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Data Note

A chromosome-level reference genome of *Ensete glaucum* gives insight into diversity and chromosomal and repetitive sequence evolution in the Musaceae

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Abstract

Background: Ensete glaucum (2n = 2x = 18) is a giant herbaceous monocotyledonous plant in the small Musaceae family along with banana (Musa). A high-quality reference genome sequence assembly of *E. glaucum* is a resource for functional and evolutionary studies of Ensete, Musaceae, and the Zingiberales.

Conclusions: The advance towards a Musaceae pangenome including *E. glaucum*, tolerant of extreme environments, makes a complete set of gene alleles, copy number variation, and a reference for structural variation available for crop breeding and understanding environmental responses. The chromosome-scale genome assembly shows the nature of chromosomal fusion and translocation events during speciation, and features of rapid repetitive DNA change in terms of copy number, sequence, and genomic location, critical to understanding its role in diversity and evolution.

Keywords: centromeres, chromosome-scale assembly, *Ensete glaucum*, Musaceae evolution, Nanopore, pangenome, repetitive DNA, retrotransposons, synteny, translocations

Background

The genus Ensete Bruce ex Horaninow (Musaceae) includes 10 species of giant, herbaceous monocotyledonous plants, native to tropical Africa and Asia [1]. Among them, the African species Ensete ventricosum (Welw.) Cheesman (enset) is an important food crop for >20 million people in Ethiopia [2]. Its sister genus Musa, grown throughout the tropics for food and fibre, includes diploid species, triploids, and hybrids of Musa acuminata and Musa balbisiana, with banana cultivars. Sisters to the grasses (Poales) and palms (Arecales) in monocots, both Ensete and Musa, along with a third genus Musella, belong to Musaceae in the order

Zingiberales (gingers and bananas) [3]. Following rapid diversification of the Zingiberales at the Cretaceous/Tertiary boundary (>65 million years ago [Mya]) the crown node age of the Musaceae family soon appears, with the Musa genus diverging from Ensete and Musella \sim 40 Mya [4, 5].

Ensete glaucum (Roxb.) Cheesman (NCBI: txid482298), like other species in Ensete, is monocarpic with a dilated and characteristically glaucous basal pseudo-stem (Fig. 1A–E), with a small number of large seeds (10 mm in diameter) in elongated, banana-like fruits, borne in hands with a terminal flower and is diploid with 2n = 2x = 18 chromosomes [1, 6–10]. E. glaucum is widely distributed

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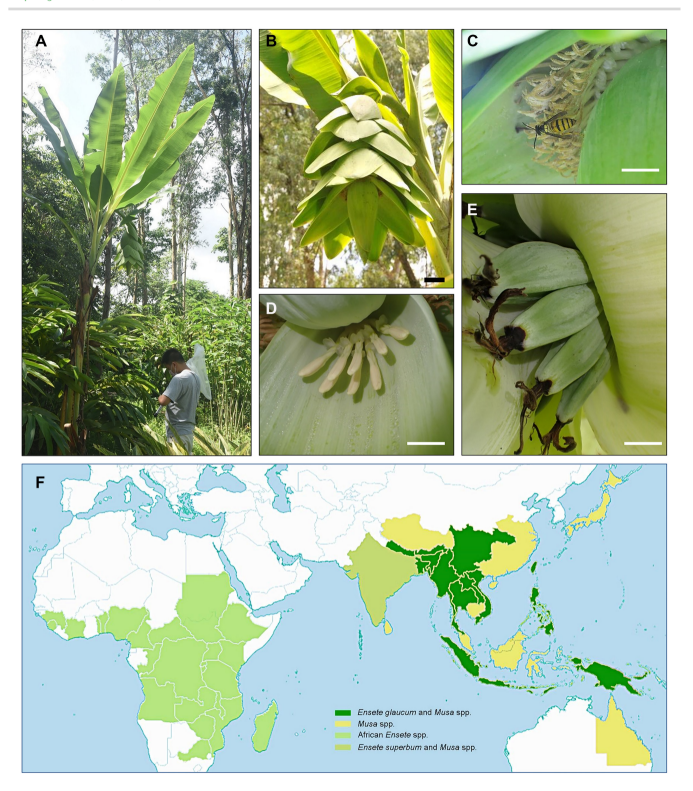


Figure 1: Ensete glaucum plant morphology and distribution map. (A) E. glaucum in South China Botanical Garden, Chinese Academy of Sciences. The pseudo-stem of this plant is ~3.5 m tall. (B) Inflorescence with male and female flowers showing bracts and flowers alternately arranged along the main axis. (C) Staminate flowers, and visiting black shield wasp (Vespa bicolor, Vespidae, Hymenoptera). (D) Female flowers. (E) Fruits. Bars represent 5 cm in B, 2 cm in C and D, and 1 cm in E. (F) Native distribution of Ensete glaucum, E. superbum, Musa, and Ensete species by countries or provinces (for China, India-Assam, and Australia). Musaceae are not currently native in the Americas, although Ensete is present in the fossil record [126]. E. glaucum always occurs in the same provinces as Musa and sometimes with other Asian Ensete species. Map adapted from POWO [62].

in Asia (Fig. 1F) and has records from Burma, China, India, Indonesia, Laos, Myanmar, Vietnam, Philippine, Papua New Guinea, Thailand, and Solomon Islands [11].

Originating in the tropics and subtropics at lower elevations, most species in Musaceae lack cold acclimation. Cold stress is one of the key limitations in extending banana planting and production to higher altitudes and beyond the tropics [12]. In contrast to other Musaceae species, Ensete glaucum can be found above 1,000 m in the mountains of Yunnan in China, where the temperature often drops lower than 0°C, with limited rainfall in winter. As one of the most cold-resistant and perhaps the most drought-tolerant species in Musaceae, E. glaucum is a potential gene and germplasm resource for abiotic stress tolerance in banana breeding, likely to be required for the adaptation to a more variable and extreme climate in the future.

Whole-genome assemblies (genome sequences) are published for some species of Musa with pseudo-chromosome-level data [13-17] using long-molecule sequencing with an N50 of >42 Mb for M. acuminata [17], M. balbisiana [15], and Musa schizocarpa [13]). The assemblies and annotations are available on the Banana Genome Hub, a community website that brings together genomic data, with genome browsers, extensive search facilities, and comparison features [18]. Draft genome assemblies in Ensete species are limited to accessions of E. ventricosum, but these are with tens of thousands of contigs with N50 lengths mostly between 10,000 and 21,000 bp and no pseudo-chromosome assignments [19, 20]. Effective analysis, introduction, and utilization of genetic resources present in wild species of Ensete, based around genome assemblies, are a need for banana improvement and understanding the genome evolution in Musacea. We applied Illumina, Oxford Nanopore Technologies (ONT), and chromosome conformation capture (Hi-C) sequencing to generate a high-quality chromosome-level assembly of the Ensete glaucum genome. We aimed to use the chromosome sequence to show the genome structure and gene composition, as well as revealing the repetitive DNA organization. The structural variations of E. glaucum (x = 9) were studied in a comparative context with Musa (x = 11) species, showing the evolutionary history of the family. The study aims to be useful in expanding the gene pool available not only to banana and enset breeder, but also for plant conservation of biodiversity in ecologically sensitive or threatened areas, and for fundamental research on chromosome and genome evolution.

Analyses, Results, and Discussion

De novo chromosome-scale genome assembly

A *de novo* chromosome-level assembly of *Ensete glaucum* was made by combining high-coverage ONT long-read sequencing, Illumina 150 bp paired-end sequences, and Hi-C chromosome conformation capture sequence data (Table 1). From the initial assembly (with N50 of 10.256 Mb, Table 2 and Supplementary Table S1), we assembled 9 pseudo-molecules, eg01–eg09 (Fig. 2 and Supplementary Table S2), corresponding to the chromosome number (2n = 18) and observed chromosome morphology.

A Hi-C/ONT-only assembly was constructed first, by using an OLC (overlap layout-consensus)/string graph method with corrected reads. Contigs were refined using Illumina short reads, and after discarding redundant contigs, the final genome as-

sembly was 481 Mb long, with 9 pseudo-chromosomes between 42,457,113 and 67,484,389 bp long. BUSCO analysis [21] was used to assess the assembly in "genome" mode showing 98.3% complete single and duplicated Embryophyta core gene sets from the embryophyta_odb10 database (Table 2, Supplementary Table S3a): of the 1,614 genes tested for, 1,526 are complete and single-copy BUSCOs (S), 61 are complete and duplicated BUSCOs (D), 12 are fragmented BUSCOs (F), 15 are missing BUSCOs (M). Few genes were fragmented or missing. *E. glaucum* chromosome designations were chosen to follow major regions of synteny with M. acuminata chromosomes [16, 17].

Genome size, heterozygosity, and organization

The contig-level assembly size is 495,175,598 bp, and 97.2% of these contigs are anchored to 9 pseudo-chromosomes after Hi-C scaffolding, resulting in a 481,507,213-bp final chromosomelevel genome assembly. Some arrays of tandem repeats, including the ribosomal DNA (rDNA) (see below) and telomeres, were collapsed and chromosome termini were not fully assembled. Approximately 55% of the assembled genome was estimated to be repeat sequences (RepeatMasker; Table 2). The genome size was estimated as 563,295,571 bp (highest 17-mer peak frequency). Presumably because of sensitivity of parameters to the evolutionary whole-genome duplications (WGDs) (Fig. 1, centre) and more recent duplications, slightly higher estimates were made by findGSE software (k = 21: 588,939,614 bp; range from k = 17 to 25, 582–591 Mb), and lower estimates by GenomeScope (468,990,370 bp for k = 21; or 407,601,233 bp for k = 17, Supplementary Fig. S1). MGSE [22] gave a mean coverage of reference regions of 62.948-fold and median coverage of 75.00-fold,, corresponding to genome size estimates of 587,786,744 and 493,333,333 bp. The total genome size of E. glaucum (x = 9) is similar to that of the x = 11 Musa species (see [17]) using sequencing methods, and to estimates of both genera by flow cytometry [23].

The heterozygosity rate of E. glaucum was 0.164% (Supplementary Fig. S1 estimated with k = 21 using GenomeScope). Heterozygosity in plants is influenced by mating systems and pollination [24], life span, habitat fragmentation, and cultivation [25]. Relatively little is known about the breeding system and pollination of Ensete species (see [26]), although we observed insects (including the hornet Vespa bicolor, a widespread pollinator in southern China) visiting flowers (Fig. 1C). Our low level of heterozygosity is within the range found in individual plants in populations of M. acuminata ssp. banksii (0.02-0.34% in 24 individuals [27]; and 0.13-0.23% [28]), and in other wild monocotyledonous species including 2 (most likely self-pollinating) diploid oat species (0.07% heterozygosity in Avena atlantica and 0.12% Avena eriantha [29]); it is, however, low compared to other species (e.g., walnut, Juglans nigra 1.0% [30]; Nyssa sinensis 0.87% [31]) and, in particular, many Musa species, some with known hybrid genome composition [28]. The low value seen in species including E. glaucum here is consistent with frequent self-pollination and inbreeding, or a population bottleneck of this monocarpic tropical

Genes were unevenly distributed along chromosomes (Fig. 2 circle b), and generally depleted in broad centromeric regions; few genes were found on the short arm of the more acrocentric chromosome eg04 and the nucleolar organizing region (NOR) bearing chromosome arm of eg06 (see rDNA below). The centromeric, gene-poor regions are rich in repeats (Fig. 2 circle c) and transpos-

Table 1: Statistics of whole-genome sequence assembly and transcriptome analysis of Ensete glaucum using Illumina, ONT, and Hi-C

Туре	Method	No. of reads	Clean data (Gb)	Read length (bp)	Assembly coverage (x)
Genome	Illumina	245,852,534	36.88	2 × 150	74
	ONT	4,357,035	109	38,885 (N50)	220
	Hi-C	319,793,734	48	2 × 150	
Transcriptome	Illumina (Leaf)	62,410,840	94	2 × 150	
	Illumina (Root)	13,712,542	21	2 × 150	

Sizes and coverage are based on the unreplicated haploid genome (1C).

Table 2: Statistics of Ensete glaucum genome assembly and annotation

Genome assembly	Value 563,295,571 bp	
k-mer estimation of genome size (17-mer)		
Total contig length	495,175,598 bp	
Percentage of estimated genome	87.9%	
Anchored into chromosomes	481,507,213 bp	
GC content	38.21%	
Contigs		
Number	124	
N50 length	10,255,891 bp	
Longest	31,226,749 bp	
Pseudo-chromosomes		
Number	9	
Shortest	42,457,113 bp	
Complete BUSCOs of genome	98.3%	
RepeatMasker repetitive DNA		
Transposable elements		
LTR retroelements	37.20%	
Copia	17.64%	
Gypsy	19.25%	
LINEs	0.77%	
Class II DNA transposons	7.18%	
Unclassified dispersed repeats	8.74%	
Simple repeats and low complexity	1.13%	
Total repeats (RepeatMasker)	55.02%	
Tandem repeat content		
45S rDNA	1.21%	
5S rDNA	0.08%	
Egcen satellite	1.32%	
Microsatellites (<8 bp motif)	0.59%	
Protein-coding genes		
Number	36,836	
Average number of exons per gene	4.86	
Average exon length per gene	1,114 bp	
Average intron length	2,816 bp	
Average length of predicted proteins	371 aa	
Complete BUSCOs of predicted genes	94.7%	
Functional annotation	04 500 (05 555)	
NR	31,599 (85.78%)	
InterPro	30,160 (81.88%)	
GO	24,436 (66.34%)	
KO	11,192 (30.38%)	
Total	31,804 (86.34%)	

RepeatMasker did not identify satellite sequences. 5S and 45S rDNA and the centromeric sequence Egcen were identified manually in assemblies and the abundance measured in raw read data; microsatellite abundance was calculated from the assemblies (see Supplementary Tables S1, S2, S3, S10, S12, and S14).

able elements (TEs) (Copia and Gypsy long terminal repeat [LTR] retroelements, Fig. 2 circles d, e, and, less markedly, DNA transposons, Fig. 2 circle f), as observed in many species (including Musa [14, 15]).

The Ks (the synonymous rates of substitution) between genes in paired collinearity gene groups were calculated between E. glaucum and M. acuminata to see whether they share the same 3 WGD events [14]. The 2 genomes have a nearly identical Ks density distribution (Fig. 3A), both having 2 peaks at ~0.55 and ~0.9. This result indicates that Musaceae share the same WGD events. The more recent peak at 0.55 most likely represents the α and β duplications, while the peaks at 0.9 may represent the more ancient γ duplication event [14]. Figure 2 (centre) links the genomic locations of paralogous gene clusters: most chromosome regions show shared relationships with 2 other chromosome regions, reflecting the α and β WGDs, as shown by D'Hont et al. [14] (their Supplementary Fig. 12).

Gene identification

Genes and gene ontology

In total, 36,836 genes were predicted (BUSCO score: C: 94.7%; Supplementary Table S3B) with 31,804 (86.34%) functionally annotated with protein domain signatures and 24,436 (66.34%) associated with GO terms (Table 2; Supplementary Table S4). E. glaucum has a similar gene space (Supplementary Table S5) to the sequenced Musa species M. acuminata (35,264), M. balbisiana (35,148), M. itinerans (32,456), and M. schizocarpa (32,809). In the Musaceae (i.e., M. acuminata, M. balbisiana, M. schizocarpa, E. ventricosum, and E. glaucum), we identified a total of 29,639 orthogroups including 173,025 (88.1%) assigned genes and 23,355 (11.9%) unassigned genes (Fig. 3B and Supplementary Table S5). Between all species, the analysis showed 48% (n = 14,523) of orthogroups were shared (core or softcore genes; increasing to 66%, n = 19,583, if orthogroups missing in only 1 species are discounted as possible annotation artefacts). The analyses highlighted 5% (1,471) of orthogroups that are Ensete genus specific and not found in Musa. A total of 162 orthogroups were found only in E. glaucum (Fig. 3B; lower than the value for E. ventricosum, but the latter is a draft genome status without RNA support and with fragmented contigs with likelihood of a large number of redundant predicted genes). The predicted genes of E. glaucum were compared to their orthologous genes in M. acuminata and Ka/Ks values between orthologous pairs were calculated. Genes with Ka/Ks > 1 were under positive selection (Supplementary Table S6), and GO enrichment was used to summarize the gene functions (Supplementary Fig. S2A), showing that many regulatory biological processes have been positively selected.

Gene family expansion and contraction

Using Musa species and 2 other monocotyledonous species (in the same clade of the Commelinids), Phoenix dactylifera (Arecaceae) and Oryza sativa (Poaceae), we explored gene family expansion and contractions in E. glaucum (Fig. 3C and Supplementary Table S7). Among 12,384 gene families shared by the MRCA (most recent common ancestor) of these monocotyledons, there

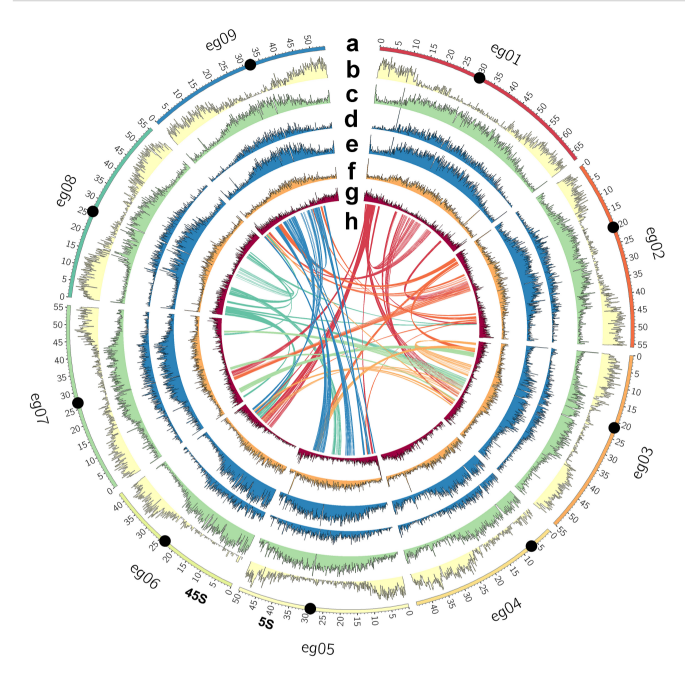
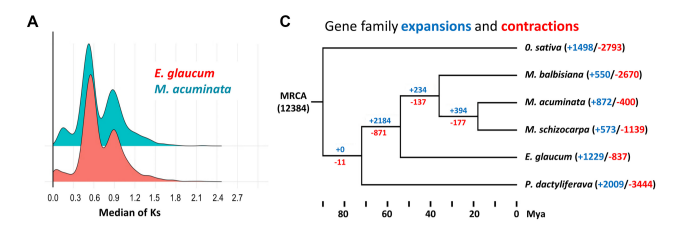
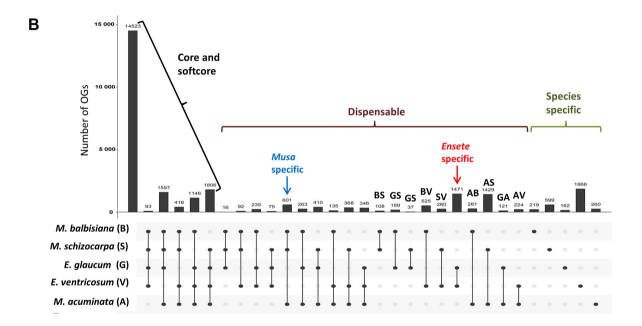


Figure 2: Ensete glaucum chromosome assembly and genome features. Circos plot of (a) The 9 pseudo-molecules (eg01 to eg09) of the EGL assembly corresponding to the 9 chromosomes. 5S and 45S rDNA loci are indicated, and centromere positions are shown by black dots; scale in Mb; (b) gene density; (c) repeat density; (d) Copia LTR retroelement density; (e) Gypsy LTR retroelement density; (f) DNA transposon density; (g) simple sequence repeat (microsatellite) density; (h) Syntenic genomic blocks, linked by curved lines (arbitrary colour) in middle of the plot.

were large numbers of gene families expanding (1,498–2,184) or contracting (817–3,444) between the genomes of Musaceae, *Phoenix*, and *Oryza* (Fig. 3C), presumably reflecting substantial differences in plant form between them. Similar, although slightly lower, figures were reported between, e.g., dicotyledons as diverse as *Arabidopsis* (Brassicaceae), *Solanum* (Solanaceae), and *Cuscuta* (Convolvulaceae) [33]. Notably, though, our results show the largest expansion of gene families in the Musaceae (2,184), likely reflecting the WGD events not shared with the Poaceae or Arecaceae (see also [34] in pineapple), and we find additional expansion in *E. glaucum*. Large gene family losses were noted in *O. sativa*, *P. dactylifera*, and *M. balbisiana* (Supplementary Table S7).

Overall, E. glaucum showed enrichment of several GO biological processes (Supplementary Fig. S2B, Supplementary Table S8) compared to Musa. Among them, "monosaccharide transmembrane transporter" (equal top hit), "carbohydrate transmembrane transport," and "carbohydrate transport"; and among molecular functions, "monosaccharide transmembrane transporter activity," "sugar transmembrane transporter activity," and "carbohydrate transmembrane transporter activity" were all included in the top 20 enrichments. The genus Ensete is notable for its accumulation of starch in the pseudo-stem and leaf bases, with E. ventricosum cultivated as a staple starchy food in East Africa [2], and perhaps this is reflected in the enrichment of certain carbohydrate transport GO terms.





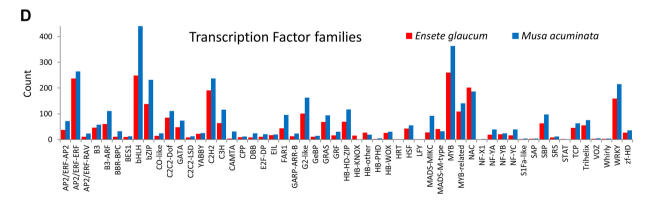


Figure 3: Gene family evolution and conservation. (A) The synonymous substitutions (Ks) frequency density distributions of orthologs within EGL or MAC, whose peaks indicate whole-genome duplications (WGDs). (B) Intersection diagram showing the distribution of shared orthogroups (OGs) (≥2 sequences per OG) among Musa and Ensete genomes. E: E. glaucum; V: E. ventricosum; A: M. acuminata; B: M. balbisiana; S: M. schizocarpa. (C) Gene family expansion and contraction with a phylogenetic tree showing timeline of divergence of monocot species. MRCA: most recent common ancestor. Numbers denote the gene family expansion (red) and contraction (blue). (D) Histogram of the comparative abundance (number of genes) of transcription factors between M. acuminata and E. glaucum.

Transcription Factors

In total, 2,637 putative transcription factor (TF) genes were identified in the E. glaucum assembly, representing 7% of all genes (Supplementary Table S9), which were classified by their signature DNA binding domain into 58 TF families (Fig. 3D). Similar to M. acuminata, the MYB (myeloblastosis) superfamily of TFs (including 260 MYB TFs plus 109 MYB-related) was the largest family, with between 140 and 210 copies of each of the bHLH, AP2/ERF, NAC, C2H2, WRKY, and bZIP families. The identification and classification of the TFs here provides a framework to explore regulatory networks in plants [35] with their target genes and to identify specific factors involved in important responses. Cenci et al. [36] analysed TFs involved in the regulation of tissue development and responses to biotic and abiotic stresses and, particularly, the NAC plant-specific gene family, while Xiao et al. [37] discuss the importance of an HLH factor involved in starch degradation during fruit ripening. Ensete and Musa differ in these characteristics, so it will be interesting to analyse differences in TFs responsible.

Repetitive DNA analysis

Repeat identification

A range of different programs were applied for repeat analysis, and, as has been considered previously [38], there were differences in the repeats identified between approaches, and small changes in parameters and reference sequences give substantial changes. Repeated elements in the genome assembly were identified by RepeatMasker (Table 2 and Supplementary Table S10) and amounted to 55% of the genome assembly, the same range as other plant species with similar DNA amount and, particularly, the genus Musa [14, 15]. For assembly-free identification of repeats, we used RepeatExplorer [39] to generate graph-based clusters of similar sequence fragments: Illumina sequence reads are available from 6 Musaceae species, allowing assembly-free comparisons (Supplementary Table S11 and Supplementary Fig. S3); while there was a little more variation in proportion of reads in the most abundant clusters, all had between 33% and 46% in the top clusters (>0.01% $\,$ genomic abundance, as defined in [37]). TEs including LTR and non-LTR retroelements, and class II DNA transposons, were found (Fig. 4 and Supplementary Table S11). Microsatellites and other repeats were further characterized by mining and dot plot analysis, as well as fluorescent in situ hybridization (FISH) to chromosomes (Supplementary Tables S12-S14 and Figs 5-7; see below). The organization of repetitive regions in the assembly was sometimes verified by mapping individual ONT long reads to assembled repeat regions (e.g., Fig. 5A), and organization was generally confirmed, except for some long tandem arrays that seem to be collapsed in the assembly owing to high homology between repeat units. A few ONT reads were found that included reversals of tandem arrays (head-to-head or tail-to-tail junctions), potentially artefacts from both strands of the DNA molecule passing sequentially through 1 pore, and these junctions need further investigation.

Figure 4A compares the abundance and species distributions of major repeat classes in the Musaceae using the comparative genome analysis function of RepeatExplorer. All species shared many transposons and rDNA sequences (Fig. 4A, central region). However, genus-specific retroelement variants were identified in Musa (Fig. 4A, left) and Ensete-with-Musella (Fig. 4A, right), showing the separation of the 2 phylogenetic branches, supported by extensive divergence of the repetitive sequence subfamilies, and evolution in copy number. Notably, satellite sequences (Fig. 4A centre-right) were much more abundant and some sequences (see centromere sequence below) were present exclusively in Ensete.

Transposable elements

The most abundant class of repetitive elements were TEs, in particular LTR retroelements. The distributions of *Copia* and *Gypsy* LTR retroelements along assembled pseudo-chromosomes (Fig. 2 circles d and e) show greater abundance in proximal chromosome regions. Approximately equal numbers of *Copia* and *Gypsy* elements (18 and 19% of the genome assembly, respectively, Table 2) were found. This result contrasts with *M. acuminata*, where *Copia* elements were considerably more frequent (29%) compared to *Gypsy* elements (11% [14]; Supplementary Table S11). The relative change in proportions of the 2 element families, while the overall abundance remains the same, has implications for genome evolution and the expansion or contraction of retrotransposon families, which can be explored in detail using the high-quality genome sequences where the elements are neither truncated nor collapsed.

Analysis of reverse transcriptase (RT) domains identified subfamilies of LTR retroelements, with the families showing different abundances in E. glaucum and M. acuminata (Supplementary Fig. S4). Insertion times of LTR retroelement subfamilies (Fig. 4B and C and Supplementary Fig. S5) were calculated based on LTR divergence for E. glaucum and recalculated for Musa to allow for identical software settings (see Material, Methods, and Validation). In E. glaucum, both Copia and Gypsy families show relatively constant activity over the past 2.5 Mya, with the major peak of insertion activity 3.5-5.5 Mya (Fig. 4B and C), corresponding to the halflife of LTR-elements [14]. The dynamic amplification of these elements is emphasized by individual subfamilies having bursts of amplification (Fig. 4B for E. glaucum and Supplementary Fig. S5 for Musa), with rounds of expansion of different elements. As shown by Wang et al. [15], M. balbisiana has the most extensive LTR activity in the past 500,000 years, and M. acuminata activity peaks \sim 1.5 Mya (Fig. 4C), in both cases with greater activity of Copia elements (Supplementary Fig. S5), contrasting with E. glaucum with equal activity of both Gypsy and Copia elements, leading to a higher proportion of Gypsy elements within the genome of E. glaucum compared to Musa (see above, and Supplementary Tables S10 and S11). This is also evidenced by the larger number of Musa-specific clusters identified as Copia Angela or Sire elements while Ensete with Musella-specific LTRs include more Gypsy Reina and Retand elements (Fig. 4A). Wu et al. [40] discuss the rounds of amplification in M. itinerans with an amplification burst after separation from M. acuminata ~5.8 Mya suggesting high turnover of the elements. The results suggest a burst of retroelement amplification (the older ones), sometime after the split of Musa and Ensete, and again more recently, perhaps after E. glaucum split from other Ensete species.

Tandem (satellite) repeats and centromeric sequences

The repeat analysis revealed the presence of an abundant tandemly repeated sequence with a monomer length of \sim 134 bp (Fig. 5A). The sequence, named Egcen (Ensete glaucum centromere), represents \sim 1.3% of the E. glaucum genome (45,000 copies) (Table 2, Supplementary Table S12; GenBank: OL310717) and forms arrays that are at places interspersed by the long interspersed nucleotide element (LINE) Nanica (described in M. acuminata [14]) and other sequences (Fig. 5B, see below). There were 1 or 2 major arrays of Egcen repeats found in the assemblies of all 9 chromosomes (Fig. 5C). In situ hybridization of Egcen showed that it was located around the primary, centromeric, constrictions as seen by DAPI (4´,6-diamidino-2-phenylindole) staining (Fig. 5D, see also Fig. 7A and C). The hybridization pattern of the FISH signal on all chromosomes showed variable strength and several sites grouped closely together, corresponding to the pattern seen in the

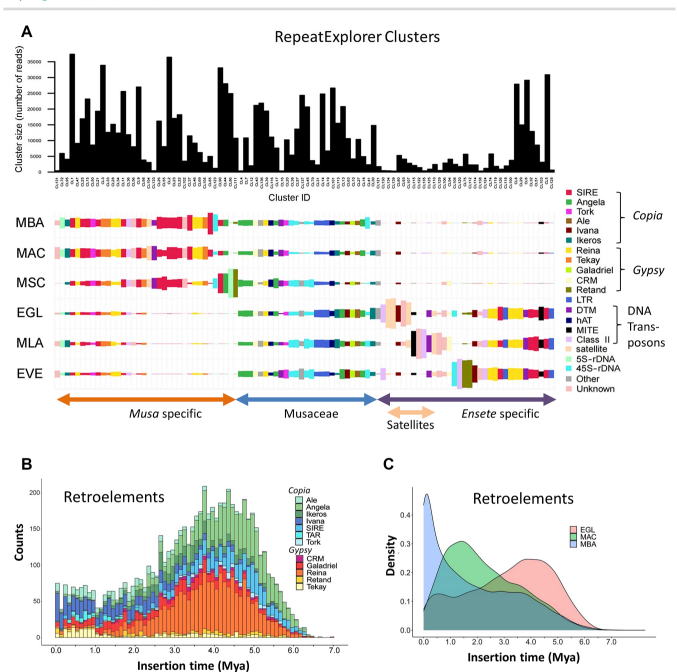


Figure 4: Comparative analysis of repetitive DNA in Musaceae using RepeatExplorer. (A) Bar chart showing the sizes (numbers of reads) of the most abundant individual graph-based read clusters (upper part; black bars) and display of their distribution among 6 Musaceae species (coloured rectangle sizes in lower part proportional to the number of reads in a cluster for each species, based on the annotation of the clusters). Clusters and species were sorted by using hierarchical clustering. EGL: Ensete glaucum; EVE: E. ventricosum; MAC: Musa acuminata; MBA: M. balbisiana; MLA: Musella lasiocarpa; MSC: M. schizocarpa. (B) The distribution of insertion times of LTR retroelements (members of Copia and Gypsy classes) in E. glaucum. (C) The ages of total LTR-retroelement insertions in E. glaucum, M. acuminata, and M. balbisiana. Mya: million years ago.

assembly. The location of the Egcen arrays was therefore used to infer the centromere mid-point position in the *E. glaucum* chromosome assemblies (Fig. 2 outer circle, Fig. 8A, Supplementary Table \$13)

Egcen was also detected at the centromeres of all *E. ventrico-sum* and *Musella lasiocarpa* chromosomes (Fig. 5E and F), showing similar distribution patterns with stronger and weaker signals as in *E. glaucum* (Fig. 5D), but it was not seen on *Musa* chromosomes by in situ hybridization (example of *M. balbisiana*, Supplementary Fig. S6) nor found in analysis of assemblies of *M. acuminata*, *M. balbisiana*, or *M. schizocarpa* (Supplementary Fig. S7A). The compara-

tive RepeatExplorer clustering shows multiple satellite sequences found only in the Ensete and Musella genomes that are not present in the 3 Musa species tested (Fig. 4A) and supports Egcen being part of the tandem repeat birth and amplification that has occurred in Ensete and Musella after the split from Musa, and contrasts with the younger insertion times found for Musa retroelements (Supplementary Table S12 and Fig. 4C).

Tandem repeats or satellite DNA sequences are found around the centromeres of many plant (and animal) species [41, 42] and may be "centromeric" or "pericentromeric." No equivalent tandem repeats were found in Musa [14, 17, 43], and the E. glaucum Egcen is

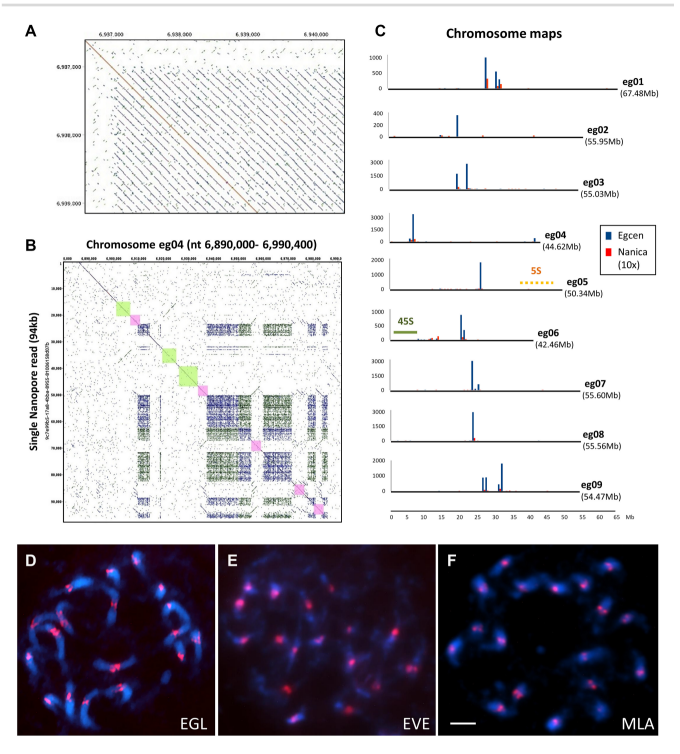


Figure 5: Ensete glaucum centromeric repeat structure. (A) Dot plot (self-comparison of sequences) showing start of a 134-bp Egcen tandem array. (B) Dot plot showing part of a chromosomes assembly (eg04) plotted against part of a single ONT read with blocks of the Egcen tandem repeat (appearing as dense rectangles at this scale) interspersed with Nanica elements (pink; 5 homologous copies in both orientations) and LTR retroelements (green; 2 non-homologous subfamilies). (C) Bar chart showing frequency distribution of the Egcen centromeric tandem repeat, Nanica transposable elements (x10 on axis), and locations of 45S and 5S rDNA along the assemblies for each pseudo-chromosome. Long Egcen arrays occur at 1 or more sites at the centromeric regions of all chromosomes. (D–F) In situ hybridization of Egcen probe detected by red fluorescence to cyan-fluorescing DAPI-stained chromosomes of (D) EGL, Ensete glaucum; (E) EVE, E. ventricosum; and (F) MLA, Musella lasiocarpa. The red Egcen signals collocate with the primary centromeric constriction on all 9 pairs of chromosomes. Bar = 5 μ m.

not present either in Musa (Supplementary Figs S6 and S7A). However, the centromeric regions of all M. acuminata chromosomes have been shown to include multiple copies of a LINE non-LTR retroelement, Nanica, both by in situ hybridization and bioinformatic analysis [14, 17]. Nanica-related sequences were also identified in the E. glaucum assembly but with less abundance than in Musa (Supplementary Fig. S7B); ~350 copies were mapped to chromosomes, mostly (but not exclusively) present interspersed within, or adjacent to, Egcen arrays (Fig. 5B and C and Supplementary Fig. S7B).

Assembly across centromeric regions including abundant repeats is difficult, and normally the tandem repeat elements are collapsed. The ONT long-molecule sequences allowed detailed examination of parts of the centromere region of chromosomes. A dot plot of an ONT read (coded 9c7e99b5, 96,300 bp long) aligned to the assembly of eg04 shows the complex organization of the Egcen array (Fig. 5B): this 100-kb region includes a total of 6 Egcen tandem blocks with between 3 and 126 repeats (a total of 385), 5 copies of *Nanica* (some rearranged, pink boxes), and 3 diverse retroelements flanked by LTRs (green boxes). Further copies of the Egcen tandem repeat occur in larger blocks over the following 450,000 bp of the assembly, and no genes were identified in the region.

A characteristic 17-bp long sequence, the canonical CENP-B box, is found within a monomer of a tandem repeat at centromeres of many species including human [44] and has been postulated to be necessary for binding of the centromeric CENP-B proteins regulating formation of centromere-specific chromatin. Within the Egcen sequence, there was a CENP-B related motif:

Egcen	ggctaaTACGTTGGTAATGGccagt
CENP-B	TNNNTTCGTTGGAAACGGGA
WheatCCS1	TCGGTTGCATGCAGGA
ATcon	CtttgtCTTTGTATCTTCTAACAACAA

Although similar CENP-B motifs have been found from wheat and Brachypodium (CCS1 [45]) to Arabidopsis (ATcon [46]; see also review [47]), the relevance of the CENP-B-related box to centromere function remains uncertain, particularly when no similar tandem repeat is present in other species or the related genera such as Musa (see above). As is the case in many other species, the relative roles of retroelements, tandem repeats, and interspersed centromeric sequences, leading to recruitment of the centromeric proteins, are uncertain: the exact sequence or sequences that mark the functional centromeres remain enigmatic. The identification of a centromeric tandem repeat and the assembly across all centromere regions in E. glaucum together with data from Musa will allow protein-binding studies (with chromatin immunoprecipitation [ChIP] analysis) to resolve the functional centromere.

Microsatellites (simple sequence repeats)

Microsatellites were searched using Phobos and a simple sequence repeat (SSR) mining pipeline ([48]; perfect SSRs from mono to hexa-nucleotide repeats with 11 to 3 repeat numbers, respec-

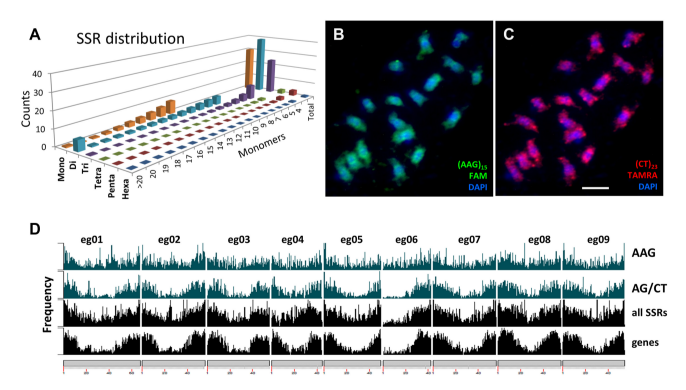


Figure 6: Microsatellite (SSR) distribution in Ensete glaucum. (A) Abundance (count) and total number of monomers of microsatellites (SSR) with motifs between 1 and 6 bp long. (B, C) In situ hybridization of synthetic microsatellite probes to DAPI-stained (blue) chromosomes, showing (B) AAG is relatively uniformly distributed along chromosomes compared to (C) where the greater abundance of AG/CT in distal chromosome regions is seen. (D) Abundance of AAG, AG, all microsatellites, and genes along the chromosome assemblies. In agreement with the in situ hybridization result, AAG is more uniformly distributed, while AG (along with genes and all the microsatellites pooled) show greater abundance in distal chromosome regions except for the arm of chromosome eg06 carrying the 45S rDNA (NOR). Bar = 5 μm.

tively). SSR abundances, array lengths, and nucleotide base composition (70% were AT-rich) are shown in Fig. 6A and Supplementary Table S14. The overall nature and abundances of microsatellites in *E. glaucum* were generally similar to *M. acuminata*, *M. balbisiana*, *M. itinerans*, and *E. ventricosum* (Supplementary Table S14) and reflect that AT-rich microsatellites and dinucleotides (in particular AG/CT) are more frequent in Musaceae, in contrast to GC-rich satellites and trinucleotides being found more often in Poaceae genomes [49].

An average of 1 SSR was found per 4,000 bp, with the density lowest around the centromere and higher at the telomeres (Fig. 6B); they are excluded from the 45S NOR chromosome arm of eg06, and their overall distribution is similar to the distribution of protein-coding genes but contrasts with the more proximal distribution of LTR retroelements and DNA transposons (Fig. 2). Individual microsatellite motifs, however, showed characteristic and different distributions. The abundant microsatellites, CT and AAG, were synthesized as labelled oligonucleotides probes and used as probes for FISH on chromosomes. Both FISH to chromosomes (Fig. 6B and C) and the bioinformatic analysis of the assembly (Fig. 6D) showed that (AAG/CTT) has a relatively uniform distribution along chromosomes, while (AG/CT) shows depletion in centromeric regions and greater abundance in distal parts of chromosomes that are gene-rich (Fig. 2). The constraints on mi-

crosatellite spread in the genome are motif-specific, and, if SSR markers were to be used for genetic mapping, those associated with genes (such as AG/CT) would potentially be more useful.

5S and 45S rDNA and rRNA genes

Tandem repeats of the rDNA were predominantly located within extended, complex loci on chromosomes eg05 (5S rDNA) and eg06 (45S rDNA) (Figs 2, 5C, 7, and 8). The 45S rDNA monomer containing the 18S rRNA gene - ITS1 - 5.8S rRNA gene - ITS2 - 26S rRNA gene - NTS (GenBank: OL310719) was 9,984 bp long, typical but slightly longer than other plant species [50-52]. The NTS region includes in most cases 16 copies of a degenerate 180-bp tandem repeat. On the basis of occurrence in the unassembled Illumina reads, there were 587 copies of the 45S rDNA monomer (1.21% of the genome, Table 2 and Supplementary Table S12). Although in the whole genome assembly the rDNA array was collapsed, the strength of the in situ hybridization signal using the rDNA sequence from wheat (Fig. 7A and B) is consistent with representing 1% of the genome. The long chromosome arm carrying the 45S NOR locus was depleted in protein-coding genes by 10-fold (mean of 12.6 genes/Mb compared with 127.6/Mb on the short arm; Fig. 7D). The single site of 45S rDNA at the NOR per chromosome set is similar to M. acuminata and other Musa species

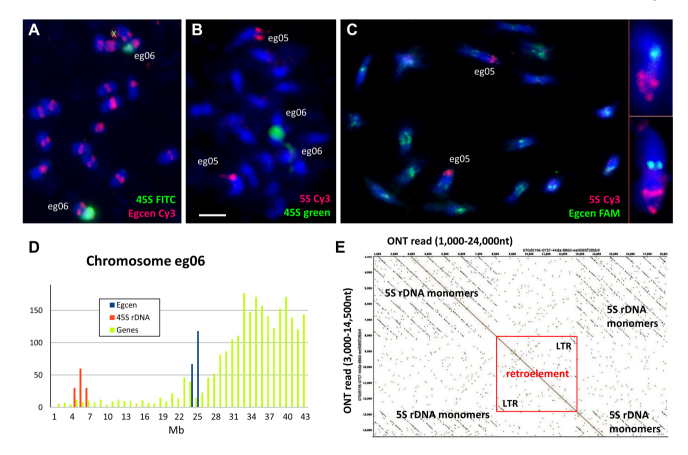


Figure 7: rDNA in Ensete glaucum. (A–C) In situ hybridization to chromosomes (stained blue with DAPI) showing locations of (A) the 45S rDNA (green) on 1 pair of chromosomes (eg06) while Egcen (red) is located at the centromeres of all 9 chromosome pairs. Unspecific signal is marked by x. (B) Chromosomes showing location of 45S rDNA loci (green on eg06; the 2 sites are on 2 chromosomes that are adjacent to each other and the 2 loci have fused); the 5S rDNA loci (red) are located near the end of 1 chromosome pair [eg05]). (C) The 5S rDNA (red on eg05) in a more dispersed pattern with Egcen (green) at all 9 pairs of centromeres; inset shows 5S rDNA chromosomes at higher magnification. The 5S rDNA signal is dispersed over a longer region of the chromosome, while the 45S rDNA locus is dense and occupies much of the chromosome arm. Bar = 5 μ m. (D) Histogram showing density of genes (light green), Egcen (blue), and 45S rDNA copies (red) on chromosome eg06. The arm carrying the 45S rDNA is depleted in protein-coding genes. (E) Part of a single ONT read covering 24 kb spanning part of the 5S rDNA array. The unusually long 1,056-bp tandemly repeated 5S rDNA monomers (14 copies) are interrupted by an LTR retroelement. LTRs, with no homology to the 5S rDNA, are seen (bottom left and top right) in the red box.

[53], although not Ensete gilletii (2n = 18), where there are 4 pairs of sites [23].

The 5S rDNA (GenBank: OL310718) comprised the 5S rRNA gene (119 bp long, typical for all plants; e.g., [51]) and intergenic spacer (937 bp), representing 0.078% of the genome or \sim 366 copies, with a complete motif length of 1,056 bp (Supplementary Table S12). The 5S rDNA locus lies in the middle of the short arm of chromosome eg05, in 3 parts ~34.5M, ~37.5M, and ~45.5M with multiple interruptions. An example of insertion of a 4.7-kb LTR retrotransposonrelated sequence in the 5S rDNA array of tandem repeats is shown in Fig. 7E. In other regions of the E. glaucum ONT reads or assembly, the retroelement-related sequence named Brep, reported in Musa [54], was also found in the 5S rDNA arrays. Garcia et al. [55] show the rather unusual and highly complex structure of 5S rDNA in M. acuminata using graph-based clusters of reads with multiple IGS and retroelement components, supporting the complexity reported here in the E. glaucum assembly. The multiple hybridization sites evident from the in situ hybridization site on 1 pair of chromosomes (Fig. 7B and C), with several, non-continuous, signals visible in the extended prometaphase chromosomes (Fig. 7C, insets), support the non-continuous nature of the 5S rDNA array.

The 5S rDNA monomer length of 1,056 bp was exceptionally long in comparison to any other plant species (typically 400-500 bp long). The first 400 bp of the intergenic spacer had no significant BLAST hits in GenBank, while the second part showed only short regions with weak homology largely to chromosome assemblies of Musa species in GenBank. There were no motifs characteristic of retroelements in the 937-bp intergenic spacer. It is unclear why the monomer length for the 5S rDNA in E. glaucum should be twice that typical in other species, including Musa, and was to be relatively homogeneous over all copies (Fig. 7E). Furthermore, in contrast to the single locus on eg04 of E. qlaucum, all species of Musa examined so far have multiple 5S rDNA sites (2, 3, or 4 per genome), and E. gilletii had 3 pairs of sites [23, 53].

Different species in the Triticeae show wide variation in numbers and locations of both 45S and 5S rDNA sites, suggesting multiple and complex evolutionary rearrangements of the chromosome arms [56] even in the absence of chromosomal rearrangements including translocations and inversions. Dubcovsky and Dvorák [57] have considered the 45S rDNA loci as the "nomads of the Triticeae genomes" given their repeated evolutionary changes in position during species radiation without rearrangements of the genes of the linkage groups. The depletion of protein-coding genes in chromosomal regions extending over most of a chromosome arm around the 45S rDNA genes is notable in Musa and Ensete, so chromosome rearrangements can lead to loci moving, although other recombination, duplication, deletion, or translocation events must occur to alter the numbers of both 5S and 45S loci observed.

Synteny and chromosome rearrangements to Musa

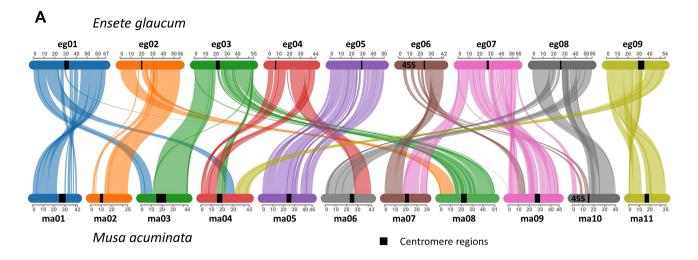
Structural comparisons of the E. glaucum genome assembly (x =9, chromosomes eg01–eg09) were performed with the high-quality assembled genomes of M. acuminata (x = 11, ma01-ma11; v4 [17, 58]) based on synteny (Fig. 8A). The comparison was also extended to M. balbisiana (mb01-mb11 [15]; Supplementary Fig. S8). Sequence dot plots (Fig. 8B) and comparative karyotypes (Fig. 8C and Supplementary Table S15) of the E. glaucum genome against the M. acuminata genome were also analysed.

Overall, the genome assemblies of M. acuminata and E. glau- cum are similar in length and gene content (Supplementary Table S5). We observed high identity between segments of the 9 chromosomes of E. glaucum and of the 11 chromosomes of the Musa (Fig. 8A, Supplementary Fig. S8). Broad centromeric regions with few protein-coding genes (Fig. 2) cannot show syntenic domains. The number of collinear genes was 48,956 between E. glaucum and M. acuminata, and 39,604 between E. glaucum and M. balbisiana (by comparison, the A and B genome of Musa show 42,854 collinear genes). Chromosomes show rearrangements, inversions, expansions, or contractions by crossed, converging, or spreading lines in the Synvisio plots (Fig. 8A). The dot plot (Fig. 8B; single chromosome comparison in Supplementary Fig. S9) shows that there are some syntenic regions distributed over the same length of chromosomes in both species (diagonal lines showing synteny at 45°, eg08/ma10). In other cases, there is expansion in 1 genome and not in the other (lines of synteny more vertical, eg04/ma04, or nearer horizontal, eg03/ma04). Many syntenic segments showed curved lines (ma03/eg03), showing relative expansion of 1 genome at 1 end of the conserved syntenic block, and expansion of the other genome at the other end.

One complete chromosome, ma05/eg05/mb05, was similar with the same gene content in all 3 species (Fig. 8A and Supplementary Fig. S8), but it showed multiple internal inversions and expansions/contractions. A nested pair of inversions was evident covering 10.4 Mb near the start of the chromosome in E. glaucum with respect to M. acuminata (8.84-Mb region) in dotp lots (Supplementary Fig. S9A) and by comparison of locations of orthologous genes (Supplementary Fig. S10). In the context of the E. glaucum inversions, we could also examine the ancestral structure of M. acuminata and M. balbisiana reported by Wang et al. [15]. Notably, a major rearrangement involving an inversion between M. acuminata ma05 and M. balbisiana mb05 [15] was the same inverted region as found in eg05, with an additional nested inversion of 3.1 Mb in eg05 with respect to ma05 (Supplementary Figs S8-S10). Using positions of orthologous genes at the boundaries of syntenic regions, the inversion structure between chromosomes eg05, ma05, and mb05 was clear. Regardless of the ancestral condition, the result indicates that closely similar inversion breakpoints were involved (at the ends of the segment, Supplementary Fig. S10) twice during evolution.

Apart from chromosome 5, an additional 3 whole chromosomes of M. acuminata are represented largely by a single, wholechromosome region/arm of E. glaucum, with some rearrangements occurring within the chromosomes (Fig. 8): ma01 is mainly the right arm of eg01; and ma02 is mainly the right arm of eg02 (Fig. 8A); ma11 is entirely the left arm of eg09 (see details in the dot plot of Supplementary Fig. S9B). The other arms of these 3 E. glaucum chromosomes (eg01, eg02, and eg09) and the remaining 6 chromosomes are related to blocks of the remaining 7 Musa chromosomes. Four Musa chromosomes have translocated fusions of segments of 2 Ensete chromosomes. ma09 has the intercalary region of eg07, with an intercalary segment of eg06 inserted within the eg07 region. ma10 includes parts of 3 Ensete chromosomes, while ma04 has 4 segments from Ensete chromosomes (Fig. 8A and C). The 45S rDNA on eg06 and ma10 are surrounded by syntenic regions but are both depleted in protein-coding genes (Fig. 2). In contrast, the 5S rDNA sites are not surrounded by other orthologous genes (see above). The non-reciprocal translocation noted by Wang et al. [15] of a terminal segment between ma03 and mb01 lies within a larger syntenic block shared between ma03 and eg03. This suggests that the translocation occurred in the M. balbisiana lineage (Supplementary Fig. S8).

In several chromosomes of both E. glaucum and M. acuminata, the breakpoints occur in the centromeric regions (e.g., in eg02,



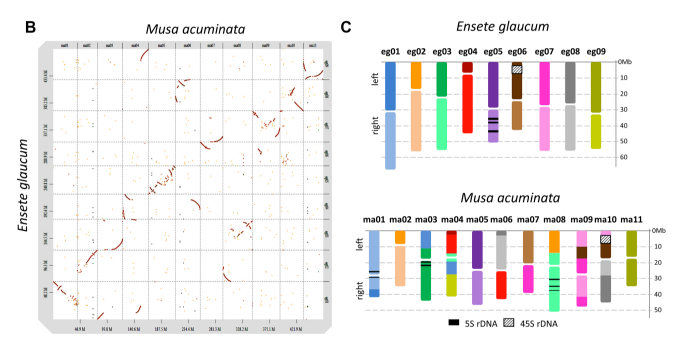


Figure 8: Synteny of Ensete glaucum and Musa acuminata. (A) Synteny plot (Synvisio) connecting syntenic genes in the 9 chromosomes of E. glaucum (egxx) and 11 chromosomes of M. acuminata (maxx). Syntenic blocks of high homology are indicated by uniformly coloured areas in the graphs. Only eg05 and ma05 maintain synteny over the full chromosome length, although there are some rearrangements. Three ma chromosomes are represented by part of 1 eg chromosome, while other ma chromosomes are fusions of > 1 eg chromosome. (B) Dot plot comparing DNA sequences of E. glaucum and M. acuminata (for more detailed dot plots see Supplementary Fig. S9). (C) Representation of the syntenic blocks in the karyotypes of E. glaucum and M. acuminata. Chromosome rearrangements are shown, complementing the Synteny plot, while inversions and relative expansions and contractions of genome regions are clear.

eg03, eg07, eg08, and eg09; ma01, ma06, and ma10; Fig. 8A). While some breakpoints occur at or adjacent to centromeres, the exact relationship of any breakpoint to the centromere and Egcen or Nanica sequences is diverse. Notably, eg03 is spanning the centromere of 3 Musa chromosomes, ma03, ma04, and ma08, and in other cases centromere regions are different despite surrounding synteny (e.g., eg07, ma07, and ma09). Telomeric or subtelomeric regions are conserved between the 2 species in 7 of the 18 E. glaucum chromosome arms (e.g., eg01/ma01, eg06/ma07, eg08/ma10; the dot plot homology lines end in the corners of the chromosomes, Fig. 8B and Supplementary Fig. S9B). In other chromosomes, telomeres in E. glaucum are in intercalary regions of Musa (see eg03/ma03 with inversion, eg08/ma06 in Fig. 8B; and in detail eg08 and ma10 in Supplementary Fig. 9C). The homology of

the whole chromosome ma11 and the left arm of eg09 (see above) indicates a fusion/fission event with loss/gain of centromere and telomere function, but they are also predicted for the other rearrangements discussed above.

Song et al. [10] review the data on the basic chromosome number of the Zingiberales, concluding that x=11 is most reasonable original basic number, with x=9 as a derived basic number. With chromosome numbers of x=9, 10, and 11 predominant in Musaceae, this family is particularly suitable to explore the nature and locations of chromosomal fusions and fissions that are predicted to often occur in similar position during karyotype evolution (e.g., in wheat [59]). The availability of high-quality genome assemblies, based on the ONT, Hi-C, and in Musa BioNano and Pacific Biosciences technologies, will allow the nature of breakpoints

in chromosome fission events to be investigated at the sequence level between the $Musa\ x=11$ and $Ensete\ x=9$ species, as well as being able to shed light on centromere and telomere function.

Conclusions

We provide a chromosome-scale assembly of Ensete glaucum, a sister genus to Musa. This assembly is valuable to infer Musa genome evolution, enabling comparison with putative last common ancestors of M. acuminata (A genome) and M. balbisiana (B genome) at protein and chromosomal levels. Most striking was the multiple rearrangements of chromosome structures between E. glaucum and the Musa A and B genomes, with only 4 of the 11 M. acuminata chromosomes (and only 3 of the 11 in M. balbisiana) showing synteny with only 1 or part of 1 E. glaucum chromosome. With the new insight into chromosome evolution here, further assemblies (in particular the Callimusa section with n=7,9, and 10) will enable resolution of ascending or descending dysploidy, in Musaceae, its sister clades in the Zingiberales, and more widely.

As well as the complex chromosome rearrangements, repetitive sequences differ extensively between the *Musa* and *Ensete* genera. There is a major tandem repeat at the centromeres of only the *Ensete* species, showing lack of conservation of this key structural element of chromosomes, although both genera have multiple copies of the *Nanica* retroelement in centromeric regions. *E. glaucum* has only 1 5S rDNA locus, with an unusually long monomer of 1,056 bp.

The complete sequence provides an accurate reference for the genus for gene identification, marker development, genotyping-by-sequencing, and genome-wide association studies and will accelerate our understanding of the molecular bases of traits such as cold tolerance and starch accumulation and allow identification of relevant genes, contributing to the aim of the Earth BioGenome Project [60] to sequence all eukaryotic species. Although not yet fully understood, the role of chromosomal structural variation and sequence copy number variation (both of genes and repetitive DNA) in genotypic and species diversity is increasingly being recognized, and our high-continuity assembly provides a reference for such studies. The work builds towards a complete pangenome of the Musaceae family, defining structural, gene, and genetic diversity, which can be used for genetic improvement across the Musaceae and more widely.

Material, Methods, and Validation

Sample collection and distribution The individual Ensete glaucum plant used for genome sequencing

The individual Ensete glaucum plant used for genome sequencing and analysis was collected from Puer city, Yunnan province, China and maintained in the South China Botanical Garden, Guangdong province, China (accession No. 19990288; Fig. 1A–E). The distributions of Ensete and Musa species were identified in databases of Flora of China, South China Botanical Garden, iNaturalist, GBIF [61] (excluding cultivation sites), and regional distribution maps. Figure 1F was then made from POWO [62] (overlaid and colouradjusted in Adobe Photoshop CC2018).

DNA extraction and sequencing

Young leaves of Ensete glaucum were collected and ground into powder in liquid nitrogen. High molecular weight genomic DNA was extracted using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany). DNA quality was assessed by agarose gel electrophore-

sis and NanoDrop 2000c spectrophotometry, followed by Thermo Fisher Scientific Qubit fluorometry.

Illumina sequencing

A genomic DNA library with 400-bp fragments was constructed using Truseq Nano DNA HT Sample preparation Kit (Illumina, USA), and 150-bp paired ends were sequenced with Illumina Novaseq (Illumina NovaSeq 6000 Sequencing System, RRID:SCR_020150) by Grandomics Biosciences Co., Ltd. (Wuhan, Hubei, China) (previously known as Nextomics, Wuhan, Hubei, China). After applying Trimmomatic v0.36 (Trimmomatic, RRID:SCR_011848) [63] to trim adaptors, filtering out low-quality reads and further quality control with fastQC v0.11.9 (FastQC, RRID:SCR_014583) [64], 246 million paired reads and 36.88 Gb of data resulted (Table 1).

Oxford Nanopore sequencing

ONT (Oxford Nanopore Technologies, Oxford, the UK) sequencing was performed by Grandomics Biosciences Co., Ltd. (Wuhan, Hubei, China): long fragments longer than 12 kb were selected with Sage Sciences BluePippin (Sage Science BluePippin system, RRID:SCR_020505), and the SQK-LSK109 kit (Oxford Nanopore, Oxford, the UK) was used to build a library that was sequenced using PromethION (PromethION, RRID:SCR_017987), flow cell R9.4.1. The base calling was performed with Guppy v2.0.8 and reads mean_q score_template (Phred) > 7 (base call accuracy >80%) were selected. A total of 129 Gb ONT reads (~250× coverage) was generated. fastp v0.19.7 (fastp, RRID:SCR_016962) [65] was used for quality control including adaptor-trimming, filtering reads with too many Ns or mean q score <7, and resulted in remaining clean data of 109 Gb (Table 1). The mean read length was ~20 kb, with the longest >120 kb (Supplementary Fig. S11).

Hi-C chromatin interaction data

Genomic DNA was extracted from E. glaucum for Hi-C analysis and generation of a contact map to anchor contigs onto chromosomes [66, 67]. First, freshly harvested leaves were cut into 2cm pieces and vacuum infiltrated in nuclei isolation buffer supplemented with 2% formaldehyde. Crosslinking was stopped by adding glycine and additional vacuum infiltration. Fixed tissue was then ground to a powder before resuspending in nuclei isolation buffer to obtain a suspension of nuclei. The purified nuclei were digested with 100 units of DpnII and tagged with biotin-14dCTP. Biotin-14-dCTP from non-ligated DNA ends was removed owing to the exonuclease activity of T4 DNA polymerase. The ligated DNA was sheared into 300-600 bp fragments and then was blunt-end repaired and A-tailed, followed by purification through biotin-streptavidin-mediated pull down. Finally, the Hi-C libraries were quantified and sequenced using the Illumina HiSeq platform (performed by Grandomics). Low-quality sequences (quality scores <20), adaptor sequences, and sequences shorter than 30 bp were filtered out using fastp v0.19.7 [65].

Genome and chromosome assembly

ONT data were corrected by Nextdenovo v2.0-beta.1 [68], with setting "read_cutoff = 3k, seed_cutoff = 25k, blocksize = 2g" and the 109-Gb filtered data were assembled by SMARTdenovo (SMARTdenovo, RRID:SCR_017622) [69] with the parameters "wtpre -J 3000, wtzmo -k 21 -z 10 -Z 16 -U -1 -m 0.1 -A 1000, wtclp -d 3 -k 300 -m 0.1 -FT, wtlay -w 300 -s 200 -m 0.1 -r 0.95 -c 1". To polish the assembly, contigs were refined with Racon (Racon, RRID:SCR_017642) [70], BWA v0.7.17 (BWA, RRID:SCR_010910) [71] was used to map the

filtered Oxford Nanopore reads to the assembly, and NextPolish v1.3.1 [72] with parameters "-consensus -w window -t 4 -m 0.5 -d 30" was used to discard possibly redundant contigs and generate a final assembly; similarity searches were performed with the parameters "identity 0.8 - overlap 0.8". Finally, BWA v0.7.17 [71] and Pilon v1.21 (Pilon, RRID:SCR_014731) [73] with setting "-changes -vcf -diploid -fix bases -threads 10 -mindepth 10" were used to further correct the assembly using the Illumina Novaseq reads, and 2 rounds of mapping back to the assembly each time with further correction were undertaken. A 494 Mb assembly with 124 contigs was achieved (Table 2 and Supplementary Tables S1 and

Read pairs from the Hi-C data were mapped to the draft assembly using bowtie2 v2.3.2 (bowtie2; RRID:SCR_016368) [74] with the settings "-end-to-end, -very-sensitive and -L 30" to select unique mapped paired-end reads. Valid interaction paired-end reads were identified by HiC-Pro v2.8.1 (HiC-Pro, RRID:SCR_017643) [75] and retained for further analysis while invalid read pairs, including dangling-end, self-cycle, re-ligation, and dumped products were discarded. The scaffolds were further clustered, ordered, and oriented onto pseudo-chromosomes by LACHESIS (LACH-ESIS, RRID:SCR_017644) [76], with parameters as follows: "CLUS-TER_MIN_RE_SITES = 100, CLUSTER_MAX_LINK_DENSITY = 2.5, CLUSTER NONINFORMATIVE RATIO = 1.4, ORDER MIN N RES IN TRUNK = 60, ORDER MIN N RES IN SHREDS = 60". Finally, regions with obvious discrete chromatin interaction were detected and their placements and orientations were manually adjusted (Supplementary Fig. S12). Validation of the assembly was performed using BUSCO v5 (BUSCO, RRID:SCR_015008) [21] to assess the completeness and presence of 1,614 genes in the embryophyta_odb10 database in "genome" mode (Supplementary Table S3a).

RNA extraction, sequencing, and transcriptome assembly

Total RNA was extracted from fresh leaves and roots of the same individual of E. glaucum that was used for genomic sequencing using RNeasy Plant Mini Kit (Qiagen, Shanghai, China). Illumina libraries were built from 1 μg total RNA of each sample with TruSeq RNA Library Preparation Kit (Illumina, USA) and were then sequenced using Illumina Novaseq platform to generate paired-end reads. A transcriptome assembly was produced using RNA-seq data using Trinity (Trinity, RRID:SCR_013048) [77] with parameters: "-genome_guided_bam EGL.star.bam -max_memory 50G -genome_guided_max_intron 10000" and mapped on the genome with PASA (PASA, RRID:SCR_014656) [78] with setting: "-MIN_PERCENT_ALIGNED = 80 -MIN_AVG_PER_ID = 80".

Genome size estimation

Using the Illumina DNA sequence, genome size was estimated from the 17-mer frequency using Jellyfish v2.0 (Jellyfish, RRID:SCR_005491) [79] with the formula k-num/k-depth (where k-num is the total number of 17-mers, 30,417,960,841; and kdepth the highest k-mer depth, 54; Table 2). Then 21-mer data were used in findGSE [80] and Genomescope 2.0 (Genomescope R, RID:SCR_017014) [81] to estimate the genome size and heterozygosity (Supplementary Fig. S1).

MGSE v0.4 [22] were also used to estimate the genomic size based on read-mapping coverage. The next-generation sequencing reads were mapped to assembly by BWA v0.7.17. The coverage of single-copy genes (BUSCO genes) was calculated by MGSE.

Gene annotation

We adopted a combination of ab initio gene prediction, homologybased gene prediction, and transcriptome-based gene prediction strategy. RepeatMasker v4.0.9 (RepeatMasker, RRID:SCR_012954) with option "-no_is -xsmall" was used to generate a repeat softmasked genome file. RNA-seg data from leaf and root tissues were mapped to the masked genome assembly with STAR v2.7 (STAR, RRID:SCR_004463) [82] with option: "-outSAMtype BAM SortedByCoordinate -outSAMstrandField intronMotif outFilterIntronMotifs RemoveNoncanonical". The RNA alignment was input into BRAKER2 v2.1.5 (BRAKER, RRID:SCR_018964) [83], a combination of GeneMark (GENEMARK, RRID:SCR 011930) [84] and AUGUSTUS (RRID:SCR_008417) [85], to perform ab initio gene predictions with the default settings. The gene models from BRAKER2 were input into MAKER v2.31.10 (MAKER, RRID:SCR_005309) [86] as model, and the RNA alignment of E. glaucum and proteins from M. acuminata v2 were used as Expressed Sequence Tag (EST) and protein evidence, respectively. We also used GeMoMa v2.3 (GeMoMa, RRID:SCR_017646) [87] to perform homology-based gene prediction using M. acuminata v2 [16] as reference annotated genome. Evidence-Modeler (EvidenceModeler, RRID:SCR_014659) [88] was used to combine de novo and homology-based predictions and our transcriptome evidence to produce the final structural gene

To annotate the function of predicted genes, we performed BLASTP (e-value = 1e-10) (BLASTP, RRID:SCR 001010) from the BLAST+ package [89] for each predicted coding sequence against the databases: UniProtKB/Swiss-Prot, UniProtKB/TrEMBL [90], and NR (non-redundant protein database at NCBI). These sequences are then processed to produce a non-identical (often referred to as pseudo non-redundant) prediction. To assign a putative function to a polypeptide we kept only the best hit on the basis of 3 parameters: (i) Qcov (Query coverage = length high-scoring segment pair [HSP]/length query), (ii) Scov (Subject coverage = length HSP/length subject), and (iii) identity. Additional functional information was added by scanning sequences with InterProScan v5.46 (InterProScan, RRID:SCR_005829) [91]. Blast2GO v6.0.1 (Blast2GO, RRID:SCR_005828) [92] was used to integrate the results of BLAST and InterProScan, and to link the GO (Gene Ontology) terms to genes accordingly (Supplementary Table S4). The functional annotation procedure is given in greater detail at [93].

BUSCO was run in mode "transcriptome" using the embryophyta_odb10 database to assess the gene annotation results and found 1,529 (94.7%) complete BUSCOs (Supplementary Table S3b).

Gene family analyses

Orthogroups identification in Musaceae

Protein-coding genes from M. acuminata [16], M. balbisiana v1.1 [15], and M. schizocarpa v1 [13] were retrieved from the Banana Genome Hub [18]. Protein-coding genes predicted from E. ventricosum were downloaded at NCBI Genome (GCA_000818735.3) to allow discrimination of Ensete-specific orthogroups and E. glaucumspecific orthogroups. Combined with E. glaucum protein-coding genes, we used OrthoFinder v2.5.2 (RRID:SCR_017118) [94] and Diamond [95] with default parameters (summary in Supplementary Fig. S5). Visualization (Fig. 3B) was produced with UpsetR [96]. Gene ontology (GO) enrichments were calculated using TopGO [97] with Fisher exact test (Supplementary Fig. S2 and Supplementary Table S8).

Gene family expansion and contraction

To identify gene family expansion and contraction, we expanded previous analyses with OrthoFinder by adding a representative of Musaceae sister clades in Palms (P. dactylifera, date palm [98]) and Poales (O. sativa v7, rice [99]; data downloaded from Phytozome [100]) but omitting E. ventricosum owing to gene redundancy. The longest transcripts were kept if alternative splicing occurred. Divergence time estimation with approximate likelihood calculation used MCMCTREE in PAML v4.9j (PAML, RRID:SCR 014932). CAFE (CAFE v4.2.1, RRID:SCR_018924) [101] was used to model the evolution of gene family sizes and stochastic birth and death processes and summarized in the phylogenetic tree (Fig. 3C).

Transcription factors

Protein-coding gene sequences for E. glaucum and M. acuminata v2 were searched in PlantTFDB v5.0 (PLANTTFDB, RRID:SCR_003362) and iTAK online v1.6 [102]. Predicted TFs were verified through a hidden Markov model (HMM) with PFAM searching tools using the cut-off E-value of 0.01 (Fig. 3D and Supplementary Table S9). Genes were verified by PFAM (Pfam, RRID:SCR_004726), CDD (Conserved Domain Database, RRID:SCR_002077), and SMART (SMART, RRID:SCR_005026) databases.

Whole-genome duplication

To identify the WGD events, we applied WGDI pipeline (wholegenome duplication identification v0.4.7 [103]). The predicted proteins of E. glaucum were blasted against themselves and then a collinearity analysis was conducted. The Ks (the synonymous rates of substitution) between genes in paired collinearity gene groups were calculated and the Ks peak was detected. For comparison, the same processes were also applied to M. acuminata v2 [16].

Synteny analyses

Structural comparisons of the E. glaucum genome were performed with M. acuminata v4 (designated the A genome) and M. balbisiana (B genome). The recent release of M. acuminata v4 assembly was preferred in this case because it improved pericentromeric regions and provided telomere-to-telomere gapless chromosomes [17]. Assemblies were aligned with minimap2 (Minimap2, RRID:SCR_018550) [104] and visualized results using D-Genies (D-GENIES, RRID:SCR_018967) v1.2.0 [105]. Protein-coding genes were processed to identify reciprocal best hits with BLASTP (e-value 1e-10) followed by MCScanX (e-value 1e-05, max gaps 25) [106] and results imported in SynVisio [58] for syntenic block visualization. Scale bars and colouring of the chromosome bars was adjusted using Adobe Photoshop CC2018.

The karyotype of E. glaucum in Fig. 8C was prepared from lengths of each pseudo-chromosome (Supplementary Table S2) with the estimated centromere position (using the Egcen array midpoints, Supplementary Table S13) to estimate the left (darker coloured) and right (lighter coloured) chromosome arms. Chromosome lengths and centromere positions for M. acuminata were taken from [17] (Fig. 2a); syntenic blocks were calculated using the SynVisio diagram (Fig. 8A).

Repetitive DNA identification and annotation

For repetitive DNA analysis, publicly available programs (see below and [38]) as well as manual searches and sequence comparisons were applied. Geneious v.10.2.6 (Geneious, RRID:SCR_010519) (Biomatters Ltd., Auckland, New Zealand) was used to produce the dotp lots of Figs 5A, B, 7E, and 8B, and Supplementary Fig. S9).

In the assembly, repeated sequences were first searched with REPET v2.5 pipeline [107]. The top 100 repeated sequences were plotted on the 4 reference Musa genome assemblies (i.e., M. acuminata, M. balbisiana, M. schizocarpa, and E. glaucum) using BlastAndDrawDensity.py script described in [17] and available on the GitHub repository [108].

Transposable elements

TEs were annotated by EDTA pipeline [109], which integrates various software to discover TEs including LTR retrotransposons [110-112], terminal inverted repeat (TIR) transposons [113], short TIR transposons or miniature inverted transposable elements (MITEs) [114], and Helitrons [115]. According to suggestions in [109], we also adopt RepeatModeler2 (RepeatModeler, RRID:SCR_015027) v2.0.1 [116] to find remaining TEs.

We also discovered repetitive elements through the REPET v2.2 [117] package including TEdenovo and TEannot. The TEdenovo procedure was used on masked assembly to produce a batch of 4,229 TE consensus sequences. From these 2,800 consensus sequences, only those with full-length fragments present in the assembly were kept for further analysis, quantification, and annotation with the TEannot procedure. A first annotation was performed using public Repbase (Repbase, RRID:SCR_021169) release 20.05, followed by Gypsy/Copia retroelement family identification using HMMs (hmmsearch version 3) to search consensus for corresponding retro-transposase PFAM domains PF04195 and PF14244, respectively. The above results were then combined and CD-HIT v4.1.8 (CD-HIT, RRID:SCR_007105) [118] was used to reduce redundancy. The LTR retrotransposons were sent to TEsorter [119] to classify into lineage level and RT domain amino acid sequences were extracted. Phylogenetic trees of Copia and Gypsy were inferred by RT domain alignment results (Supplementary Fig. S4). The proportions of TEs in the assembly are given in Table 2 (Supplementary Tables S11 and S12) and chromosomal distributions in Fig. 2.

To estimate ages of LTR retrotransposons and the time of insertion (Fig. 4B and C and Supplementary Fig. S5), complete elements were found by LTRharvest v1.6.1 (LTRharvest, RRID:SCR_018970) [110] and LTR_retriever (LTR_retriever, RRID:SCR_017623) [111] and then classified by TEsorter v1.3. The estimation of time was based on the divergence of the 5'- and 3'-end LTRs, and these 2 LTRs of every LTR retrotransposon were extracted into separate files with a custom script. The 5' and 3' LTRs were aligned by MUSCLE v3.8.1551 [120]. The divergence distances under K2P evolutionary model were calculated by R package ape v5.4-1 (ape, RRID:SCR_017343). The average base substitution rate was selected to be 11.3E-8 [121]. The insertion time T was calculated as T = K/(2r), with r as the rate of nucleotide substitution and K as the divergence distance between LTR pairs. The script to perform the analysis is on GitHub [122].

Graph-based clustering of reads using RepeatExplorer

A sample of 2 Gb of the Illumina HiSeq raw reads were used for assembly-free analysis by RepeatExplorer2 [39]. Graph-based clusters of similar sequence fragments were generated under default parameters. Clusters were assigned to repeat classes and retroelement lineages using the automated Repeat Masker and Domain hits provided by the program (Supplementary Table S11). Comparative analysis with sample Illumina sequence reads from 5 other Musaceae species, namely, M. acuminata v2 [16], M. balbisiana v1.1 [15], and M. schizocarpa v1 [13], E. ventricosum [20], and

M. lasiocarpa (Wang, Cui, Rouard, Schwarzacher, Heslop-Harrison, Liu in preparation)) were also analysed (Supplementary Fig. S3) and compared with RepeatExplorer2 following "comparative repeat analysis" protocol. The results were visualized by R script "plot comparative clustering summary.R" (Fig. 4A).

SSR tandem repeats

The genome assembly was searched for SSR (microsatellite) motifs using the SSR mining pipeline developed by Biswas et al. [48]. Searches were standardized for mining perfect SSRs from monoto hexanucleotide repeats (minimum repeat number of 12 for mononucleotides, 8 for di-, 5 for tri-, tetra-, and penta-, and 4 repeats for penta- and hexanucleotides). SSR abundance and nature was analysed on the basis of density in the genome (~1 per 4,000 bp), array length (Fig. 7B and Supplementary Table S14; SSR search parameters minimum lengths mono = 1*12 = 12nt, di = 8*2 = 16nt, tri = 3*5 = 15nt, tetra = 4*5 = 15nt, penta = 5*4 = 15nt, penta = 5*4 = 15nt, tri = 15nt, tri = 15nt, penta = 15n 20nt and hexa = 6*4 = 24nt; total SSR count 123884; Class I>20nt and Class II≤20nt), nucleotide base composition of the SSR loci (70% were AT-rich), and abundance of each motif.

Fluorescent in situ hybridization

Chromosome preparation and FISH were performed as described by Schwarzacher and Heslop-Harrison [123] with minor modifications. Plants of E. glaucum, E. ventricosum, M. lasiocarpa (purchased commercially), and M. balbisiana "Butuhan" (ITC1074) [124] were grown in the glasshouse at the University of Leicester, UK. Actively growing root tips were treated with 2 mM 8-hydroxyquinoline and fixed with 96% ethanol:glacial acetic acid (3:1). For chromosome preparations, roots were digested with a modified enzyme solution (32 U/mL cellulose, Sigma-Aldrich C1184; 20 U/mL "Onozuka" RS cellulose; 35 U/mL pectinase from Aspergillus niger, Sigma-Aldrich P4716; 20 U/mL Viscozyme, Sigma-Alderich V2010) in 10 mM citric acid/sodium citrate buffer (pH 4.6) for 3-5 h at 37°C and then kept in buffer for 12–30 h at 4°C. Meristems were dissected in 60% acetic acid and routinely 2-6 slide preparations were made from each root. Slides were stored at -20° C until FISH.

The 45S rDNA probe was labelled by random priming (Invitrogen) with digoxigenin dUTP or biotin dUTP (Roche) using the linearized clone pTa71 [125] containing the 45S rDNA repeat unit of Triticum aestivum. A 50-100 ng quantity of labelled probe was used per slide and detection of hybridization sites was carried out with fluorescein-conjugated anti-digoxigenin (Roche) or streptavidin conjugated to Alexa-647 (Molecular Probes, Invitrogen). The remaining probes were designed from the consensus sequence of the centromeric repeat Egcen (Fig. 5) and the 5S rDNA (Fig. 7) or as SSRs (Fig. 6); as directly labelled oligonucleotides (200-500 ng per slide) they needed no further detection and were as follows:

- CenCy3: EGL_2640R: [Cyanine3]GAC CGT CGC ATT TTT TGG CGA AAC CAT GCT CGT ACG ACT TCC CAT GGG CTA AAA CGT TAG GA
- CenFAM: EGL_G2640L: [6FAM]GGC CTA TAT TTT GAA ATT CCG AGA CGG TGC ATG AAA AAC CGA TCG AAA CGA AAC ATT GCG
- 5S_4M_Cy3: [Cyanine3]TCA GAA CTC CGA AGT TAA GCG TGC TTG GGC GAG AGT AGT AC
- 5S_3R_Cy3: [Cyanine3]AGT ACT AGG ATG GGT GAC CCC CTG GGA AGT CCT CGT GTT GC
- 5S_6L_Cy3: [Cyanine3]GCG ATC ATA CCA GCA CTA AAG CAC CGG ATC CCA TCA GAA CTC C
- (AAG)15_FAM: [6FAM]AAG AAG AAG

(CT)23_TAMRA: [TAMRA]CTC TCT CTC T

For hybridization, probes were prepared in 40% (v/v) formamide, 20% (w/v) dextran sulphate, 2× SSC (sodium chloride sodium citrate), 0.03 µg of salmon sperm DNA, 0.12% SDS (sodium dodecyl sulfate), and 0.12 mM EDTA (ethylenediamine-tetra acetic acid). Chromosomes and 40-50 µL of probe mixture were denatured together at 72°C for 8 mins, cooled down slowly, and allowed to hybridize overnight at 37°C. Post-hybridization washes were at 42°C in 0.1× SSC, giving a stringency of 80–85% for the short oligo probes and 70-75% for the 45S rDNA probe. Chromosomes were counterstained with 4 µg/mL DAPI and mounted in CitifluorAF. Slides were examined using Nikon Eclipse 80i microscope and images were captured with a DS-QiMc monochrome camera, and NIS-Elements v2.34 (Nikon, Tokyo, Japan) assigning colour and merging channels. Overlays of hybridization signal (shown in green or red) and DAPI images (in cyan or blue) were enhanced with Adobe Photoshop CC2018 using only cropping and functions that treat all pixels of the image. Seven FISH runs with different combinations of probes and replicates were performed, and between 5 and 15 metaphases per slide (99 metaphases in total from 15 slides) were analysed in detail.

Data Availability

All supporting data can be found in the GigaScience database [127]. Raw sequence reads (RNA-seq, Illumina HiSeq, ONT, and Hi-C) were deposited in NCBI under accession No. PRJNA736572. Specifically, ONT raw reads: SRX11350424 and SRX11350425; RNA-seq raw reads, as follows: leaf: SRX11350426; root: SRX11350427; genomic Illumina short-read data: SRX11350423; raw reads of the Hi-C library: SRX11350428 and SRX11350429. The raw read data were also deposited in the Genome Sequence Archive (GSA) of the China National Center for Bioinformation (accession code: CRA004283).

The assembled genome was also deposited to GenBank in NCBI and can be accessed via accession No. JAHSUZ000000000. Genome assembly, gene and TE annotation data, and transcriptomic data are also available on the Banana Genome Hub (http://banana-gen ome-hub.southgreen.fr/) for download or exploration via a dedicated Genome Browser (Jbrowse) and syntenic browser (SynVisio).

Additional Files

Supplementary Figure S1. Genomescope analysis of heterozygos-

Supplementary Figure S2. GO enrichment terms.

Supplementary Figure S3. RepeatExplorer clustering summary in Musaceae species.

Supplementary Figure S4. LTR retroelement trees EGL and MAC. Supplementary Figure S5. Gypsy and Copia insertion times in Musa and E. glaucum.

Supplementary Figure S6: Egcen FISH to Musa chromosomes. Supplementary Figure S7. EgCen and Nanica in assemblies of E. glaucum and Musa.

Supplementary Figure S8. Synteny of E. glaucum with Musa A and B genome.

Supplementary Figure S9. Dot plots of individual chromosomes. **Supplementary Figure S10.** Inversions on chromosome 5.

Supplementary Figure S11. Length distribution of ONT reads.

Supplementary Figure S12. Hi-C interaction contact map.

Supplementary Table S1. Contig statistics based on assembly of ONT sequencing data.

Supplementary Table S2. Chromosome lengths and number of contigs anchored in Ensete glaucum assembly.

Supplementary Table S3. Quality assessment of the gene annotation of Ensete glaucum using BUSCOs v5.

Supplementary Table S4. Complete gene list: homology and GO. **Supplementary Table S5.** Statistics for shared orthogroups (OG) and gene clustering among E. glaucum, E. ventricosum, Musa acuminata, M. balbisiana, and M. schizocarpa genomes.

Supplementary Table S6. Positively selected genes and their annotation

Supplementary Table S7. Result of gene family size change anal-

Supplementary Table S8. (a) Top 20 GO molecular function enrichments for E. glaucum and shared E. glaucum/E. ventricosum gene families; (b) Top 20 GO biological pathway enrichments for E. glaucum and shared E. glaucum/E. ventricosum gene families.

Supplementary Table S9. Comparison of transcriptional factors between Ensete glaucum and Musa acuminata.

Supplementary Table S10. Transposable elements and other repeat proportions comparison in assembly (RepeatMasker).

Supplementary Table S11. Repeat content (RepeatExplorer) comparison between different Musaceae genomes.

Supplementary Table S12. Abundance of major tandemly repeated DNA repeats in Illumina raw reads.

Supplementary Table S13. Inferred centromere positions from locations of interrupted tandem arrays of the Egcen centromeric sequence on the chromosome assemblies.

Supplementary Table S14. Comparative survey of microsatellite sequences in Ensete glaucum genome with other sister species.

Abbreviations

BLAST: Basic Local Alignment Search Tool; bp: base pairs; BUSCO: Benchmarking Universal Single-Copy Orthologs; BWA: Burrows-Wheeler Aligner; CAFE: Computational Analysis of gene Family Evolution; CTAB: cetyl trimethylammonium bromide; DAPI: 4´,6-diamidino-2-phenylindole; Egcen: Ensete glaucum centromere sequence; FISH: fluorescence in situ hybridization; Gb: gigabase pairs; GC: guanine-cytosine; GeMoMa: Gene Model Mapper; GO: gene ontogeny; Hi-C: high-throughput chromosome conformation capture; HMM: hidden Markov model; ITS: internal transcribed spacer of rDNA; kb: kilobase pairs; KEGG: Kyoto Encyclopedia of Genes and Genomes; LACHESIS: Ligating Adjacent Chromatin Enables Scaffolding In Situ; LINE: long interspersed nucleotide element; LTR: long terminal repeat; Mb: megabase pairs; ML: maximum likelihood; Mya: million years ago; NCBI: National Center for Biotechnology Information; NR: RefSeq non-redundant proteins; NOR: nucleolar organizing region; NTS: non-transcribed spacer of rDNA; ONT: Oxford Nanopore Technologies; PAML: Phylogenetic Analysis by Maximum Likelihood; PASA: Program to Assemble Spliced Alignments; rDNA: ribosomal DNA; RNA-seq: RNA sequencing; RT: reverse transcriptase; SRA: Sequence Read Archive; SSR: simple sequence repeat; TE: transposable element; TF: transcription factor; WGD: whole-genome duplication.

Consent for Publication

The origin of E. glaucum plants is given in Materials and Methods. They were collected in China and conserved in the South China Botanical Garden, Chinese Academy of Sciences, with appropriate agreements. No live material was exported out of the country. Other plants for chromosome preparations were obtained from the International Transit Centre, ITC genebank, with official Standard Material Transfer Agreement (SMTA), acknowledged in the manuscript.

Competing Interests

The authors declare that they have no competing interests.

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Authors' Contributions

Q.L. and J.S.H.H. designed the project and with M.R. and Z.W. contributed to project coordination. Z.W. and Q.H. collected samples and conducted DNA and RNA extraction. T.S. and J.S.H.H. conducted FISH experiments. Z.W. carried out genome assemblies; Z.W., M.R., G.D., and M.K.B. conducted gene annotation and comparative genomic analyses; Z.W., T.S., J.S.H.H., and F.C.B. conducted repetitive sequence analysis. All authors contributed to writing and editing the manuscript.

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