirm that global cowpea is characterized by genetic differentiation between germplasm from Western vs Eastern and Southern Africa (Huynh et al. 2013; Xiong et al. 2016; Muñoz-Amatriaín et al. 2017; Fatokun et al. 2018; Herniter et al. 2020; Muñoz-Amatriaín et al. 2021). Further, our results suggest that germplasm derived from Northern, Eastern, and Southern but not Western Africa represent the majority of ancestry in lines distributed to other parts of the world. Breeding material appears to be enriched for ancestry from the ESA cluster, potentially neglecting climate-adapted alleles from other clusters.

## Phenotypic distributions of subpopulations

Our analyses of population structure in global cowpea indicate that cowpea germplasm is broadly genetically differentiated into two germplasm groups defined by geography (i.e. the ESA and WNCA clusters). To understand the degree to which population structure is linked to phenotype, we leveraged the IITA Cowpea Characterization and Cowpea Evaluation datasets (IITA 2023) to compare the distributions of 34 quantitative phenotypes and the frequencies of 37 qualitative phenotypes between the two subpopulation clusters (Materials and Methods, Supplementary Tables 5 and 6). As a caveat, the phenotyping data we analyze here was collected through IITA's cowpea breeding program through numerous field trials conducted across both time (1967–present) and location. Therefore, the data represent reference phenotype values used for breeding efforts and do not correspond to values measured as part of a single experiment.

We identified 17 quantitative phenotypes with significantly different distributions between the WNCA and ESA subpopulation clusters (Wilcoxon test, Bonferroni-corrected  $P < 0.05$ , Fig. 3a, Supplementary Table 7). Overall, the majority of differences were found in life history and morphological traits and not in



**Fig. 2.** Population structure in global cowpea germplasm. a) Inferred ancestry for all genotyped individuals from the core collection and map of collection sites for 545 traditional cultivars with known collection sites in Africa. Accessions are grouped by subregion according to the United Nations geoscheme and each accession with a known collection site is colored according to subpopulation assignment when modeling  $K = 4$  subpopulations. Accompanying barplots specify the proportions of ancestry attributed to each subpopulation per individual when modeling  $K = 2$ ,  $K = 3$ , and  $K = 4$  subpopulations, respectively. Eur. = Europe, OC = Oceania. b-d) Principal components analysis of allele frequencies with accessions colored by subpopulation for  $K = 2-4$ .

disease susceptibility traits. Accessions in the ESA cluster generally matured earlier (Fig. 3b), were more branched (Fig. 3c), had larger lateral organs (Fig. 3d and e), and produced more seeds that were generally smaller in size compared to the WNCA cluster (Fig. 3f). Additionally, accessions in the ESA cluster were more likely to have smooth testa texture, raceme in all layers of the canopy, intermediate or semi-erect growth habit, higher twining tendency, and higher susceptibility to anthracnose compared to the WNCA cluster (Supplementary Fig. 6). Conversely, accessions in the WNCA cluster were more likely to have rough testa texture, raceme in the upper leaf layers only, prostrate or semi-prostrate growth habit, low twining tendency, and higher resistance to anthracnose.

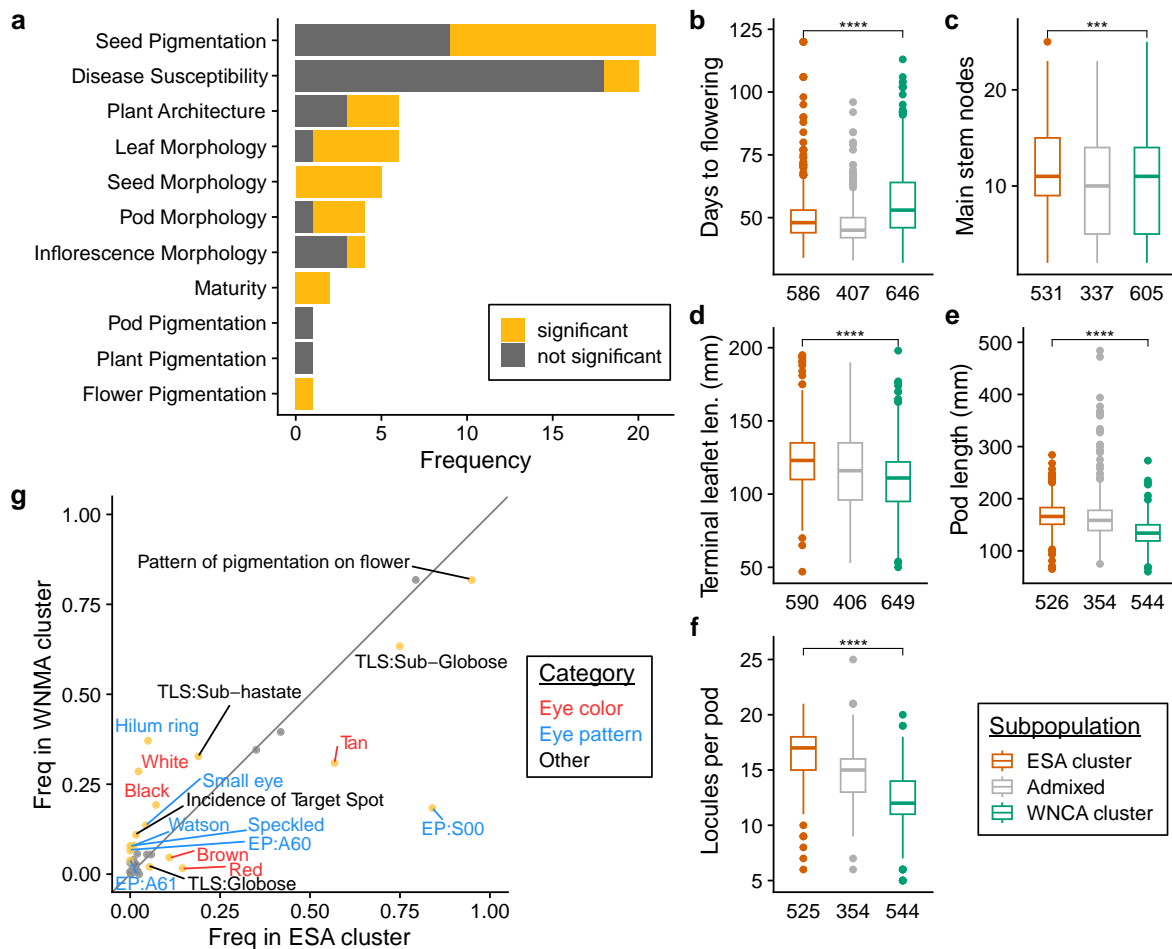
We also observed differences in the incidence of qualitative traits between the ESA and WNCA subpopulation clusters. In total, there were 17 traits with differential frequency between the two clusters (Fisher's exact test, Bonferroni-corrected  $P < 0.05$ , Fig. 2g, Supplementary Table 8). Many of the phenotypes with significant differences were seed coat pigmentation color and patterning traits. For instance, black and white eye colors and the hilum ring, small eye, Watson, and speckled pigmentation patterns were more common in the WNCA cluster compared to the ESA cluster (see below for a detailed description of these patterns).

In contrast, the brown, red, and tan eye colors were more common in the ESA cluster. Additional phenotypes with differential frequencies between the two clusters included several leaflet shapes, with globose and sub-globose leaflet shapes more common in ESA and the sub-hastate shape more common in the WNCA cluster. We also observed that flower pigmentation is more common in the ESA cluster and incidence of target leaf spot was more common in the WNCA cluster.

Taken together, the data suggest that ESA and WNCA cowpea are differentiated across a wide suite of traits spanning plant architecture, pigmentation, and life history.

### Decay of LD in the cowpea genome

To establish the IITA Core Collection as a valuable resource for identifying the genetic basis of agronomically relevant traits using GWAS in cowpea, we examined the patterns of genome-wide and chromosome-specific LD between SNP markers. Overall, we found that LD decays to  $R^2 = 0.20$  by 121.5 Kbp on average, with full decay to the genome-wide background by 2.76 Mbp (Fig. 4a). We also observed differences in the rate of LD decay between the ESA and WNCA subpopulation clusters, with more rapid decay in the WNCA cluster ( $R^2 = 0.20$  by 136 and 89 Kbp, genome-wide background by 3.57 Mbp and 2.97 Mbp, respectively).



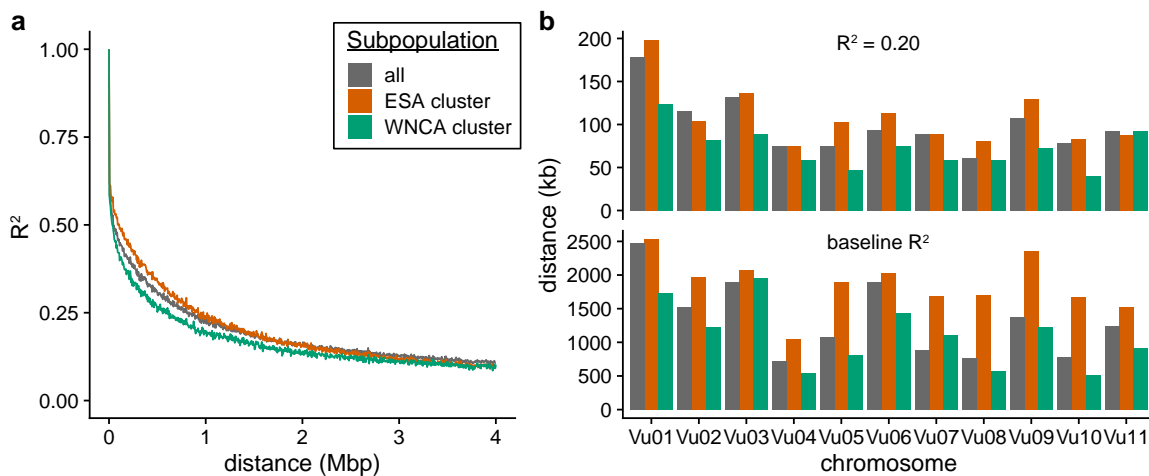
**Fig. 3.** Phenotypic distributions in subpopulation clusters. a) Number of phenotypes by category, colored by the number of phenotypes with statistically significant difference between the ESA and WNCA clusters. b-f) Tukey's boxplots of representative phenotype distributions by subpopulation cluster. Numbers below each boxplot denote N. Additional phenotypes are presented in Supplementary Fig. 6. g) Qualitative trait frequencies in the two subpopulation clusters. Grey reference line indicates equal frequency in both clusters. Points are colored according to whether there is a significant difference in trait frequency (golden rod) between the two subpopulation clusters. Labels correspond to phenotype names and are colored by category. EP, eye pattern; TLS, terminal leaf shape.

The rate of LD decay also varied between chromosomes. LD decay to  $R^2 = 0.20$  ranged from 68 Kbp on Vu08 to 178 Kbp on Vu01 and decay to chromosome-specific background levels of LD ranged from 721 Kbp on Vu04 to 2.48 Mbp on Vu01 (Fig. 4b, Supplementary Fig. 7). As expected given the genome-wide data, the rate of LD decay was more rapid across nearly all chromosomes in the WNCA cluster compared to the ESA cluster.

The resolution of genetic mapping in GWAS is determined by the recombination rate, which determines LD decay, and the density of genetic markers near an associated variant. In regions with dense markers and high recombination (i.e. rapid LD decay), there is a higher likelihood of pinpointing an associated locus within a narrow genomic window. Using our measures of LD decay, we estimate that an associated locus will contain ~25 genes and ~60 SNPs on the array on average. This estimate is based on the average distance of 5,611 bp between adjacent genes, an average gene length of 3,843 bp, and average distance of 3,997 bp between adjacent SNPs as measured in the reference genome (Muñoz-Amatrián et al. 2017; Lonardi et al. 2019). However, we acknowledge that these estimates should be considered to be coarse due to the nonuniform distribution of genes and SNPs and variation in the local recombination rate within the genome.

## The genetic architecture of seed pigmentation patterning

To demonstrate the utility of the genotyped IITA Core Collection to map the genetic basis of agronomically important traits, we focused on studying the genetic basis of seed pigmentation. Variation in the color and pattern of cowpea seed pigmentation is a particularly conspicuous example of the rich phenotypic diversity of the IITA collection. Cowpea seeds range from having no pigmentation (called “no color”) to complete pigmentation throughout the seed coat (so-called “full coat”). Between these extremes, various pigmentation patterns exist, traditionally classified qualitatively. The “small eye” or “Eye 1” pattern displays pigmentation in a narrow margin around the seed hilum with an indistinct margin, while the “large eye” or “Eye 2” pattern displays pigmentation encompassing the hilum with a wide, distinct margin and covering a significant portion of the ventral surface (Herniter et al. 2019). The “Watson” pattern, named after a corresponding variety, features pigmentation surrounding the hilum and extending to the micropylar end of the seed, with an unclear boundary (Spillman 1911). Similarly, the “Holstein” pattern, also named after a variety possessing the pattern, exhibits solid pigmentation over most of the seed coat, with patches of color on



**Fig. 4.** LD in the cowpea genome. a) Mean pairwise LD between SNPs in 500 bp windows. b) LD per chromosome. Minimum distance to  $R^2$  of 0.20 (top) and to chromosome-specific background level of LD (bottom).

the remaining portion of the seed. In addition to pigmentation patterning, seed coat color exhibits extraordinary variation across a spectrum of colors including white/cream, tan/buff, brown, red, maroon, and purple/black (Harland 1919; Harland 1920; Spillman and Sando 1930; Saunders 1960; Drabo et al. 1988).

Seed coat pigmentation patterning in cowpea has been described as being controlled by a three locus system consisting of the color factor (*C* locus), Watson factor (*W* locus), and Holstein factor (*H* locus) (Spillman and Sando 1930; Saunders 1960; Drabo et al. 1988). The *C* locus is epistatic to the other two loci and determines if pigmentation is present throughout the seed coat or just restricted to the eye. The *W* and *H* loci meanwhile act as modifiers and determine whether the pattern of pigmentation in the seed coat develops into the Watson, Holstein, or full coat patterns. Previous work based on QTL mapping in biparental populations has proposed *Vigun07g110700* as a candidate gene for the *C* locus, *Vigun09g13900* for the *W* locus, and *Vigun10g163900* for the *H* locus (Herniter et al. 2019).

We first sought to confirm the previously proposed candidate loci for the *W*, *H*, and *C* factors using genome-wide association study in the IITA Core Collection. Our mapping results for pigmentation patterning yielded 3 distinct associated loci: 1 locus on Vu09 and 2 loci on Vu10 (Fig. 5, Supplementary Table 9). The locus on Vu09 was associated with all 5 pigmentation patterns, spanned 1.38 Mbp (Vu09:30,035,583–31,416,843), and contained 116 genes (Fig. 5f). We identified 2\_01960 as the tag SNP for this locus and found that the small eye, large eye, and Holstein pigmentation patterns were associated with the A allele (*MAF* = 0.146) at this locus while the Watson and full coat patterns were associated with the major frequency G allele. Examination of the genes in this significant interval found that *Vigun09g13900*, a WD-repeat gene proposed as a candidate gene for the *W* locus (Herniter et al. 2019), is present in the region between the tag SNP and a linked upstream significant SNP. Furthermore, the candidate gene is surrounded by other significant variants that are not linked with the tag SNP, suggesting the presence of additional associated haplotypes segregating in this region.

The first locus on Vu10 was associated with the small eye, Watson, and full coat pigmentation patterns. The tag SNP for this locus is 2\_31919, with the C allele (*MAF* = 0.221) associated with the small eye and Watson patterns and the major frequency T allele associated only with the full coat pattern. The locus spans

12 KB (Vu10:38,454,820–38,467,161) and overlaps with *Vigun10g165400*, an R2–R3 MYB transcription factor (Fig. 5g). R2–R3 MYB genes are reported to have a role in transcriptional regulation of flavonoid biosynthesis as part of MYB–bHLH–WDR complexes (Xu et al. 2015).

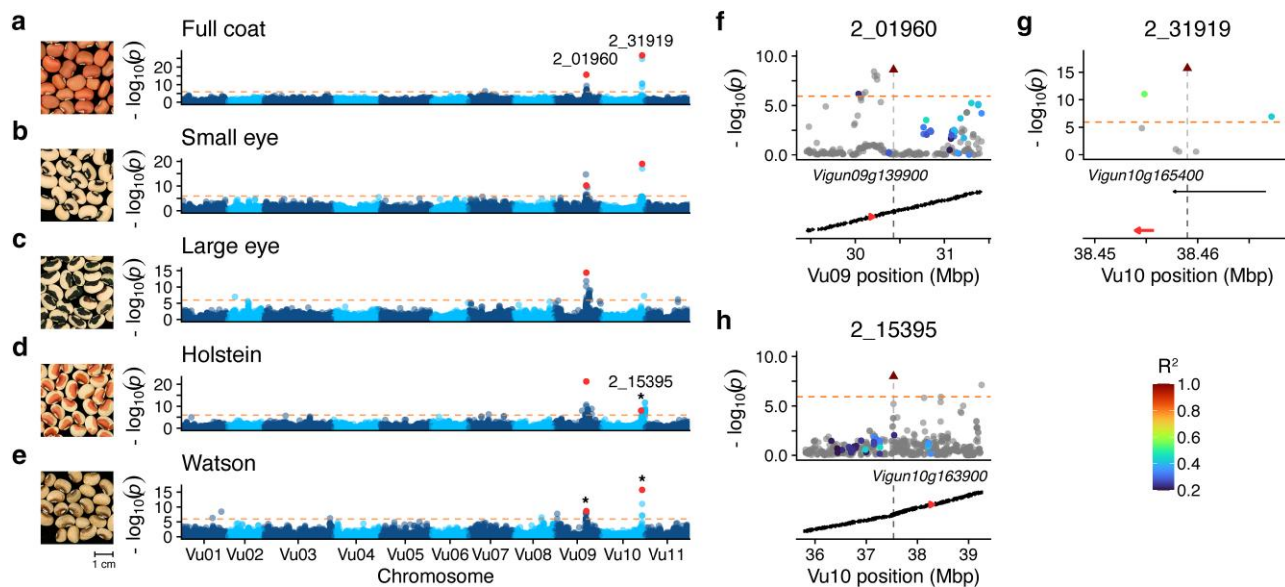
The GWAS for the Holstein pattern yielded 2 peaks on Vu10 that are several megabases apart. The first peak has tag SNP 2\_15395, which is moderately linked ( $R^2$  = 0.346) to 2\_23640, a SNP that is ~52 KB upstream of *Vigun10g163900*, an E3 ubiquitin ligase previously proposed as a candidate gene for the *H* locus (Herniter et al. 2019) (Fig. 5h). There are also several SNPs surrounding the gene that did not reach significance per the conservative Bonferroni threshold but are close to the threshold and would have likely been significant with increased power. The second peak spans 189 KB (Vu10:41,100,865–41,290,623) and contains 23 genes but does not yield a clear candidate gene.

Our results confirm that previously reported candidate genes for the *W* locus and *H* locus could be mapped with the Core Collection GWAS panel. Although we were unable to map the *C* locus on Vu07 in the five main seed pigmentation patterns, the locus was mapped in the GWAS for hilum ring and eye absent patterns (Supplementary Fig. 8).

### The genetic architecture of seed color

In cowpea, seed coat color varies on a continuous spectrum with several colors typically described qualitatively such as white/cream, tan/buff, brown, red, blue, purple, and black. Previous genetic studies of the segregation of seed coat color phenotypes revealed that coat color is inherited independently of pigmentation pattern and is likely controlled by at least five unlinked major genes (Saunders 1960; Drabo et al. 1988). To date, only the locus controlling black seed color, the *Bl* locus, has been identified (Herniter et al. 2018). To attempt to identify the loci associated with additional seed coat colors, we mapped the genetic basis of the black, red, tan, blue, and white seed coat colors using GWAS in the Core Collection (Fig. 6). In total, we found 4 major peaks associated with seed coat color: 1 each on Vu03 (red/tan), Vu05 (black/tan), Vu06 (blue), and Vu10 (No color/white).

We first sought to confirm if the Vu05 locus previously proposed to be associated with black seed coat color (Herniter et al. 2018) could be mapped via GWAS in the Core Collection. The GWAS for black seed coat yielded a major 121 KB peak on Vu05



**Fig. 5.** The genetic basis of seed coat pigmentation patterning. a–e) Manhattan plots from genome-wide association mapping of seed coat pigmentation pattern phenotypes. Orange dotted line indicates Bonferroni-corrected significance threshold. Tag SNPs are highlighted in red and labeled. Tag SNP 2\_15395 is unique to Holstein, highlighted SNP on Vu10 in Watson and Full coat correspond to 2\_31919. Photographs from UCR Cowpea Minicore as follows: Full Coat/Self-colored (TVu-3842/MinicoreUCR\_247), Small eye/Eye 1 (TVu-16278/MinicoreUCR\_197), Large eye/Eye 2 (TVu-8775/MinicoreUCR\_318), Holstein (TVu-7625/MinicoreUCR\_290), Watson (TVu-2933/MinicoreUCR\_232). f–h) Local linkage disequilibrium for tag SNPs indicated by \* in a–e. Candidate genes are highlighted in red and labeled.

(Vu05:3,104,538–3,225,877), spanning 13 genes (Fig. 6a and f). The locus was also associated with the tan seed coat color (Fig. 6c). The minor C allele at tag SNP 2\_19309 (MAF = 0.463) was associated with the black seed coat phenotype while the major T allele was associated with the tan seed coat phenotype. Among the genes in the region, we found Vigun05g039500, a R2–R3 MYB transcription factor previously identified as a candidate gene for the Bl locus, which is associated with black seed coat and purple pod tip (Herniter et al. 2018).

The peak on Vu03 is associated with both the red and tan seed colors (Fig. 6b, c, and g), spans 274 KB (Vu03:10,933,603–11,208,404) and contains 25 genes. The T allele (MAF = 0.102) at the tag SNP 2\_20787 is associated with red seed color, while the major frequency G allele is associated with the tan color. The locus maps to a pair of anthocyanin reductase (ANR) genes: Vigun03g118700/Vigun03g118800, with the tag SNP a predicted missense mutation in the former gene. ANR acts in the biosynthesis of flavan-3-ols and proanthocyanidins (condensed tannins) in plants (Xie et al. 2003). Mutations in ANR in *Arabidopsis thaliana* have been shown to result in the accumulation of anthocyanins in the seed coat (Devic et al. 1999), making this locus a compelling candidate for the red seed color phenotype.

The GWAS for blue seed color resulted in a ~155 KB peak on Vu06 (Vu06:13,491,073–13,645,944) that spans 5 genes (Fig. 6d). SNP 2\_22603, which is strongly linked to the tag SNP and lies directly upstream, is a predicted synonymous or intron mutation within the gene Vigun06g030900 (Fig. 6h). This gene is a homolog of AT3G59030 (TRANSPARENT TESTA12) which has been shown to be required for sequestration of flavonoids in the seed coat epithelium tissue of *A. thaliana* (Debeaujon et al. 2001).

Our attempt to map the genetic basis of white/cream seed color yielded one of the peaks on Vu10 (2\_31919) observed in seed coat patterning GWAS (Fig. 5). Additionally, we did not observe any significant SNPs in the GWA mapping of brown seed coat color, likely due to the fact that this phenotype can be heat induced (Pottoroff et al. 2014). In total, our GWAS was able to map a previously identified

locus for the black seed coat color and propose novel loci associated with the red, tan, and blue seed coat colors.

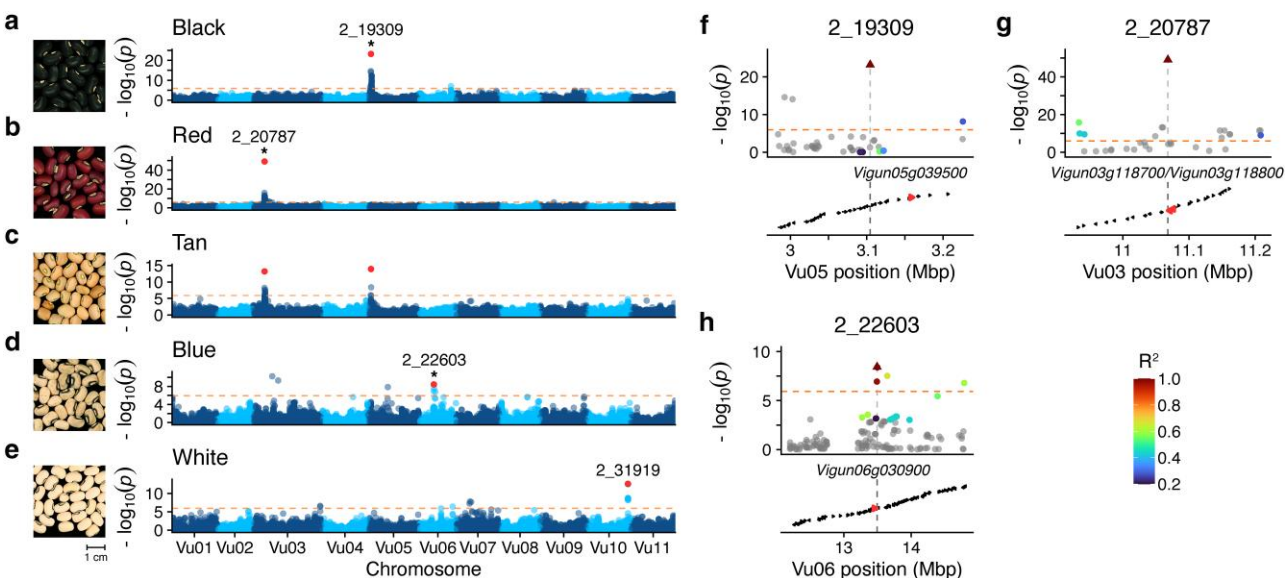
## Mapping of additional phenotypes

Including the seed coat pigmentation pattern and color phenotypes discussed previously, we attempted to map the genetic basis of 71 distinct phenotypes in total. Altogether, 42 of the GWAS had one or more significant SNPs (range 1–106 SNPs, median 8, Supplementary Table 9). Of the 748 SNPs that were associated with variation in a phenotype, 698 were associated with variation in only one phenotype and 50 SNPs were associated with more than one phenotype. P-values and minor allele effects for all GWAS are presented in Supplementary Tables 10 and 11, respectively.

We found numerous interesting association peaks with clear underlying candidate genes. For instance, GWAS for presence of flower pigmentation had a significant SNP adjacent to Vigun07g148800, an E3 ubiquitin ligase. Although E3 ubiquitin-ligases are implicated in numerous biological processes, they have been reported to be negative regulators of the flavonoid biosynthesis pathway transcriptional activation complex (Shin et al. 2015; Herniter et al. 2019). We also found candidate genes associated with disease resistance. GWAS for incidence of rust yielded 2 peaks with clear candidates: a significant SNP on Vu02 in a 12-oxophytodienoate reductase (Vigun02g192500) involved in the Jasmonic acid biosynthesis pathway and a very strong peak on Vu09 spanning 90 KB (Vu09:2,156,441–2,246,970) containing 3 TIR-NBS-LRR family disease resistance genes with substantial sequence similarity: Vigun09g027450, Vigun09g027600 (DSC1), and Vigun09g027700 (suppressor of npr1-1) (Supplementary Fig. 9). Overall, our analyses demonstrate the utility of the genotyped IITA Cowpea Core Collection for discovering variants associated with phenotypic variation in cowpea.

## Discussion

The purpose of this study was to utilize the recently reported SNP genotyping data to examine population structure and the



**Fig. 6.** The genetic basis of seed coat color. a–e) Manhattan plots from genome-wide association mapping of seed color phenotypes. Orange dotted line indicates Bonferroni-corrected significance threshold. Tag SNPs are highlighted in red and labeled. Photographs from UCR Cowpea Minicore as follows: black (TVu-13968/MinicoreUCR\_139), red (TVu-15426/MinicoreUCR\_173), tan (TVu-1036/MinicoreUCR\_108), blue (TVu-4669/MinicoreUCR\_262), and white (TVu-14401/MinicoreUCR\_153). f–h) Local linkage disequilibrium for tag SNPs (triangle) for peaks indicated by \* in a–e. Candidate genes are highlighted in red and labeled.

potential for genome-wide association mapping in the IITA Cowpea Core Collection. The accessions in this collection are representative of the genotypic and phenotypic diversity of extant cultivated cowpea germplasm. To our knowledge, the dataset constitutes the largest cohort of cowpea that have been genotyped genome-wide with a single technology, and we anticipate that the resources linked to this work will be of considerable utility to the cowpea research and breeding community.

### Genetic relationships in global cowpea

Our analysis of population structure revealed that global cowpea is broadly genetically differentiated into two major subpopulations defined by geography, with separation between germplasm collected in Eastern and Southern Africa (ESA cluster) from germplasm collected in West, Central, and North Africa (WNCA cluster). The genetic differentiation between germplasm from West Africa and East Africa has been observed previously in smaller subsets of globally collected cowpea germplasm (Huynh et al. 2013; Xiong et al. 2016; Muñoz-Amatriaín et al. 2017; Fatokun et al. 2018; Herniter et al. 2020; Muñoz-Amatriaín et al. 2021), and our observations confirm that this is the major organizing principle of cowpea diversity. However, we did observe that between 9 and 12% (depending on the number of subpopulations considered) of accessions collected in West Africa were assigned to the ESA cluster (cluster 1). While this pattern is consistent with the proposed center of domestication in West Africa and potentially reflects the historical migration of a subset of West African germplasm to the South and East, it could also be explained by the modern migration of germplasm from East to West Africa. Therefore, we were unable to rule out the possibility of a separate domestication center in East Africa given our data. Examining the genetic diversity of wild cowpea populations across West and East Africa could provide valuable insight into the mode and tempo of domestication of this crop.

Although the IITA Cowpea Core Collection primarily consists of traditional cultivars from West Africa, the population structure

analyses did provide some insight into the dispersal of cowpea germplasm around the globe. The majority of germplasm from Latin America and the Caribbean, North America, and Oceania clustered within the ESA cluster, suggesting dispersal to these locales from Eastern and Southern Africa. In contrast, germplasm collected from East Asia, Melanesia, and Southeast Asia was mostly assigned to cluster 3 and thus likely represents migrants dispersed out of North Africa, perhaps through the Sabaean Lane, an ancient trade route connecting East Africa with modern-day India (Murdock 1959; Herniter et al. 2020). Further, our analyses indicated a limited role of cluster 2 (West Africa specific) ancestry in the dispersal of cowpea out of Africa, with only a few accessions, all present in Latin America and the Caribbean, assigned to this cluster when considering more than 2 subpopulations. Our analysis of phenotypic differences between the ESA and WNCA clusters indicates that ESA germplasm is associated with earlier flowering, suggesting that this phenotype was important for the worldwide dispersal of cowpea.

Compared to other surveys of global population structure in cowpea, our analysis yielded fewer “optimal” subpopulation clusters. We attribute this result to our use of the Evanno method (Evanno et al. 2005) to decide the number of subpopulations, which has been shown previously to be biased to select  $K = 2$  (Janes et al. 2017). Therefore, we considered up to 2 additional subpopulations, yielding a cluster of accessions collected from the Nile Delta in Northern Egypt in close proximity to the Mediterranean Sea and a cluster of early-flowering accessions in West Africa, both of which were also discovered in the UCR Minicore, a mini-core subset of a cowpea diversity collection held at the University of California, Riverside (Muñoz-Amatriaín et al. 2021). In contrast to previous studies, we did not observe subpopulation clusters consisting of only breeding lines or clusters consisting solely of germplasm collected outside of Africa. It is possible that these signals are also present in the data and would have been resolved if we had improved sampling from continents beside Africa, if we had considered additional subpopulations, or if

we had conducted the analysis omitting the traditional cultivars from West Africa that constitute the bulk of the collection. Our analysis also did not identify the cluster of yardlong bean (*ssp. sesquipedalis*) previously identified in other collections of global germplasm (Muñoz-Amatriaín et al. 2021) as the IITA Cowpea Core Collection only has 3 accessions known to be from this group.

In addition to the patterns of population structure, we found that a substantial number of accessions are nearly identical to one or more other accessions in the Core Collection. Our analysis indicated that the majority of clusters of identical accessions consisted of lines collected across geopolitical boundaries, with many clusters containing collections transcending continents. A primary objective when selecting accessions for the Core Collection was to ensure that the chosen lines were representative of the geographic and phenotypic variation present in the full IITA Cowpea Collection (Mahalakshmi et al. 2007). Consequently, the occurrence of dispersed identical lines within the Core Collection may be a result of sampling across geographic gradients (i.e. sampling an accession that had been dispersed across a geopolitical border in both the original and dispersed location in an effort to have representatives from all regions where cowpea is found) or due to sampling of lines subject to phenotypic plasticity upon selection of the core collection subset.

### A genome-wide association mapping panel for cowpea

A primary objective of this study was to determine whether the IITA Cowpea Core Collection constitutes a suitable panel for genome-wide association mapping in domesticated cowpea. The rate of decay of LD dictates the precision with which significant associations can be linked to specific candidate genes. The decay of LD in the cowpea core collection is slow when compared to other plant systems (1.5 KB in maize (Remington et al. 2001) and 10 KB in *A. thaliana* (Kim et al. 2007)), which is consistent with observations in smaller previously genotyped collections (Muñoz-Amatriaín et al. 2021; Sodedji et al. 2021). The slow decay of LD may be a feature common to legumes as similar rates have been reported in cultivated soybean (Zhou et al. 2015). However, the slow decay of LD in cowpea is partly due to strong population structure, as LD decays somewhat more quickly within the WNCA population cluster compared to the ESA cluster.

Despite the slow decay of LD in cowpea, we were able to use genome-wide association to identify hundreds of loci associated with variation in dozens of traits. We note that the phenotype values used for our analyses were collected across many field studies spanning years and locations, leaving us unable to control for the contribution of environmental variance to phenotypic variance. We anticipate that GWAS of phenotypes collected in a single experiment will have even greater success in identifying the genetic basis of trait variation using the IITA Cowpea Core Collection.

### The genetic basis of seed coat pigmentation in cowpea

In the present study, we focused on mapping the genetic basis of several seed pigmentation patterning and color phenotypes using GWAS in the IITA Cowpea Core Collection. Seed pigmentation traits are ideal for genetic study due to their high heritability and Mendelian inheritance pattern (Harland 1919, 1920). These traits are also important for consumer preference, with regional variation in preferred patterns or colors driven by perceptions of seed quality and intended culinary use (Langyintuo et al. 2003, 2004; Boukar et al. 2019). For instance, seeds with black eyes are highly desirable in Ghana but are undesirable in Cameroon

(Langyintuo et al. 2004). Moreover, brown or red seed coat color is preferred over white seed coat color in some regions of Nigeria (Langyintuo et al. 2003). As such, identifying markers associated with seed pigmentation traits is crucial for cowpea breeding and improvement to align with regional consumer preferences and market demands.

Our efforts to map the genetic basis of seed pigmentation patterning using GWAS yielded several candidate loci that were previously discovered using QTL mapping in F2 and advanced RIL populations (Herniter et al. 2019). The consistency between the loci mapped as part of our work and previous reports suggests that the loci proposed previously likely regulate seed pigmentation pattern more generally in global cowpea and support the three locus model for the genetic control of seed pigmentation patterning (Spillman and Sando 1930; Saunders 1960; Drabo et al. 1988).

Through GWAS, we were able to map a second locus on Vu10 that likely contributes to seed pigmentation patterning (Fig. 5g). For this locus, we propose Vigun10g165400, a R2-R3 MYB transcription factor, as a candidate gene. Analysis of patterns of LD at this locus suggests that this locus is distinct from the candidate gene at the *H* locus despite being in close proximity (~110 KB downstream). This locus has been observed in a previous attempt to map the genetic basis of seed pigmentation patterning (Herniter et al. 2019) but was disqualified due to not being observed in the associated haplotype blocks of all the tested F2 populations. Our results support the hypothesis that multiple genes in close proximity on Vu10 likely regulate seed pigmentation patterning. Further work is needed to determine the molecular basis of this trait and to demonstrate the efficacy of the proposed candidates.

Seed coat color has been studied for over 100 years in cowpea (Harland 1919; Harland 1920) and the existence of at least 5 genes affecting this trait have been proposed (Saunders 1960; Drabo et al. 1988). However, the full genetic control of seed coat color has yet to be elucidated. Here, we propose that the *R* locus controlling red seed color is an ANR, either Vigun03g118700 or Vigun03g118800. Although both genes lie under the associated peak, the tag SNP is a predicted missense variant in Vigun03g118700, which is also expressed at a high level during early seed development in the reference cultivar IT97K-499-35, which has a black-eyed phenotype. In contrast, Vigun03g118800 is expressed at a low level across many tissues and developmental timepoints (Supplementary Fig. 10) (Yao et al. 2016). Thus, we consider Vigun03g118700 to be the likely causal locus for the red seed coat color phenotype. This candidate gene was also independently proposed as regulating red seed coat color based on the results of QTL mapping in a biparental F2 population and GWAS in the UCR minicore population (Herniter et al. 2024).

ANR is active in the flavonoid biosynthesis pathway, where it competes with 3-O-glucosyltransferase to convert cyanidin into condensed tannins or anthocyanins, respectively (Kovinich, Saleem, Arnason, et al. 2012; Kovinich, Saleem, Rintou, et al. 2012). Mutations in ANR have been shown to result in the rapid accumulation of anthocyanins during early seed development in *A. thaliana* (Devic et al. 1999) and knockdown of ANR in soybean yields a red seed coat phenotype (Kovinich, Saleem, Arnason, et al. 2012; Kovinich, Saleem, Rintou, et al. 2012). Our results suggest that a nonfunctional ANR gene is responsible for the red seed coat phenotype in cowpea.

We also propose Vigun06g030900, a homolog of TRANSPARENT TESTA12 (TT12), as a candidate gene for the blue seed color phenotype. TT12 is a MATE transporter which mediates anthocyanin deposition in *A. thaliana* seeds (Debeaujon et al. 2001; Marinova et al.

**Table 1.** Genetic model of seed pigmentation pattern and color. Proposed genotypes at pigmentation patterning and color quantitative trait loci. QTL correspond to SNPs as follows: Pat9.1 = 2\_01960, Pat10.1 = 2\_31919, Pat10.2 = 2\_15395, Col3.1 = 2\_20787, Col5.1 = 2\_19309, Col6.1 = 2\_22603.

Phenotype	Genotype at patterning loci	Genotype at color loci
Full coat	Pat9.1 <sup>g</sup> , Pat10.1 <sup>t</sup> , Pat10.2 <sup>c</sup>	
Small eye (Eye 1)	Pat9.1 <sup>a</sup> , Pat10.1 <sup>t</sup> , Pat10.2 <sup>t</sup> or Pat10.2 <sup>c</sup>	
Large eye (Eye 2)	Pat9.1 <sup>a</sup> , Pat10.1 <sup>t</sup> , Pat10.2 <sup>c</sup>	
Holstein	Pat9.1 <sup>a</sup> , Pat10.1 <sup>t</sup> , Pat10.2 <sup>c</sup>	
Watson	Pat9.1 <sup>g</sup> , Pat10.1 <sup>c</sup> , Pat10.2 <sup>t</sup> or Pat10.2 <sup>c</sup>	
Black		Col3.1 <sup>g</sup> , Col5.1 <sup>c</sup> , Col6.1 <sup>t</sup>
Red		Col3.1 <sup>t</sup> , Col5.1 <sup>c</sup> or Col5.1 <sup>t</sup> , Col6.1 <sup>t</sup>
Tan		Col3.1 <sup>g</sup> , Col5.1 <sup>t</sup> , Col6.1 <sup>t</sup>
Blue		Col3.1 <sup>g</sup> , Col5.1 <sup>c</sup> , Col6.1 <sup>c</sup>
White	Pat10.1 <sup>c</sup>	Col3.1 <sup>g</sup> , Col5.1 <sup>c</sup> or Col5.1 <sup>t</sup> , Col6.1 <sup>t</sup>

2007). It remains to be seen whether blue seed color might be driven by changes in activity or regulation of *Vigun06g030900*.

A full genetic model of seed pigmentation patterning and color in cowpea based on our GWAS is presented in Table 1.

## Future directions

The genotyped IITA Cowpea Core Collection represents a valuable tool for dissecting the genetic basis of agronomic traits in cowpea. We anticipate that the resources produced by this work will be useful for marker-assisted breeding and will facilitate the production of new cultivars in response to consumer preference and changing environmental conditions. To facilitate the latter aim, this collection could be used to identify climate-associated alleles, as many of the traditional cultivars have known collection locations. This knowledge could be especially useful for ensuring the resilience of cowpea into the future.

## Data availability

The genotyping data has been published previously on Dryad (<https://doi.org/10.6086/D19Q37>). Information about the IITA Cowpea Collection, including phenotyping data, is available at <https://my.iita.org/accession2/collection.jspx? id=1>. All other data necessary for reproduction of this work is present within the manuscript, supplemental figures, and supplemental tables.

Supplemental material available at G3 online.

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## Conflicts of interest

The author(s) declare no conflict of interest.

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