Population genomic and genome-wide association studies of agroclimatic traits in sorghum


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Accelerating crop improvement in sorghum, a staple food for people in semiarid regions across the developing world, is key to ensuring global food security in the context of climate change. To facilitate gene discovery and molecular breeding in sorghum, we have characterized ∼265,000 single nucleotide polymorphisms (SNPs) in 971 worldwide accessions that have adapted to diverse agroclimatic conditions. Using this genome-wide SNP map, we have characterized population structure with respect to geographic origin and morphological type and identified patterns of ancient crop diffusion to diverse agroclimatic regions across Africa and Asia. To better understand the genomic patterns of diversification in sorghum, we quantified variation in nucleotide diversity, linkage disequilibrium, and recombination rates across the genome. Analyzing nucleotide diversity in landraces, we find evidence of selective sweeps around starch metabolism genes, whereas in landrace-derived introgression lines, we find introgressions around known height and maturity loci. To identify additional loci underlying variation in major agroclimatic traits, we performed genome-wide association studies (GWAS) on plant height components and inflorescence architecture. GWAS maps several classical loci for plant height, candidate genes for inflorescence architecture. Finally, we trace the independent spread of multiple haplotypes carrying alleles for short stature or long inflorescence branches. This genome-wide map of SNP variation in sorghum provides a basis for crop improvement through marker-assisted breeding and genomic selection.

Sorghum bicolor | quantitative trait locus | adaptation

Agricultural production and food security in the developing world face numerous threats, particularly in semiarid regions, which are acutely vulnerable to climate change (1). Sorghum [Sorghum bicolor (L.) Moench.] is an important crop species for farmers in semiarid and arid regions because relative to other cereal crops it can sustain high yields where precipitation is low or erratic. Thus, sorghum has become the major cereal crop in semiarid regions and a dietary staple for more than 500 million people, predominantly in sub-Saharan Africa and south Asia (2). Worldwide, sorghum is grown for food (grain and syrup), animal feed, fiber, and fuel in both subsistence and commercial agriculture systems. Because rising temperatures and reduced precipitation due to climate change make some areas unsuitable for maize and rice production, the importance of drought-tolerant crops like sorghum is likely to increase (1). Current breeding priorities in sorghum seek to mitigate climate-dependent stressors, both abiotic (e.g., drought and acid soils) and biotic (e.g., insect pests and fungal diseases) (2). To meet the projected doubling of global food demand over the next few decades in the context of global change, the pace of crop improvement must be accelerated (3).

Sorghum has a wide range of adaptation, and traditional varieties from across Africa and Asia provide a rich source of morphological and physiological traits for crop improvement (4–6). The primary domestication of sorghum occurred near present-day Sudan approximately 10,000 y ago, and diffusion occurred to diverse climates across Africa, India, the Middle East, and east Asia between 8,000 and 1,500 y ago (7). Because of this ancient origin and diffusion, adaptation to local climates and cultural practices is reflected in morphological and physiological variation among and within the five major types (races) of domesticated sorghum (8). For instance, in parts of West Africa where rainy periods are long and erratic, open panicle guinea types are preferred to reduce grain mold and insect damage. Conversely, in parts of South and East Africa where rainy seasons are relatively short and predictable, dense panicle kafir and durra types are preferred to increase grain yield per plant (2). Further natural and human selection has occurred in the United States over the past ∼150 y as temperate and tropical germplasm from Africa and Asia has been adapted for use in combine-harvested commercial agriculture (9).

Genomic analysis of diverse populations is increasingly being used to uncover the genetic basis of complex traits, including agroclimatic traits of crop species. Genome-wide single nucleotide polymorphism (SNP) scans of population genetic parameters in crops have been used to identify loci under selection (10, 11) and dissect quantitative traits (11). In addition, genome-wide association studies (GWAS) have been used to elucidate the genetic basis of agronomic traits in rice (12) and maize (10). Nucleotide diversity scans (13, 14) and association studies (5, 15) have been carried out in sorghum, but the resolution and sensitivity of these studies has been limited by the small number of markers (14). Thus, compared with maize and rice, less is known about the genetic basis of agronomic traits in sorghum. Among the four classical dwarfing loci that have been studied in sorghum for more than 70 y (9), only one has been cloned (Dw3/SbPGP1) (16). Recently, it has become feasible to genotype thousands of markers rapidly and at low cost through the application of barcode multiplexing and high-throughput sequencing (17). To better understand the diversity of sorghum, facilitate the genetic dissection of agroclimatic traits, and accelerate marker-assisted breeding, we characterized 971 sorghum accessions at 265,487 SNPs by using genotyping-by-sequencing (GBS). Here, we describe a genome-wide map of SNP variation, trace patterns of crop diffusion to diverse agroclimatic regions, and use GWAS to identify genes underlying natural variation in agroclimatic traits.
Results and Discussion

Genome-Wide Map of SNP Variation. To represent the genetic, geographic, and morphological diversity of sorghum, we used 971 accessions from the world germplasm collections, combining three previously defined sorghum diversity panels (Dataset S1) (4–6). The majority of these accessions consist of source-identified landraces or traditional cultivars from across Africa and Asia (Fig. 1A). Of these accessions, 238 are landrace-derived sorghum conversion lines, in which alleles for short stature and early maturity were introgressed into tropical landraces to facilitate the use of tropical germplasm in temperate breeding programs (18). The remainder consists of wild/weedy relatives or elite lines and breeding materials, many of which have unknown geographic origin and/or mixed ancestry. For each accession, we constructed ApeKI-reduced representation libraries and generated a total ∼21 Gbp of sequence on the Illumina Genome AnalyzerHi/HiSeq by using GBS (Dataset S2) (17). In total, 6.13 million unique 64-bp tags were identified across all sorghum accessions. Eighty-five percent of these tags aligned to the reference sorghum genome (19), and 384,561 putative SNPs were identified. After filtering for local linkage disequilibrium and tag coverage (>10% of taxa), 265,487 SNPs were retained, with an average density of one SNP per 2.7 kbp. Of 27,412 annotated genes in the reference sorghum genome, 72% were tagged by a SNP within the gene and 99% were tagged by a SNP within 10 kb. Importantly, this genome-wide map of SNP variation is of sufficient resolution for GWAS in sorghum, given >100,000 SNPs are estimated to be required (14). Additionally, because of simultaneous SNP discovery and genotyping, this sequencing-based SNP map will have little ascertainment bias and greater power for mapping studies (20).

Linkage Disequilibrium and Recombination Rates. Characterizing patterns of linkage disequilibrium (LD) is critical for the design of association studies (21, 22), interpretation of association peaks (12), and the transfer of alleles in marker-assisted selection (23). To characterize the mapping resolution for genome scans and GWAS, we quantified the average extent of LD decay and localized patterns of LD for each chromosome (Table S1 and Fig. S1). On average, LD decays to 50% of its initial value by 1 kb and to background levels ($r^2 < 0.1$) within 150 kb. These LD decay estimates are higher than previously published values in sorghum of 15–20 kb (24) and 50–100 kb (14). This difference may be attributed to low genome coverage of markers and fewer genotypes in previous studies. Because sorghum is a predominantly selfing species, but readily outcrosses, we expect a greater extent of LD than in out-crossing species (25). Accordingly, the extent of LD in sorghum is similar to that in rice (∼75–150 kb) (22), another self-pollinated crop, but much greater than in maize (∼2 kb) (26), which is an outcrosser. Sliding window (1 Mb) estimates of pairwise LD show that telomeric regions have lower LD than centromeric regions (Fig. S1B). This pattern is likely due to higher historical recombination rates in telomeric regions compared with centromeric regions (Fig. 2C). The average recombination rate in sorghum (1.4 $\rho$/kb) is intermediate relative to recent estimates in plants such as Arabidopsis (0.8 $\rho$/kb) (21) and maize (2.2 $\rho$/kb) (26). Based on these results, we expect mapping resolution to range widely across the genome, from single-gene resolution in some telomeric regions to megabase-level resolution near the centromeres.

Population Structure and Geographic Differentiation. To understand the geographic structuring of genetic diversity, we contrasted genome relatedness among 971 sorghum accessions to the stated location of origin and morphological descriptors in the worldwide germplasm database. The resulting neighbor-joining trees (Fig. 1B) and Bayesian clustering analysis (Fig. S2 and Dataset S3) show population structuring along both morphological type and geographic origin, confirming previous analyses (4, 14) and providing additional insights into the fine-scale patterns of ancestry resulting from crop diffusion. Of the five morphological types, the kafir sorghums that predominate in southern Africa show the strongest pattern of population subdivision relative to other races (Fig. 1B and Fig. S2). Durra type sorghums, found in warm semi-arid or warm desert climates of the Horn of Africa, Sahel, Arabian peninsula, and west central India, form a distinct cluster that is further

![Fig. 1.](https://www.pnas.org/cgi/doi/10.1073/pnas.1215985110 Morris et al.)
differentiated according to geographic origin. Bicolor types are not notably clustered, except those from China (known as kaoliang), which forms a distinct subgroup and shows genetic similarity to durra types, particularly those from Yemen. Caudatum types, which are primarily found in tropical savanna climates of central Africa, are diverse and show only modest clustering according to geographic distribution. Finally, guinea types, which are widely distributed in tropical savanna climates, show five distinct subgroups, four of which cluster according their geographic origin (far west Africa, west Africa, eastern Africa, and India). A fifth guinea subgroup, which includes guinea margaritiferum types, forms a separate cluster along with wild genotypes from western Africa (Fig. 1B) and may represent an independent domestication (4).

The structure of sorghum populations provides insight into historical processes of crop diffusion within and across agroclimatic zones of Africa and Asia. Diffusion across agroclimatic zones is expected to be rare relative to diffusion within agroclimatic zones (27). Indeed, the patterns of relatedness among sorghum populations suggest that agroclimatic constraints have been at least as important as geographic isolation in shaping the diffusion process (Fig. S3). Among the four phylogenetically supported sorghum types (kafr, durra, guinea, and caudatum), there is the least population structure among caudatum types, which range primarily in the ancestral region of domestication or adjacent areas with similar climate (Fig. L4). The one geographically structured subpopulation of caudatum is the latitudinal-diffused subpopulation from highland areas in east Africa. Although durra types diffused widely across Africa and Asia, they are restricted to regions with semiarid and desert climates (Fig. S3). This diffusion included kaoliang sorghums, which are likely derived from durra populations of the Middle East, but are found in cold semiarid regions of northern China (Fig. 1 and Figs. S2 and S3). Similarly, although guinea types have diffused over long distances, from western Africa to southeastern Africa and eastern India, they remain restricted to tropical savanna climates. Interestingly, Bayesian clustering analysis suggest that the temperate/subtropical-adapted kafr type is derived from (or at least shares ancestry with) guinea types of east African populations ($k = 3$ through $k = 6$). In this case, the kafr type may represent major phenotypic divergence and genetic bottlenecks resulting from a shift to a contrasting agroclimatic zone.

Genomic Patterns of Nucleotide Variation. To investigate the genomic signatures of domestication and diversification in sorghum, we quantified genome-wide nucleotide variation across sorghum landraces. Overall, average nucleotide diversity ($\pi$) was 0.00037/kb, $\theta = 0.00017$/kb, and Tajima’s $D$ value of 3.6. A scan of expected heterozygosity ($H_e$) values across the genome revealed many megabase-scale regions of low heterozygosity including a ∼40-Mb region of reduced nucleotide variation around the centromere of chromosome 7 (Fig. 2A). Of six starch-related genes previously studied as a priori candidates for domestication loci (15, 28), two are found at regions with low heterozygosity (Fig. 2A). The starch biosynthesis enzyme brittle endosperm 2 (b2) gene, which has been shown to be a likely domestication locus in maize (28) and sorghum (15), is at the base of the low heterozygosity region on chromosome 7. The size of this low diversity region and extensive LD (Fig. S1) may be due to low recombination rates in this pericentromeric region (Fig. 2C) or the presence of additional loci under selection. Another a priori domestication candidate from starch metabolic pathways, transcription factor opaque2 (15), is found at the base of a low heterozygosity region on chromosome 2. Lastly, another recently identified domestication locus, the shattering gene Sh1, is found at the edge of a region with moderately, but not strikingly, lower heterozygosity, consistent with the observation that non-shattering alleles are found on at least three haplotypes in domesticated sorghums (29). The large footprints of selection we observe here (up to several megabases) are consistent with the predominance of inbreeding in sorghum. Selective sweeps in out-crossing maize left smaller footprints (<100 kb) (30) than in self-pollinating rice (250 kb to 1 Mb) (31).

In sorghum conversion lines that carry introgressions of early maturity and short stature alleles, we also observed major reduction in heterozygosity in several genomic regions (Fig. 2B). These regions colocalize with previously mapped height [Dw2 (32), Dw3 (16), and Dw1/SbH1T9.1 (33)] and maturity loci [Ma1/SbPRR37 (34)] that are recessive in the introgression donor B73x406. In contrast, another classical maturity locus, Ma2/phyB (35), which is wild type in B73x406 and therefore was not under selection during the conversion process, shows no such reduction in heterozygosity. On chromosome 6, the low heterozygosity region extends from approximately 6.6 Mb to 42 Mb (the Ma1/Dw2 locus), suggesting that another height or maturity locus may be localized at 6.6 Mb (SI Results and Discussion). As was seen in the landraces, we find that...
large LD blocks result when selection occurs in low recombination regions around the centromere (Fig. 2 B and C).

GWAS. The genome-wide map of SNP variation we generated permits the dissection of complex traits in sorghum by using GWAS. To elucidate the genetic basis of plant height in sorghum, we determined associations between SNPs and plant height components by using data from 336 lines in the sorghum association panel (SAP; Fig. 3 and Fig. S4) (5, 33). Plant height is an important component for many agroclimatic traits such as competitive growth with weeds, resistance to lodging, and, in the case of temperate-adapted grain sorghums, the efficiency of combine harvest (2). Because this panel incorporates a large fraction of sorghum conversion lines with introgressions of dwarfing alleles, we know that much of the variation for height in this panel has a common genetic basis. We identified SNPs associated with total plant height, and two height components: preflag height, which quantifies elongation in the lower portion of the stem, and flag-to-apex length, which quantifies elongation in the upper portion of the stem (Fig. S4). The Dw3/SbPGP1 gene provides a positive control for GWAS (16). It is known that the reduced height of dw3 mutants is due to reduced elongation of lower internodes, therefore we considered preflag leaf height as a measure of lower internode elongation (33).

The third most significant association peak for preflag height is found at the dw3 locus [within 12 kb for general linear model (GLM) and 22 kb for compressed mixed linear model (CMLM)]. We also refined the mapping location of dw1 and dw2 and identified a possible location of dw4 (SI Results and Discussion).

Because the SAP includes a large fraction of converted lines, with large introgressions around height and maturity loci, the previous analysis does not reflect a typical GWAS case. To validate the broader applicability of GWAS in sorghum, we also sought to dissect a trait that was not a target of selection in the sorghum conversion program. Inflorescence architecture is a major agroclimatic trait that, in part, defines the major morphological types in sorghum (8). Moreover, because the genetic basis of inflorescence architecture is well-studied in maize, rice, and Arabidopsis, there are many a priori candidate genes that can be considered to evaluate the mapping approach. Indeed, several of the significant association peaks for inflorescence branch length were located in or near a priori candidate genes for inflorescence architecture, which are homologous to known maize, rice, or Arabidopsis floral regulators (Fig. 4 A and Table S2). For example, two peaks are in, and one is near (47 kb), C2H2 zinc finger transcription factors homologous to the classical maize floral development gene INDETERMINATE 1 (ID1) (Fig. 4 A and Table S2) (36).

Another association peak was found in a sorghum ortholog of Arabidopsis UNUSUAL FLORAL ORGAN (UFO) and rice ABERRANT PANICLE ORGANIZATION 1 (APO1). In rice, apo1 mutants exhibit small panicles and fewer branches (37).

In sorghum, strong population structure among the morphological types presents a challenge for mapping the genetic basis of the inflorescence architecture and other population-associated traits. Although statistical controls for population structure have proven effective here, the effects of population structure can be better addressed by the experimental design of mapping popu-
lizations, using regional mapping (20) or nested-association mapping (NAM) (38) approaches. Because of the use of sorghum conversion lines, the SAP captures some aspects of the NAM approach. The introgressions reduce the confounding effects of maturity differences in diverse germplasm (5) and, for height and maturity loci, improve mapping power by increasing the frequency of rare alleles (33). However, because the introgressions originate from the same donor line (BTx406), the large blocks of linked non-causative variation reduces the resolution of the association analysis (Fig. 2B). Also, the low diversity around height and maturity loci on chromosomes 6, 7, and 9 may prevent the mapping of QTL for other traits in these regions, especially on chromosome 6 where most of the chromosome has been introgressed in sorghum conversion lines (Fig. 2B). In some cases, therefore, mapping without converted lines will be more effective for association studies.

**Geographic Distribution of Haplotypes.** To gain further insight into the origin and spread of haplotypes linked to agronomic traits, we characterized the geographic distribution for SNPs of interest in 330 source-identified landraces that are independent of the lines used for GWAS. As expected given the role of *Ma1/SlBPRR37* in temperate zone adaptation (34), a previously identified functional variant (K162N) is near fixation in high-latitude kafir accessions from southern Africa and rare (<5%) elsewhere (Fig. S5). The three major height QTL identified by genome scan and GWAS, *dw2*, *dw3*, and *SlHT9.1*, have distinct allelic distributions across Africa and Asia (Fig. 3E–G). One of the alleles at the SNP closest to the putative *SlHT9.1* causative gene (*GA2-oxidase*) (39) is found at high frequency (>90%) in East African durra, Indian durra, and Chinese accessions and low frequency (<10%) in all other accessions (Fig. 3G). Likewise, one allele at the *Dw2* QTL peak is common in northeast Africa and Asia and rare elsewhere (Fig. 3E). This pattern is consistent with a genetic basis for semidwarfsim in East African and Asian sorghums, conferred by at least two causative polymorphisms, and originating from ancestral East African durra populations. Taken together, the evidence suggests that the classical dwarving alleles were likely selected from standing variation in durra (*dw1, dw2, dw4* and kafir (*dw3*) landraces (SI Results and Discussion).

The geographic distribution of alleles at inflorescence branch-length QTL also reveals evidence of the independent spread of multiple alleles controlling branch length (Fig. 4B–D). In general, the minor allele associated with longer branches is found at high frequency in West African guinea populations, and in a number of cases it also is found in other geographically distant populations. Interestingly, none of the alleles at top branch length association peaks were restricted to durra accessions (Table S2 and Fig. 4B–D), suggesting that we were able to identify QTL for the long-branch phenotype in guinea types, but not QTL for the short-branch phenotype in durra types. This result may be attributed to stronger population structuring of durra populations, compared with guinea, which can confound mapping of traits (20). Given the statistical correction for population structure, we did not map QTL underlying the branch length differences among major morphological types, rather we mapped QTL for branch length segregating within the morphological types, which are globally rare but locally common (10) (Table S2 and Fig. 4B–D).

**Conclusion**

A better understanding of genetic diversity in sorghum will support in situ conservation efforts, enhance the use of germplasm collections, and guide ongoing collection efforts (40). This genome-wide map of SNP variation will accelerate molecular breeding by expanding the diversity of germplasm accessible to crop improvement programs and increasing the resolution of GWAS, marker-assisted selection, and genomic selection (23, 41). By facilitating crop improvement in locally adapted and locally improved cultivars, genomic analysis of diverse crop germplasm can play an important role in supporting sustainable agriculture in Africa, Asia, and semiarid regions worldwide.

**Materials and Methods**

**Plant Materials.** Diverse sorghum germplasm from worldwide collections were used, combining three diversity panels: the US sorghum association panel (SAP) (5), the sorghum mini core collection (MCC) (6) and the Generation Challenge Program sorghum reference set (RS) (www.icrisat.org/
what-do-we-do/crops/sorghum/Sorghum_Reference.html. We were able to obtain appropriate plant material for 971 accessions (Dataset 5). The SAP was obtained from GRIN (www.ars-grin.gov). Country of origin, and approximate latitude and longitude for source-identified accessions, were obtained from the SINGER crop germplasm database (www.genesys-pgr.org).

**Genotyping by Sequencing.** DNA from MCC and RS lines (5–6 plants per accession) was isolated from 12-d-old seedlings by using the CTAB protocol (42). SAP DNAs were isolated by using DNeasy Plant Mini Kit (Qiagen). Genomic DNAs were digested individually with ApeKI (recognition site: G[T/A]GC) and 96- or 384-plex GBS libraries were constructed (Dataset 52) (17). DNA sequencing was performed either on the Illumina Genome Analyzer IIx or HiSeq2000. Sequences were mapped to the BTx623 sorghum reference genome (19) by using BWA (43), and SNPs were called with the TASSEL 3.0.9 GBS pipeline (www.maizegenetics.net/tassel). Sequence tags, 64-bp sequences that included a leading 4-bp C[T/A]GC signature from the cut site, were identified, and tags with at least 10× total coverage were retained. Missing data were imputed with NPUTE (44).

**Population Genetic Analysis.** Hierarchical population structure was estimated by using the ADMIXTURE program (45), a model-based estimation of ancestry in unrelated individuals using maximum-likelihood method. The neighbor-joining trees were built and heterozygosity calculated by using the ape package in R (46). Pairwise LD was calculated (r2) separately for each chromosome by using TASSEL 3.0 (47). Recombination rates were inferred by using Bayesian reversible-jump MCMC under the cross-over model of rhomap in LDhat (48) with 1 million iterations and 1 million burn-ins. Rates were estimated separately for each subgroup identified by ADMIXTURE at k = 10. To avoid confounding effects of shared introgressions, SAP lines were not included in recombination and LD analyses.

**GWAS.** Published phenotypes for plant height components and inflorescence branch length for the SAP were used for GWAS (33). GWAS was carried out in Genomic Association and Prediction Integrated Tool (49) by using a (i) GLM; or, to control for population structure, (ii) CMLM with population parameters previously determined (50) with the first three principal components as covariates. Bonferroni correction was used to identify significant associations.

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Supporting Information

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SI Results and Discussion

Genome Scan and GWAS on Plant Height. Because *dw3* mutants are known to show increased upper stem elongation, we additionally mapped *dw3* based on associations with flag leaf to apex distance (flag-to-apex). In this case, the top association peak is found near the *dw3* locus (114 kb from peak; \( P < 10^{-8} \)). The strongest association peak for plant height traits (Fig. S4) is a narrow peak on chromosome 9 around 57.2 Mb, which colocalizes with previously described plant height locus *dw1*/*ShH91* (1). The top association for plant height is 29 kb from a GA2-oxidase, a catabolic enzyme in the gibberellin pathway, which has been proposed as the gene underlying the plant height QTL *ShH91.1/Dw1* (2). Overexpression of GA2ox in rice leads to semidwarf phenotypes (3).

The second most significant peak maps to chromosome 6 between 39.7 Mb and 42.6 Mb near the classical dwarfing locus *Dw2* (4, 5). *Dw2* has been mapped adjacent to *Ma1* on chromosome 6 (6), to a region of ~100 kb around 42.2 Mb, but the gene underlying this QTL has not been cloned. The association peak for total plant height and preflag leaf height maps to a histone deacetylase (*SH06g015420*), which is homologous to well-studied global transcriptional regulators in plants (*hda*) (7, 8). In maize and Arabidopsis, down-regulation of closely related histone deacetylases (*hda101* and *AihHD1*, respectively) results in reduced plant height and a variety of changes in inflorescence architecture (7, 8). In rice, overexpression of *OshDAC1* increases plant height (9), whereas the knockdown of many genes in the *OshDAC* gene family lead to semidwarf phenotypes (10). Therefore, we propose that *dw2* phenotype is a result of loss of function in a sorghum histone deacetylase.

The fourth classical dwarfing locus in sorghum, *dw4*, has not been genetically mapped but is known to be linked to the other dwarfing loci (11). Based on the location of the next most significant peak in the height GWAS and heterozygosity scan, a potential physical position of the *Dw4* locus is at ~6.6 Mb on chromosome 6 (Fig. 3B and Fig. S4). Note, because of the simultaneous progression of maturity and dwarfing alleles in conversion lines, it may not be possible to distinguish between them using genome-wide scans, and additional loci may have been involved.

Origin of Dwarfing Alleles. Did the mutations that underlie classical dwarfing alleles arise de novo in the early US grain sorghums, as is suggested in classical breeding literature (12), or were they recruited from standing variation present in African or Asian landraces? The haplotypes associated with dwarfism at *dw1*/*ShH91.1*, *dw2*, and *dw3* are widely distributed among African and Asian landraces (Fig. 3), but these haplotypes could represent ancestral haplotypes on which new dwarfing mutations occurred. The classical literature, however, confirms that dwarfing alleles were already present in African landraces at the time that dwarf alleles were being adopted in US grain sorghums. For instance, dwarf durra varieties collected near Khartoum, Sudan, c. 1920 carry the *dw4* allele (*Gahan dura*) or both *dw1* and *dw4* (hegari) (11).

GWAS on Inflorescence Branch Length. CMLM GWAS on inflorescence branch length identified candidate genes homologous from several known regulators of inflorescence development or cell elongation (Table S2). These genes include *LEUNIG* (13), *The-seus1* (14), *Short panicle1* (15), Lost meristems (16), Dwarf in light (17), Dwarf8 (18), Tesintete brachidi (19), GIGANTEA (20), Clavata1 (21), Bearded ear1 (22), Indeterminate1 (23), Gibberellin dependent dwarf1 (24), and Aesthetic panicle organization1 (25). Compared with plant height components, mapping of inflorescence branch length QTLD depended more on methods for controlling population structure (Fig. S4). The top association peak for branch length (whether population structure is controlled) is a SNP found in another ID1 homolog (Table S2). The minor allele at this SNP is restricted to the three broomcorn varieties in the panel, which display the most extreme branch length phenotypes, with inflorescence branches more than 0.5 m in length or >8 SDs above the species-wide mean.

Fig. S1. Genome-wide patterns of linkage disequilibrium. (A) LD decay curves for each chromosome. (B) Average level of linkage disequilibrium (LD) in 1-Mb windows along each chromosome.
Fig. S2. Bayesian hierarchical clustering of sorghum accessions based on 265,000 SNPs. Posterior probability of membership (Q) in each population at various values of K. Color-coding of Q-value bar plots (upper section) is arbitrary, whereas color-coding for rug plots (lower section) indicates morphological type as given in the legend. For clarity, only African and Asian source-identified accessions are displayed. The lowest cross-validation error was observed at K = 16.
Fig. S4. Genome-wide association studies for plant height components and inflorescence branch length. Manhattan plots and quantile-quantile plots for GLM and CMLM GWAS with Bonferroni significance threshold of 0.05 \[−\log_{10}(p) \sim 7\] noted with the horizontal line for plant height (A and B), flag-to-apex distance (E and F), preflag leaf height (C and D), and inflorescence branch length (G and H).
Fig. S5. Worldwide allelic distribution for a functional SNP (K162N) in the Ma1/SbPRR37 gene.

<table>
<thead>
<tr>
<th>Table S1. Decay of linkage disequilibrium in sorghum diversity panels</th>
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Distance in kilobases until the linkage disequilibrium decays to the given $r^2$ value. The ranges reflect the resolution of the analysis given the window sizes used.
Table S2. SNPs with significant association to inflorescence branch length

<table>
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<th>Chromosome</th>
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<th>Minor allele frequency</th>
<th>Effect size</th>
<th>Candidate gene</th>
<th>Distance to peak SNP</th>
<th>Description (Putative function)</th>
<th>Ref(s.)</th>
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<td>0.04</td>
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<td>30 kb, nearest</td>
<td>Receptor-like kinase, homolog of Theseus1 (Cell elongation)</td>
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<td>Sb01g027730</td>
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Chr., chromosome.

Other Supporting Information Files

Dataset S1 (XLSX)
Dataset S2 (XLSX)
Dataset S3 (PDF)
Dataset S3: High-resolution version of Bayesian hierarchical clustering of sorghum accessions based on 265,000 SNPs. Posterior probability of membership (Q) in each population for K=2 to K=19. Color-coding of Q-value bar plots (upper section) is arbitrary, while color-coding for rug plots (lower section) indicates morphological type (see legend). For clarity, only African and Asian source-identified accessions are displayed. The lowest cross-validation error was observed at K=16.