

Genome biology of the paleotetraploid perennial biomass crop *Miscanthus*

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Miscanthus is a perennial wild grass that is of global importance for paper production, roofing, horticultural plantings, and an emerging highly productive temperate biomass crop. We report a chromosome-scale assembly of the paleotetraploid *M. sinensis* genome, providing a resource for *Miscanthus* that links its chromosomes to the related diploid *Sorghum* and complex polyploid sugarcanes. The asymmetric distribution of transposons across the two homoeologous subgenomes proves *Miscanthus* paleo-allotetraploidy and identifies several balanced reciprocal homoeologous exchanges. Analysis of *M. sinensis* and *M. sacchariflorus* populations demonstrates extensive interspecific admixture and hybridization, and documents the origin of the highly productive triploid bioenergy crop *M. × giganteus*. Transcriptional profiling of leaves, stem, and rhizomes over growing seasons provides insight into rhizome development and nutrient recycling, processes critical for sustainable biomass accumulation in a perennial temperate grass. The *Miscanthus* genome expands the power of comparative genomics to understand traits of importance to Andropogoneae grasses.

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In addition to its historical roles in paper production and as ornamentals, varieties of the wild grass *Miscanthus* can produce high yields of harvestable vegetative biomass while maintaining and potentially increasing soil carbon¹. These features, enabled by C4 photosynthesis, perenniality, and related high efficiencies of light, nutrient, and water use, make *Miscanthus* and its close relatives (including sugarcanes and energy canes) promising candidates for economically feasible and sustainable bioenergy crops^{2–4}. Continued genetic improvement of bioenergy feedstocks is needed to enhance productivity and ensure that these crops remain robust in the face of ongoing biotic and abiotic stresses. This is particularly true for perennial grasses, where the advantages in economic and environmental sustainability relative to annuals depend on the longevity of the crop once established. Although perennial crops have tremendous potential for maximizing agricultural yields and minimizing environmental impacts, our knowledge of their biology and ability to manipulate their genetics lags well behind that in annual crops⁵.

A key limitation to the genetic improvement of perennial bioenergy grasses is the complexity of their genomes, which hinders the application of modern breeding approaches⁶. *Miscanthus sinensis* is a genetic diploid ($2n = 38$) with a genome size of $1C = 2.4–2.6$ Gb⁷; the related *M. sacchariflorus* occurs in both diploid ($2n = 38$) and tetraploid ($2n = 76$) forms. The $n = 19$ monoploid chromosome set of *Miscanthus* arose by ancient doubling of a sorghum-like $n = 10$ ancestor, with a single chromosomal fusion^{8–10}. Interspecific hybrids of *Miscanthus* form readily, even between individuals of different ploidy^{11,12}. Indeed, the predominant commercially grown miscanthus bioenergy variety is the high-yielding, sterile, asexually propagated triploid hybrid *M. × giganteus* “Illinois” ($3n = 57$). It is a clone of the taxonomic-type specimen, holotypus 1993–1780 Kew^{13,14}. Polyploidy is also common within the *Saccharum* complex, a group of closely related and highly productive perennial C4 grass species in the subtribe Saccharinae that includes sugarcanes (*Saccharum* spp.) and miscanthus. Intergeneric hybrid “miscanes” have been made by crossing miscanthus with hybrid sugarcanes¹⁵, suggesting that natural genetic variation in these two genera could be combined in order to blend desirable traits (e.g., cold tolerance and disease resistance).

Here we establish miscanthus as a genomic model for perenniality and polyploidy, and develop a foundation for genomic variation that will enable the future improvement of perennial biomass crops. We describe a draft chromosome-scale genome sequence for *M. sinensis*, prove that miscanthus is a paleoallotetraploid by analyzing the distribution of transposable elements across its genome, and establish the timing of key evolutionary events. By mRNA sequencing, we identify genes preferentially expressed in rhizomes, stems, and leaves, and explore the unique transcriptional dynamics of nutrient mobilization in this rhizomatous perennial grass. Unlike most perennial Andropogoneae, which are restricted to tropical or subtropical regions, the *Miscanthus* genus comprises species that naturally range from tropical to subarctic regions. Genomic analysis of 18 miscanthus accessions sequenced for this study, in addition to reduced representation genotyping of over 2000 accessions collected in the wild from east Asia, reveals extensive population structure and interspecific introgression, which further contributes to the genomic diversity of the genus *Miscanthus*.

Results

Genome sequence and organization. We assembled the *M. sinensis* genome into $n = 19$ chromosomes by combining short-read whole-genome shotgun (WGS) and fosmid-end data with

in vitro¹⁶ and in vivo¹⁷ chromatin proximity libraries (Supplementary Fig. 1, Supplementary Table 1, and Supplementary Notes 1, 2). The reference accession is the previously characterized⁸ doubled haploid DH1, which as expected is homozygous throughout. The genome assembly anchors 1.68 Gb of contigs to chromosomes, with a contig N50 length of 33.1 kb and pre-HiC scaffolding N50 length of 190 kb (Supplementary Table 2). An additional 0.20 Gb of contig sequence in scaffolds is not yet placed on linkage groups; highly repetitive sequences are problematic and missing from the assembly (Supplementary Fig. 1b). We validated the assembly at chromosome scale by comparison with an integrated genetic map with 4298 assignable markers (Supplementary Note 3).

We predicted the structure of 67,967 protein-coding genes based on several lines of evidence, including homology with other grasses and deep transcriptome data for miscanthus and sugarcane¹⁸. These predicted genes account for an estimated 98% of protein-coding genes, with 94% assigned to a chromosomal position (Supplementary Tables 3–5, Supplementary Fig. 5, and Supplementary Note 4). These genes are embedded within a sea of transposable element relicts and other repetitive sequences, which account for 72.4% of the *M. sinensis* genome assembly. The most common class of assembled transposons are gypsy long-terminal-repeat (LTR) retrotransposons (Supplementary Table 6 and Supplementary Note 5).

The paleotetraploidy of miscanthus is evident at the sequence level, since each sorghum chromosome aligns to a pair of *M. sinensis* chromosomes, after accounting for the chromosome fusion of ancestral sorghum 4- and 7-like chromosomes⁸ that reduces the karyotype from $n = 20$ to $n = 19$ (Fig. 1a). As expected from earlier genetic maps^{8–10} (Supplementary Fig. 3), the miscanthus and sorghum genomes show extensive 2:1 conserved collinear synteny (Fig. 1a and Supplementary Fig. 4a), consistent with a whole-genome duplication in the *Miscanthus* lineage. While it has been suggested¹⁹ that this duplication could be shared with sugarcane, comparison of *M. sinensis* and *S. spontaneum*²⁰ genomes shows that the duplications in the two lineages are distinct (Supplementary Note 7 and Fig. 2). Although the doubled genome and disomic genetics of miscanthus is suggestive of an allotetraploid history, neither a mechanism nor timing for paleotetraploidy has been described, in part due to the absence of known diploid progenitor lineages. We address this further below.

Regarding the more than twofold difference in bulk genome size between sorghum and miscanthus, we find that lengths of coding sequence and introns are generally similar (Supplementary Fig. 4b, c), with overall differences arising from increased intergenic spacing in miscanthus due to transposon insertion, as well as by the expansion of repetitive pericentromeric regions, which are only partially captured in the assembly (Supplementary Fig. 4b). The chromatin conformation contact map (Supplementary Fig. 2a) exhibits an enrichment of centromeric and telomeric contacts, respectively, consistent with the interphase nuclear “Rabl” conformation as seen in the barley genome²¹. We identified locally interacting chromosomal compartments (Supplementary Fig. 2b and Supplementary Note 2) for which A compartments have a higher gene density and B compartments have lower gene density (one-sided *t*-test *p* value $< 2.2 \times 10^{-16}$) and tend to occur predominantly in the pericentromeric region, as observed in other plants²².

Allotetraploid origin of *Miscanthus*. An allotetraploid (i.e., hybrid) origin for a paleotetraploid species is commonly demonstrated by showing that one set of its chromosomes (a subgenome) is more closely related to some diploid lineages to the

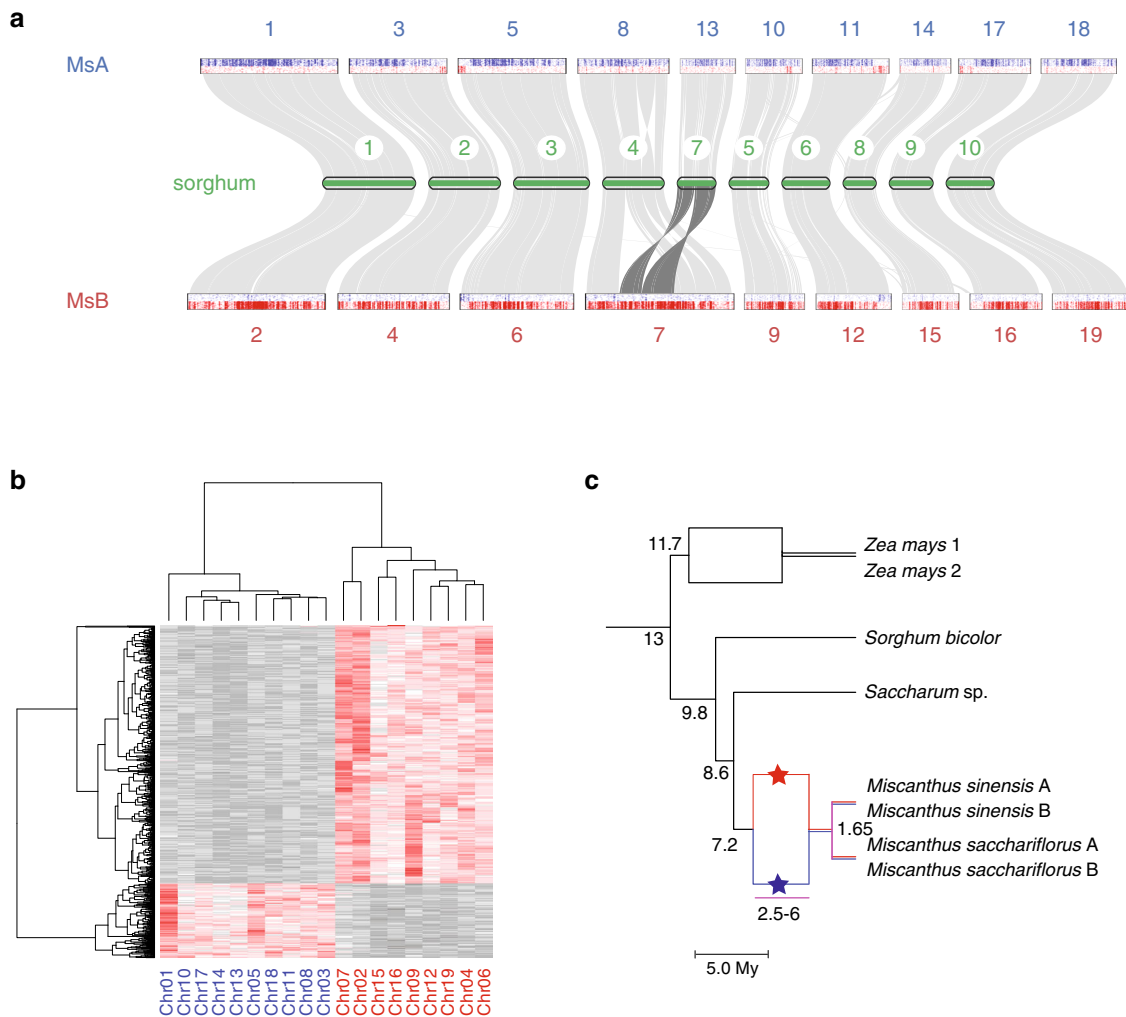


Fig. 1 Allotetraploidy in miscanthus. **a** Syntenic relationships between sorghum and *M. sinensis* subgenomes MsA and MsB. Distribution of subgenome-specific 13-mer sequences (blue for MsA, red for MsB) is shown for each *M. sinensis* chromosome (see text and Supplementary Note 7.1). **b** Clustering of counts of 13-mers that differentiate homeologous chromosomes enables the consistent partitioning of the genome into two subgenomes. Blue chromosome names correspond to the A subgenome, red chromosome names correspond to the B subgenome. **c** Timetree of Andropogoneae showing the timeline of allotetraploidy in the *Miscanthus* lineage, with divergence and hybridization times of the A and B progenitors estimated from sequence comparisons (Supplementary Note 8). Source Data underlying Fig. 1b are provided as a Source Data file.

exclusion of others²³. Because there are no known candidates for the diploid progenitors of tetraploid miscanthus, this approach cannot be used here. Instead, we used a new method that relies on the chromosomal distribution of repetitive elements, which can provide robust markers for subgenome ancestry²⁴. We sought repetitive sequences whose presence is enriched on one member of each homeologous chromosome pair (Supplementary Note 6). Such sequences are definitive markers of allotetraploidy, and occur as relicts of repetitive elements that were active in only one of the two diploid progenitors prior to hybridization and genome doubling²⁴. Importantly, the method does not require access to or even knowledge of living representatives of the progenitor lineages. We found 1187 13-bp sequences (13-mers) whose pairwise enrichment pattern consistently partitions homeologous chromosome pairs between distinct A and B subgenomes (Fig. 1a, b). This observation establishes the past existence of distinct A and B progenitor lineages (which remained separate for millions of years, see below), and the allotetraploid origin of miscanthus.

Although we can use these markers to assign each miscanthus chromosome in bulk to the A or B subgenome, we find evidence for the balanced reciprocal exchange of distal segments between

homeologous chromosomes such that dosage remains intact (e.g., the ends of chromosomes 5–6, 11–12, and 16–17; Figs. 1a, 3a, Supplementary Fig. 6, and Supplementary Note 6). Based on consistency with our dense genetic map, these are clearly bona fide homeologous exchanges rather than misassemblies. The observed distal reciprocal exchanges likely occurred either by mitotic recombination in the vegetative tissue of an AB F1 hybrid founder prior to genome doubling, or by aberrant homeologous recombination after allotetraploidy. The concentration of these exchanges toward the ends of chromosomes is consistent with the proximity of these regions in a telomeric bouquet conformation. The maintenance of discrete A/B patterns of diagnostic 13-mers in these distal segments implies that these exchanges occurred by single crossover events rather than recurring recombination throughout the distal regions of the chromosomes, which would blur the distinctive A/B 13-mer signature.

Discrete homeologous exchanges are often observed in newly formed allotetraploids and are thought to occur in response to a new meiotic environment²⁵. In studies of other polyploids, homeologous replacements that alter the balance between A and B alleles are common; when such variants are segregating in a

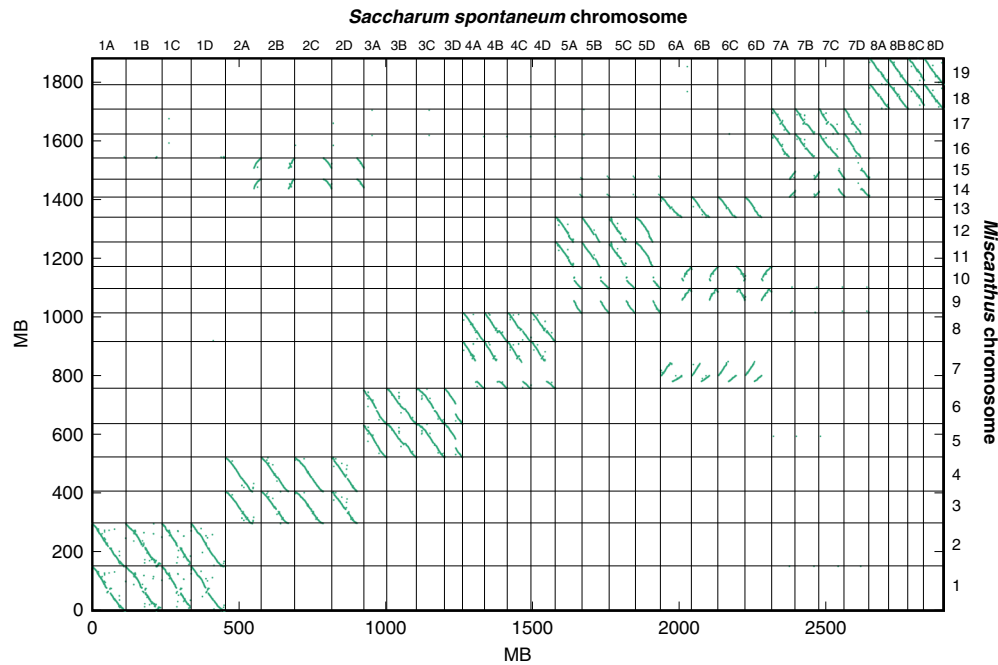


Fig. 2 Miscanthus-Saccharum synteny. Dotplot showing co-orthologs between *Miscanthus sinensis* and *Saccharum spontaneum*. All syntenic genes with c score ≥ 0.7 are shown using the mscscan ortholog algorithm. Note the 2:4 ratio of miscanthus:sugarcane chromosome segments. Source Data are provided as a Source Data file.

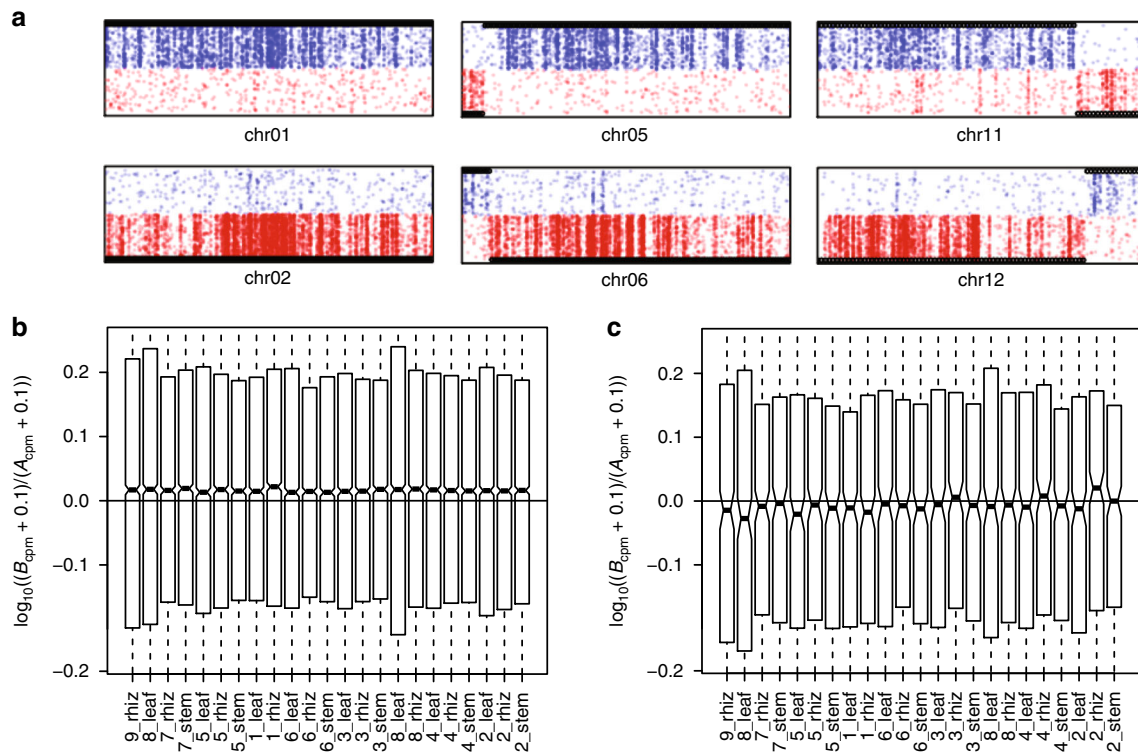


Fig. 3 Post-allotetraploidy reciprocal exchanges. **a** Example of a chromosome pair without reciprocal exchange (chr01–chr02), and two chromosome pairs with distal reciprocal exchanges (chr05–chr06 and chr11–chr12). Red and blue dots represent occurrences of subgenome-specific 13-mers. Black bars identify A and B ancestry inferred from a Hidden Markov Model (Supplementary Note 7.2). **b** Relative expression of homeologous gene pairs. Across tissues and seasonal sampling times, there is a 3.8% median bias toward the expression of the B member of the pair. **c** Homeologous gene pairs within reciprocally exchanged regions show the expression bias of their ancestral location. Source Data underlying Fig. 3b, c are provided in the Source Data file.

population, the resulting genetic variation can underlie quantitative trait loci^{26,27}. In contrast to these studies, however, in *Miscanthus*, we find (1) predominantly balanced reciprocal exchanges that alter chromosomal linkage, but do not change A/B dosage, and (2) no evidence that these segmental exchanges are segregating in our sequenced samples, suggesting that the reciprocal homeologous exchanges are the result of ancient events that have become fixed in *Miscanthus* (and therefore cannot be causal for any phenotypic variation in the genus) (Supplementary Note 6)). In addition to these long fixed reciprocal exchanges, there are several shorter internal homeologous segments (Supplementary Note 6) that could correspond to nonreciprocal or recurrent exchange. These segments will be interesting to study further.

From the identification of distinct A and B subgenomes, we see that the sorghum-7 and -4-like chromosomes that fused⁸ to form miscanthus chromosome 7 were both derived from the B progenitor. While it is possible that the fusion occurred in the B progenitor itself prior to hybridization, the absence of other Saccharinae with $n=9$ chromosomes, and the likelihood of chromosome instability in the aftermath of allotetraploidization, suggests that the fusion occurred after allohybridization.

The timeline of paleotetraploidy in miscanthus can be established through inter- and intra-subgenome comparisons (Fig. 1c and Supplementary Note 7). We estimate that the A and B progenitors diverged from their common ancestor ~7.2 Mya (million years ago), based on the synonymous differences between homeologous protein-coding genes (Supplementary Fig. 7). After this divergence but before hybridization, the two (now likely extinct) progenitors evolved independently; evidence of their species-specific transposable element activity appears in the contemporary *Miscanthus* genome as subgenome-specific repeats²⁴. Consistent with this hypothesis, we find several LTR-retrotransposon families within only one of the two subgenomes, and estimate that they were actively inserting during the period ~2.5–6 Mya (Supplementary Note 7). In contrast, transposon activity after the allotetraploidy event should be distributed across the entire *Miscanthus* genome without regard to subgenomes. Also, consistent with this picture, we find a burst of transposon activity that is not subgenome-specific starting ~2.5 Mya, which serves as our best estimate for the allotetraploid origin of *Miscanthus* (Supplementary Note 7 and Supplementary Fig. 7c). Finally, the interfertile sister species *M. sinensis* and *M. sacchariflorus* diverged ~1.65 Mya (Fig. 1c), consistent with speciation occurring after allotetraploidy. Chromosome-level comparisons of repetitive elements and protein sequences confirm that the polyploidies of *Miscanthus* and sugarcane occurred independently (Supplementary Note 7).

Common hallmarks of allopolyploidy are asymmetric gene loss (or conversely, retention) and biased gene expression between subgenomes, which are both thought to arise from epigenetic asymmetries in the aftermath of allohybridization^{28,29}. Comparing miscanthus and sorghum genes, we find that ~29% of sorghum genes have been lost on one of two subgenomes; conversely, ~71% have co-orthologs on both subgenomes (Supplementary Note 6). Gene retention in *M. sinensis* shows a small but statistically significant bias toward the B subgenome (87.1% genes retained on B vs. 83.9% on A, Supplementary Table 7; Fisher's exact p value, two-sided = 1.2×10^{-9}). The level of homeologous gene retention in *M. sinensis* is nearly twice that of maize (71% vs. 36%), presumably because the miscanthus allotetraploidy is more recent. The subgenome retention bias in *Miscanthus* is also smaller than in maize²⁸ (80.6% in maize 1 vs. 55.4% in maize 2), which may reflect differences in the degree of genomic differentiation between maize versus *Miscanthus* progenitors prior to hybridization.

Similarly, for retained homeolog pairs, we find a weak but significant expression bias (median B/A expression ratio 1.038, without strong variation across tissues or season, Fig. 3b). Although most pairs of homeologous genes have similar expression levels, there are ~10% more pairs with higher B-subgenome expression than vice versa (Supplementary Table 8). This is again notably weaker than the expression bias in maize²⁸. Interestingly, genes in regions of homeologous exchange show (on average) the bias of their source subgenome (Supplementary Note 8 and Fig. 3c), indicating that subgenome expression bias arises from local effects and/or became fixed early in the allotetraploid evolution. This observation is consistent with experiments that show rapid development of subgenome bias in neoallopolyploids^{25,30,31}. The weaker subgenome expression and retention bias seen in the more recent miscanthus allotetraploidy versus the older maize suggests that these effects may become amplified over time, and may also be influenced by the relative genomic divergence of progenitors.

Seasonal dynamics of gene expression. As a rhizomatous perennial, miscanthus provides a model for studying the biology of rhizomes, which are modified underground stems that enable temperate perennial grasses to overwinter by their capacity to (1) store nitrogen, carbon, and other nutrients from senescing leaves and stems, and (2) mobilize these reserves in the spring to feed new vegetative growth. Amino acids, particularly asparagine with its high N:C ratio, are the primary form of nitrogen cycled among plant tissues³². Monitoring free asparagine concentrations (Fig. 4a) from stem, leaf, and rhizome tissues of *M. × giganteus* sampled throughout the growing season (May to October) over 3 years revealed high concentrations in the spring rhizome, low levels in all tissues during the summer period of rapid growth, followed by increasing accumulation in stem and rhizomes after flowering. Elevated asparagine levels mark periods of active nitrogen remobilization from rhizome to shoot in spring, and from the shoot to rhizome in autumn.

To characterize the seasonal dynamics of gene expression and regulatory programs associated with perenniality in *Miscanthus*, we performed RNA-seq from the same tissue samples collected for profiling nitrogen cycling (Supplementary Note 8 and Supplementary Data 1). Principal component analysis (PCA) identified the two largest sources of variation as tissue type, followed by sampling time (Fig. 4b). Comparisons among tissues produced a catalog of organ-preferred genes (Supplementary Fig. 8 and Supplementary Data 1–9). As expected, leaf-preferred genes are significantly enriched in genes functioning in carbon fixation and metabolism, and stem-preferred genes include those associated with phenylpropanoid biosynthesis and amino acid metabolism. Gene expression in rhizomes is more similar to stems than leaves, consistent with their developmental origin as modified stems (Supplementary Fig. 8a, b). Relative to stems and leaves, rhizomes preferentially express transcription factors that regulate growth and metabolic processes, and genes that respond to stimuli such as water and stress (Supplementary Fig. 8e). We identified 35 genes that are preferentially expressed in the rhizome, including homologs of genes like *GIANT KILLER (GIK)* and *SHORT INTERNODE (SHI)* implicated in organ patterning, differentiation, and cell elongation^{33–36}. Overexpression of *SHI*-like genes results in compact plants with shorter stem internodes^{37–39}, which is consistent with the morphological differences between miscanthus rhizomes and stems.

We identified and characterized the transcriptional network regulating seasonal nutrient mobilization in miscanthus (Supplementary Note 8), which is central to the perennial lifecycle and efficient recycling of resources. Although tissue identity

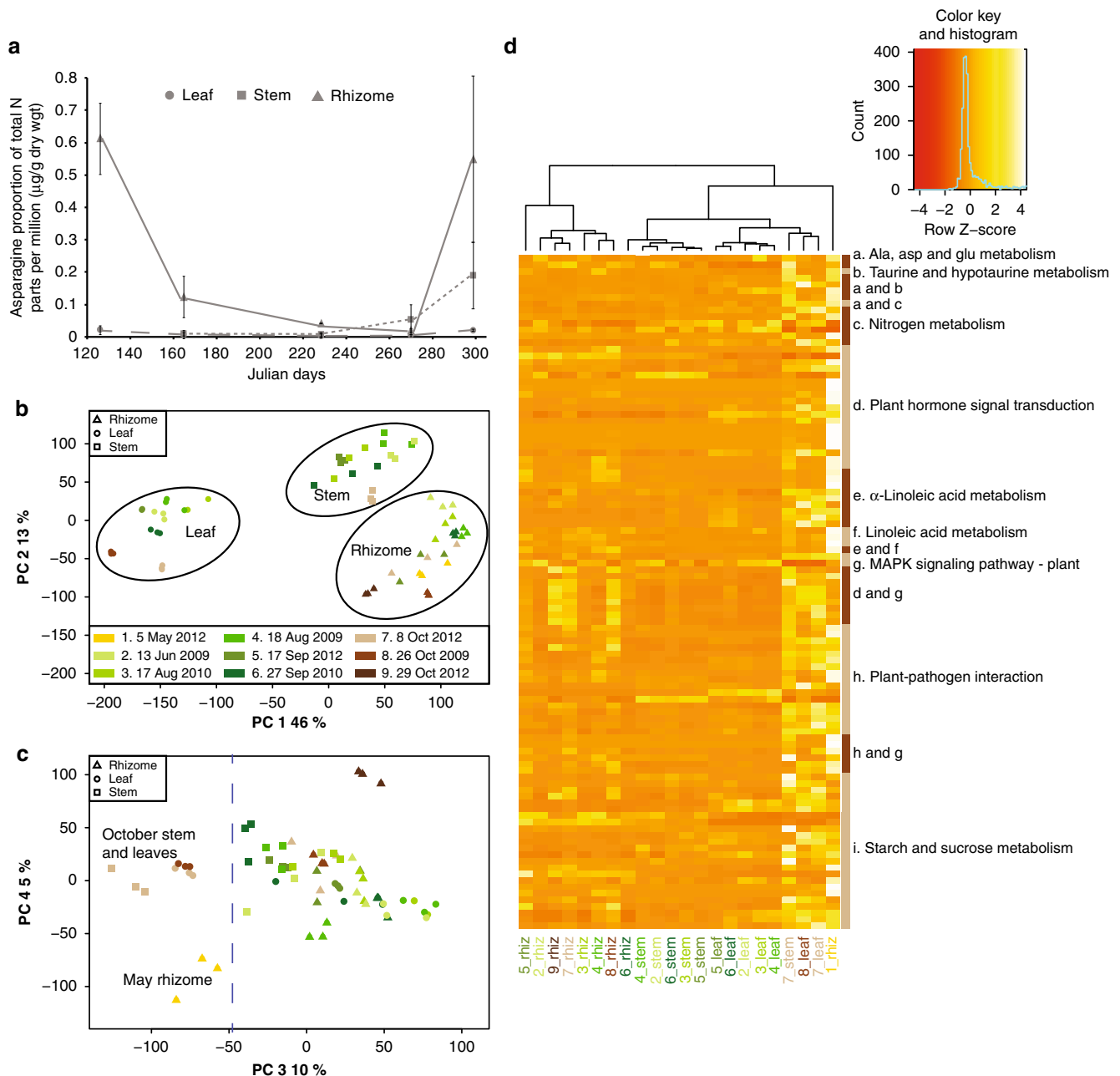


Fig. 4 Seasonal gene expression changes in miscanthus. **a** Shows asparagine in rhizome, stem, and leaves over the growing seasons normalized to total nitrogen in the sample. The error bars represent the standard deviation. **b** Principal component analysis of RNA-seq read counts normalized using the DESeq2 variance-stabilizing transformation method. PC1/2 distinguishes the three tissues from each other. **c** PC3/PC4 separates samples based on their nutrient mobilization status. The color scheme for the organs and dates matches **b**. **d** Heatmap across all tissues in the study comparing the expression of a subset of genes expressed in tissues that are actively remobilizing nutrients. Source Data are provided as a Source Data file.

dominates the first two principal components of gene expression, the third component (PC3) separates the spring rhizomes, fall leaves, and fall stems from the other tissues (Fig. 4c). Differentially expressed genes contributing to the pattern in PC3 (Supplementary Note 8) comprise a dynamic network differentiating the fall rhizome that is storing nitrogen from the spring rhizomes that are releasing nitrogen to promote new growth (Supplementary Data 1). Of these genes, 104 had a functional or KEGG assignment, including a suite of transcription factors and genes with known important roles in nitrogen mobilization⁴⁰ like *ASPARAGINE SYNTHETASE (ASN1)*, *GLUTAMATE DEHYDROGENASE (GDH2)*, and *GLUTAMATE DECARBOXYLASE (GAD1)*. Remarkably, the most prominent (“hubby” or central)

transcription factors within the network are a subset of JASMONATE ZIM DOMAIN (JAZ) family proteins that regulate jasmonic acid biosynthesis (e.g., *ALLENE OXIDE SYNTHASE, AOS*) and signaling, a pathway recently shown to activate nitrogen remobilization in rice⁴¹ (Fig. 4d). These data reveal a group of regulators and enzymes that may be key for promoting the nitrogen remobilization in spring.

Inter- and intraspecific variation and introgression. Breeding to improve miscanthus for biomass and other applications can draw upon extensive wild germplasm from multiple species and ploidy levels. We therefore investigated the genetic diversity of *Miscanthus* and the distribution of inter- and intraspecific variation

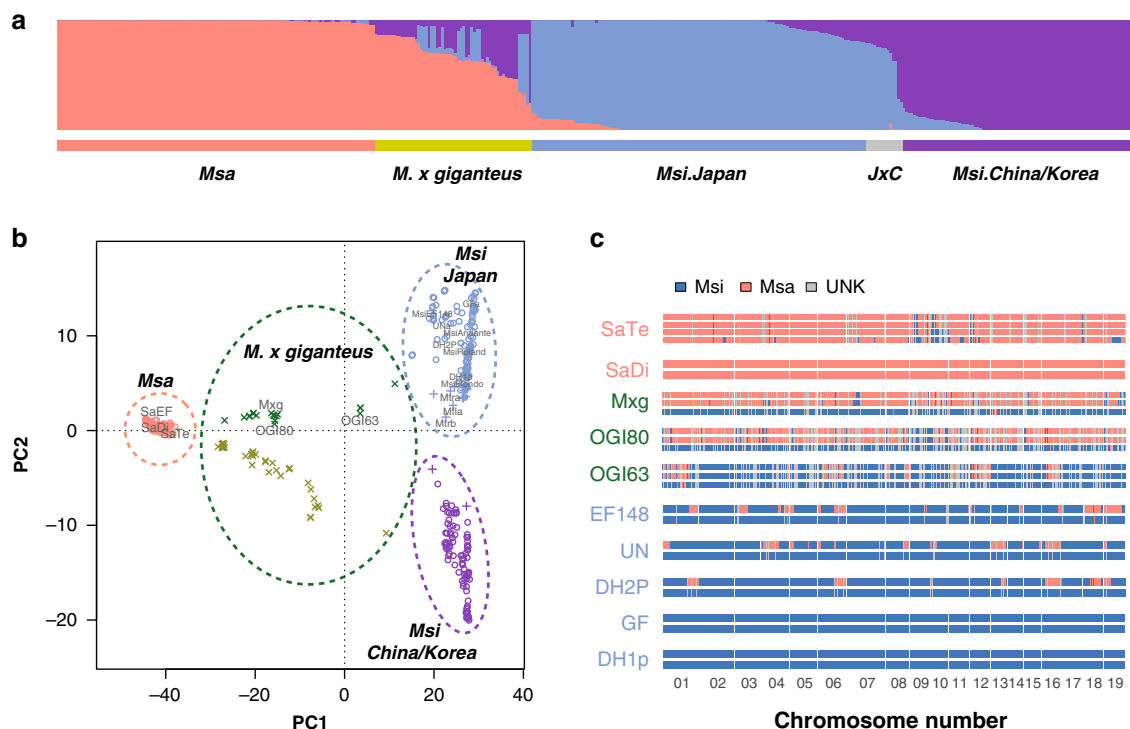


Fig. 5 *Miscanthus* population structure and segmental ancestry. **a** Population structure of 407 *Miscanthus* accessions, including 57 *M. × giganteus*, 120 *M. sacchariflorus* (*Msa*), and *M. sinensis* (*Msi*) from China (75), Korea (15), and Japan (140). **b** Principal component analysis of 407 *Miscanthus* accessions where “x” marks admixtures of *Msa* and *Msi*. Such hybrids are collectively referred to as *M. × giganteus* (*Mxg*), and can be diploid, triploid, or tetraploid. Separation of Japanese and mainland Asian populations is largely consistent with structure analysis in **a**. Whole-genome shotgun (WGS)-sequenced accessions are labeled. **c** Segmental ancestry of *Miscanthus* accessions based on WGS sequencing. Each horizontal bar denotes one (imputed) haploid chromosome set; red and blue indicate *Msa* and *Msi* ancestry, respectively. The number of bars represents ploidy. Introgression of *M. sacchariflorus* into *M. sinensis* (*Msi*EF148, Undine, DH2, DH2P) is common among cultivated European types (Supplementary Fig. 10). Source Data underlying Fig. 5c are provided as a Source Data file.

in admixed populations. We combined new WGS sequencing of 18 accessions of varying ploidy, including the triploid biofuel cultivar *M. × giganteus* “Illinois” (see Supplementary Note 9 and Supplementary Table 9) with previously generated genotyping-by-sequencing data from primarily wild accessions with broad geographic coverage^{11,12,42,43}, spanning the native range of *Miscanthus* across north- and south-east China, Korea, Russia, and Japan. Genome-wide admixture (Fig. 5a) and PCA (Fig. 5b) readily differentiate two species, *M. sinensis* and *M. sacchariflorus*. Other named *Miscanthus* accessions, such as *M. transmorrisonensis* and *M. floridulus*, lie within the range of genetic variation of *M. sinensis*, suggesting that these taxa should more properly be considered subtypes of *M. sinensis*. The accession in our collection named *Miscanthus junceus*, however, is clearly distinct and appears to be more closely related to sugarcane than *Miscanthus* (Supplementary Fig. 9). It is African, sometimes classified in a separate genus *Miscanthidium*, and clearly separate from *Miscanthus sensu stricto*⁴⁴.

Our chromosome-scale genome assembly allows us to investigate patterns of admixture in interspecific hybrids (Fig. 5c). While all *M. sinensis* × *M. sacchariflorus* hybrids and admixtures are taxonomically characterized as *M. × giganteus*, this nothospecies has rich diversity due to the occurrence of diploid, triploid, and tetraploid accessions (Supplementary Fig. 10). We find that many ornamental diploids, especially many bred by Ernst Pagels in Germany, contain chromosomal segments of *M. sacchariflorus* introgressed into an *M. sinensis* background, consistent with prior admixture studies^{11,12}. Mainland Asian and Japanese *M. sinensis* are distinct subpopulations (Fig. 5a) that diverged ~500,000–1,000,000 years ago based on chloroplast DNA (Supplementary Note 9).

Our data confirm that the highly productive triploid biofuel *M. × giganteus* genotype, “Illinois,” is an interspecific hybrid of tetraploid *M. sacchariflorus* and diploid *M. sinensis*^{14,45}. We find a predominant 2:1 ratio of *M. sacchariflorus*:*M. sinensis* alleles across the entire genome, consistent with this hypothesis; however, we also observed that the *M. sacchariflorus* ancestor had interspecific admixture (Fig. 5c and Supplementary Fig. 10c, e), which indicates that the most productive *Miscanthus* genotype currently grown is the product of more than one cycle of introgression from *M. sinensis* into *M. sacchariflorus*. Hybrids between *M. sacchariflorus* and *M. sinensis* are frequently highly vigorous and high-yielding, regardless of whether they are diploid, triploid, or tetraploid^{46,47}. Thus, understanding how prior introgression of *M. sinensis* alleles into a primarily *M. sacchariflorus* genetic background affects the yield potential of subsequent interspecific hybrids will be important for optimizing breeding strategies. In particular, *M. × giganteus* combines the tufted habit (many stems per area; short rhizomes) of its *M. sinensis* parent with the spreading rhizomatous habit (few stems per area; long rhizomes) of its *M. sacchariflorus* parent, typically in an intermediate form, and optimizing the number of stems per area is critical to breeding for high yield in *M. × giganteus*⁴⁸. The recently collected Japanese *M. × giganteus* triploid⁴⁹ “Ogi80” has a similar pattern to “Illinois,” with both including several short blocks containing two or three *M. sinensis* alleles. These regions could be due to segmental gene conversion or loss during the propagation of this sterile triploid, or interspecific introgression prior to triploid formation. Another natural triploid, “Ogi63,” shows a distinct pattern, highlighting the diversity of natural polyploid *Miscanthus* hybrids (Supplementary Fig. 10).

Miscanthus is a promising perennial biomass source and candidate biofuel crop with efficient C4 photosynthesis that is highly adaptable. Its ability to grow on marginal lands with limited inputs, and its high drought and chilling tolerance make it suitable for both tropical and temperate climates. The genome sequence and genomic analysis presented here provides a foundation for systematic improvement of *Miscanthus* to optimize its productivity and robustness. Comparative analyses among the Andropogoneae⁵⁰, which unites *Miscanthus* with maize, sorghum, and sugarcane, promise to reveal the genetic basis for innovations that contribute to the high productivity and wide adaptation of this tribe of grasses.

Methods

Genome sequencing and chromosomal assembly. We shotgun-sequenced the *M. sinensis* genome at ~90× redundancy with Illumina paired-end and mate-pair data, augmented by fosmid-end pairs and in vitro and in vivo chromatin conformation capture (HiC) as described in Supplementary Note 1. Illumina shotgun assembly was performed with Meraculous⁵¹ and organized into chromosomes with HiC data using HiRise (Dovetail Genomics, Scotts Valley, CA) followed by manual curation with Juicebox⁵², and confirmation of internal self-consistency as described in Supplementary Note 2. The assembly was further corroborated and assigned to chromosomes using a genetic map derived from four crosses, with 4298 uniquely assignable 64-bp markers, as described in Supplementary Note 3.

Protein-coding gene and transposable element annotation. Protein-coding gene structures were annotated using the DOE Joint Genome Institute annotation pipeline⁵³ that incorporates transcriptional evidence, homology support from related grasses, and ab initio methods, as described in Supplementary Note 4. RNA-seq data from three tissues and 57 timepoints for *M. × giganteus* and *M. sinensis* DH1 leaf and rhizome (PRJNA575573, SRP017791) were used, and these data are summarized in Supplementary Note 8 including accession numbers. Genome completeness was estimated using BUSCO⁵⁴, and orthologous gene families identified using OrthoVenn⁵⁵ as described in Supplementary Note 4.

Transposable elements were identified de novo using RepeatModeler⁵⁶ to augment existing catalogs of grass repeats from rebase⁵⁷ and MIPS⁵⁸ using RepeatMasker⁵⁹, and identified intact retrotransposons with LTRHarvest⁶⁰, as described in Supplementary Note 5. LTR families were defined by clustering these LTRs with those of sorghum and sugarcane by BLAST score using 90% identity and 90% length cutoffs as described in Supplementary Note 5.

Subgenome and homeologous exchange identification. We partitioned the *M. sinensis* genome into subgenomes A and B by a modification of methods described in Session et al.²⁴ and described more fully in Supplementary Note 6. Importantly, this method can be applied without requiring sequences from extant A and B diploids. Briefly, we identified 1187 13-bp sequences (13-mers) that (1) occurred at least 100 times across the genome, and (2) were at least twofold enriched in one member of each homeologous chromosome pair (excluding the case of fused homeologs). 13-mers were counted using Jellyfish⁶¹. Homeologous chromosomes were determined based on conserved synteny to each other and to sorghum (Fig. 1). These 13-mers allowed chromosomes to be clustered by subgenome, and were found to overlap with subgenome-specific repeats as described in Supplementary Note 6. To identify cases of homeologous exchanges, we sought chromosomal regions whose 13-mer identity differed from the overall identity of the chromosome, using a hidden Markov model whose observed state was the number of A- and B-specific 13-mers and whose emitted state is A or B, as described in Supplementary Note 6.

Determination of biases in subgenome gene retention. We used two methods to determine orthology between *M. sinensis* genes and sorghum in order to assess differential retention of gene duplicates after allotetraploidy, using sorghum as the outgroup representing the ancestral (preduplicated state). For the first method, gene families were constructed using OrthoVenn⁵⁵. For the second method, we used BLAST-based clustering. Subgenome-specific retention is defined as the number of genes on a given subgenome divided by the number of inferred ancestral (i.e., preduplication) gene number. Details of this analysis can be found in Supplementary Note 6.

Timing of events associated with allotetraploidy. We estimated the timing of speciations in the Andropogoneae using a set of 1:1 orthologs for species shown in Fig. 1c with *P. hallii* and *S. italica* as outgroups, as described in Supplementary Note 7. Briefly, concatenated multiple-sequence alignments were produced using Dialign-TX⁶² and Gblocks⁶³. *M. sinensis* and maize genes were partitioned into A and B subgenomes, and 1 and 2 subgenomes, respectively, with 1–2 assignments as determined by Schnable et al.²⁸. The dataset included *M. sacchariflorus* A and B genes predicted by mapping diploid *M. sacchariflorus* shotgun sequence to the *M.*

sinensis assembly. *M. sacchariflorus* has the same karyotype as *M. sinensis*, and hybrids are fertile, indicating that they share the same A/B ancestral tetraploidy. Phylogenies were produced from the resulting 28,887 nucleotide alignment using PhyML⁶⁴. Timetrees were estimated using r8s⁶⁵ with a smoothing parameter of 0.1, and constraining the *Setaria/Panicum* node to 12.8–20 Mya and the *Sorghum/maize* split to 13–21.2 Mya⁶⁶.

We estimated the period during which the A and B progenitors were separate species using phylogenies of five subgenome-specific LTR families with ≥100 members that contain a subgenome-enriched 13-mer, as described in Supplementary Note 7. Subgenome-specific LTR families have been active when the two progenitors were separate species, but before allotetraploidy. To calibrate the rate of LTR substitution in *Miscanthus*, we used LTR families that are (1) found in high copy number in *Miscanthus* across both the A and B subgenomes, and so were active after allotetraploidy, and (2) have parallel activity in the sorghum genome, and used a *Miscanthus*–sorghum divergence time of 10 My as determined from protein-coding genes. We used the median substitution rate of these families (2.1×10^{-8} substitutions per My) to infer the timing of subgenome-specific activity based on Jukes–Cantor distance. Details are provided in Supplementary Note 7.

Analysis of gene expression. We analyzed RNA-seq data using Tophat2.1.1⁶⁷, HTSeq⁶⁸, DESeq2⁶⁹, and the NOISeq R package^{70,71} to extract expression levels and further analyze the RNA-seq data as described in Supplementary Note 8. To identify genes that were constitutively expressed in any one organ type, we considered only genes with a count per million (cpm) of 5 or greater within all samples of an organ type. KEGG enrichment analysis using keggseq⁷² was performed on genes that were preferentially in leaves, stems, and rhizomes, respectively, to determine if they clustered into specific pathways or functional categories. Enriched pathways with a *q* value ≤ 0.01 are shown in Supplementary Fig. 8c–e.

For the purposes of comparing gene expression of homeologs, we measured gene expression using cpm, after combining replicates, as described in Supplementary Note 8. In order to measure subgenome expression bias, for each homeolog pair, we considered only experiments where one or both homeologs have nonzero expression (cpm > 0.5). This condition is necessary because the majority of genes are not expressed in every tissue, leading to a large number of uninformative comparisons. We considered expression bias using a variant of the approach of Schnable et al.²⁸, identifying homeolog pairs where one member of the pair was expressed *X*-fold relative to the other, where *X* = 2, 5, and 10, again requiring both members to be expressed at a minimal level (cpm > 0.5) to avoid uninformative comparisons.

Analysis of genetic variation. WGS sequences of 18 *Miscanthus* accessions (Supplementary Table 9) were aligned to the haploid *M. sinensis* DH1 reference sequence using bwa mem⁷³, and variants called using GATK⁷⁴ version 3.6, as described in Supplementary Note 9. Restriction site-associated DNA-sequencing (RAD-seq) data from 2819 *Miscanthus* individuals were used to obtain a snapshot of genetic diversity, as described in Supplementary Note 9.

For PCA with the RAD-seq data genotypes, we retained SNPs with a maximum of 30% missing data and a minimum minor allele frequency of 0.01, resulting in a set of 144,337 SNPs. From this dataset, individuals with 50% or more missing data were removed, leaving 2492 out of the original 2819 individuals. By filtering SNPs and individuals in this way, the remaining data were primarily derived from *PstI* sequencing libraries, as this was the enzyme most commonly used across the dataset. Genotypes were coded on a numeric scale from 0 to 1, indicating copy number for the nonreference allele, i.e., 0, 0.5, and 1 for diploids, 0, 0.33, 0.67, and 1 for triploids, and 0, 0.25, 0.5, 0.75, and 1 for tetraploids. PCA was performed using probabilistic PCA method implemented in the Bioconductor package pcaMethods⁷⁵. All SNPs were centered and scaled to unit variance before PCA.

The genomic makeup of the accessions was analyzed with ADMIXTURE⁷⁶. Figure 5a shows the result for *K* = 3, which was used to analyze the populations. To resolve admixture along chromosomes, we identified 1283,756 species-specific SNPs in the nonrepetitive regions of 19 chromosomes from fixed differences between the two species as represented by 4 diploid exemplar genomes without evident admixture as described in Supplementary Note 9. These ancestry-informative markers were used to obtain a high-resolution admixture map for the WGS accessions (Fig. 5c), following the method of Wu et al.⁷⁷. A subset of these ancestry-informative markers that overlapped RAD-seq variants were used to infer the segmental ancestry of the RAD-seq accessions. Further details are provided in Supplementary Note 9 and Supplementary Data 10.

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

Data availability

Data supporting the findings of this work are available within the paper and its Supplementary Information files. A reporting summary for this article is available as a Supplementary Information file. The datasets generated and analyzed during the current study are available from the corresponding author upon request. Genomic reads for the *M. sinensis* DH1 genome assembly can be found at PRJNA346689, transcriptomic reads

at [PRJNA575573](https://doi.org/10.1038/s41467-020-18923-6) and [SRP017791](https://doi.org/10.1038/s41467-020-18923-6). The genome, annotation, transcriptomic, and variation data are available on [Phytozome](https://phytozome.jgi.doe.gov/). Source data are provided with this paper.

Code availability

All custom scripts used for parsing and analyzing transposable elements, gene families, and gene expression, as described in Supplementary Notes, are available at GitHub [<https://github.com/miscanthus-paper/Miscanthus-genome.git>] and [<https://bitbucket.org/bredeson/artisanal.git>].

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Author contributions

D.S.R., K.S., T.M., S.P.M., and M.E.H. provided project leadership. D.S.R., K.S., T.M., S.P.M., A.M.S., B.T.J., G.A.W., and L.V.C. provided figures and wrote the paper. T.M. and J.V.B. assembled the genome and conducted the comparative genomics analysis. N.H.P. generated the HiC assembly. A.M.S. conducted the repeat and allotetraploidy analysis. B.T.J. and M.B.B. conducted the transcriptomic analysis. G.A.W. analyzed the genetic diversity and introgression patterns. S.S. provided the protein-coding gene annotation. H.D. and S.L. provided genetic map data. AB collected samples and provided the transcriptomic, amino acid, and nitrogen data. J.R.H., A.O., and J.E.M. processed samples and extracted nucleic acids for the project. K.F. and I.S.D. contributed the *M. sacchariflorus* whole-genome sequencing data. J.Gr., J.S., and K.B. coordinated the genome sequencing. K.G. created the double-haploid line and S.J. provided the line. W.B.C. and J.Gi. generated the mapping populations, and J.A.J. and E.J.S. oversaw the generation of the mapping populations. T.Y. provided Ogi63 and Ogi80 triploid lines. J.D.V., L.V.C., S.B., and E.J.S. contributed to RAD-seq data, and J.D.V. and L.V.C. used these data to call variants. M.K., T.H., L.L., X.J., J.P., C.Y.Y., K.H., J.H.Y., and B.K.G. provided miscanthus germplasm.

Competing interests

Dovetail Genomics LLC is a commercial entity developing genome assembly methods. N.H.P. was an employee of Dovetail Genomics, and D.S.R. is a scientific advisor to and minor investor in Dovetail.

Additional information

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