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Genome sequence of the cultivated cotton *Gossypium arboreum*

Fuguang Li^{1,11}, Guangyi Fan^{2,11}, Kunbo Wang^{1,11}, Fengming Sun^{2,11}, Youlu Yuan^{1,11}, Guoli Song^{1,11}, Qin Li^{3,11}, Zhiying Ma^{4,11}, Cairui Lu¹, Changsong Zou¹, Wenbin Chen², Xinming Liang², Haihong Shang¹, Weiqing Liu², Chengcheng Shi², Guanghui Xiao³, Caiyun Gou², Wuwei Ye¹, Xun Xu², Xueyan Zhang¹, Hengling Wei¹, Zhifang Li¹, Guiyin Zhang⁴, Junyi Wang², Kun Liu¹, Russell J Kohel⁵, Richard G Percy⁵, John Z Yu⁵, Yu-Xian Zhu³, Jun Wang^{2,6-10} & Shuxun Yu¹

The complex allotetraploid nature of the cotton genome (AADD; 2n = 52) makes genetic, genomic and functional analyses extremely challenging. Here we sequenced and assembled the *Gossypium arboreum* (AA; 2n = 26) genome, a putative contributor of the A subgenome. A total of 193.6 Gb of clean sequence covering the genome by 112.6-fold was obtained by paired-end sequencing. We further anchored and oriented 90.4% of the assembly on 13 pseudochromosomes and found that 68.5% of the genome is occupied by repetitive DNA sequences. We predicted 41,330 protein-coding genes in *G. arboreum*. Two whole-genome duplications were shared by *G. arboreum* and *Gossypium raimondii* before speciation. Insertions of long terminal repeats in the past 5 million years are responsible for the twofold difference in the sizes of these genomes. Comparative transcriptome studies showed the key role of the nucleotide binding site (NBS)-encoding gene family in resistance to *Verticillium dahliae* and the involvement of ethylene in the development of cotton fiber cells.

The genus *Gossypium* includes 46 diploid (2n = 2x = 26) and 5 wellestablished and 1 purported tetraploid (2n = 4x = 52) species^{1,2}. It has been proposed that all diploid cotton species may have evolved from a common ancestor that subsequently diversified to produce eight groups, including groups A-G and K³. Allopolyploid cotton may have appeared in the last 1-2 million years through hybridization and subsequent polyploidization events between the A- and D-subgenome progenitors. All tetraploid cotton species came from interspecific hybridization between the A-genome species G. arboreum (A₂) and the D-genome species G. raimondii $(D_5)^3$. The A-genome species are cultivated, whereas the D-genome species do not produce spinnable fiber. Although G. arboreum (A2) and G. raimondii (D5) are the putative donor species for the A and D chromosome groups, respectively, tetraploid cotton species differ greatly with respect to plant morphology as well as economic characteristics, including fiber production, oil content and disease resistance. Furthermore, G. arboreum (1,746 Mb/1C) has a genome size that is almost twice that of G. raimondii (885 Mb/1C)⁴.

In the current study, assisted by a high-resolution genetic map, we anchored and oriented 90.4% of the *G. arboreum* assembled scaffolds on 13 pseudochromosomes. This assembly was compared with that of *G. raimondii* to understand possible paths for genome

evolution and species divergence. A highly homozygous single-seed descendant, derived from 18 successive generations of self-fertilization, of the cultivated diploid cotton *Shixiya1* (*SXY1*) was used for DNA sequencing. Assembly of the *G. arboreum* genome and comparative studies with the genome of *G. raimondii*^{5,6} may provide new insights into the process of divergence among polyploid species^{7–9}. We believe that the *G. arboreum* genome offers a diploid reference for the analysis of cotton agronomic traits, such as fiber quality^{10–13} and disease resistance^{14,15}.

RESULTS

Genome sequencing and assembly

We sequenced and assembled the *G. arboreum* genome using the wholegenome shotgun approach. In brief, a total of 371.5 Gb of raw pairedend Illumina reads was generated by sequencing genome shotgun libraries with different fragment lengths that ranged from 180 bp to 40 kb (**Supplementary Table 1**). After filtering out low-quality reads, 193.6 Gb of high-quality sequence was obtained and used for the *de novo* assembly process (**Supplementary Table 2**). We next used 33,454 BAC end sequences (16,727 pairs; approximately 1-fold physical coverage) to improve the assembly. Over 90% of our BAC clones contained inserts of

¹State Key Laboratory of Cotton Biology, Institute of Cotton Research of the Chinese Academy of Agricultural Sciences, Anyang, China. ²BGI-Shenzhen, Shenzhen, China. ³State Key Laboratory of Protein and Plant Gene Research, College of Life Sciences, Peking University, Beijing, China. ⁴Key Laboratory for Crop Germplasm Resources of Hebei, Agricultural University of Hebei, Baoding, China. ⁵Crop Germplasm Research Unit, Southern Plains Agricultural Research Center, US Department of Agriculture–Agricultural Research Service (USDA-ARS), College Station, Texas, USA. ⁶Department of Biology, University of Copenhagen, Denmark. ⁷King Abdulaziz University, Jeddah, Saudi Arabia. ⁸Macau University of Science and Technology, Macau, China. ⁹Department of Medicine, University of Hong Kong, Hong Kong. ¹⁰State Key Laboratory of Pharmaceutical Biotechnology, University of Hong Kong. ¹¹These authors contributed equally to this work. Correspondence should be addressed to Y-X.Z. (zhuyx2@pku.edu.cn), S.Y. (yu@cricaas.com.cn) or Jun Wang (wangj@genomics.org.cn).

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Table 1 Global statistics of *G. arboreum* genome assembly and annotation

		N50	Longest	Size	Percent of the
Category	Number	(kb)	(Mb)	(Mb)	assembly
Total contigs	40,381	72.0	0.8	1,561	NA
Total scaffolds	7,914	665.8	5.9	1,694	100
Anchored and oriented scaffolds	3,740	790	5.9	1,532	90.4
Genes annotated	41,330			105	6.2
miRNAs	431			0.05	< 0.01
rRNAs	10,464			1.2	0.07
tRNAs	2,289			0.2	0.01
snRNAs	7,619			0.8	0.04
Repeat sequences	NA			1,160	68.5

NA, not applicable or not analyzed.

 \geq 50,000 bp in length (**Supplementary Fig. 1**). In-depth analysis showed that all scaffolds were supported by multiple (>80) paired-end linkages (**Supplementary Fig. 2**). The number of scaffolds decreased in libraries with increasing insert size (>2,000 bp). Using *K*-mer distribution analysis, the genome size was estimated to be 1,724 Mb (**Supplementary Fig. 3**), which is consistent with a previous report⁴.

Our final assembly, performed with SOAPdenovo^{16,17}, showed that the *G. arboreum* genome is 1,694 Mb in total length (Table 1). Ninety percent of the assembly fell into 2,724 superscaffolds that were >148 kb in length, with the largest scaffold being 5.9 Mb. N50 (the size above which 50% of the total length of the sequence assembly can be found) for the contigs and scaffolds was 72 kb and 666 kb, respectively. Data quality was assessed by comparison with Sanger sequencing-derived G. arboreum sequences and by mapping available ESTs to the assembly. Nineteen of the twenty completely sequenced G. arboreum BAC clones available from GenBank were recovered in our assembly, with >98% sequence identity (Supplementary Table 3). Of the 55,894 Sanger sequencing-derived G. arboreum ESTs (>200 bp in length) available from NCBI, 96.37% were detected in our assembly (Supplementary Table 4). Using a restriction site-associated DNA (RAD) linkage map that we constructed during the current study based on 24,569 codominant SNP markers, 1,532 Mb or 90.4% of the assembled genome was anchored and oriented on 13 pseudochromosomes (Fig. 1, Supplementary Fig. 4 and Supplementary Table 5), which contained 97.1% of the predicted gene models. Collinear relationships existed between the SNP markers and most of the assembled G. arboreum pseudochromosomes, which indicates that our assembly is of high quality (Supplementary Fig. 5). Collinearity was also observed between our G. arboreum assembly and the tetraploid genetic map reported previously¹⁸ (Supplementary Fig. 6 and Supplementary Table 6). The GC content of the G. arboreum genome was comparable to those for the genomes of G. raimondii⁵, Theobroma cacao¹⁹ and Arabidopsis thaliana (Fig. 1e).

Figure 1 Characterization of the *G. arboreum* cotton genome.
(a) Percent coverage of TEs in non-overlapping windows (window size = 500 kb). Outer tick marks show the calculated lengths of 13 *G. arboreum* pseudochromosomes. (b) Gene density estimated on the basis of the number of genes in non-overlapping 500-kb windows.
(c) Transcription state. The transcript level for each gene was estimated by averaging values of reads per kilobase of mapped cDNA per million reads (RPKM) from different tissues in non-overlapping 500-kb windows.
(d) Marker density represented by the number of SNPs in non-overlapping 500-kb windows.
(e) GC content estimated on the basis of the percentage of G+C nucleotides in 500-kb non-overlapping windows.

Genome annotation

We performed genome annotation by combining results obtained from ab initio prediction, homology search and transcriptome alignment. As much as 68.5% of the G. arboreum genome was composed of various types of repeat sequences (Fig. 1a, Table 1 and Supplementary Table 7). We believe that this genome has the greatest amount of repeat-containing sequences among sequenced eudicots¹⁹⁻²⁴. Long terminal repeat (LTR) retrotransposons accounted for 95.12% of all repeat sequences. In comparison with the G. raimondii genome⁵, the G. arboreum genome had noticeable proliferation of Gorge elements, whereas LTR Copia elements tended to accumulate in the G. raimondii genome (Supplementary Table 8), as has been suggested previously²⁵. LTR retrotransposons in G. arboreum appeared to have inserted randomly along each chromosome (Fig. 1a), in a pattern substantially different from the ones observed for their insertion sites in soybean and potato, in which LTRs are clustered near the centromere and are found less frequently near telomeres^{20,21}.

A total of 41,330 protein-coding genes were identified in the *G. arboreum* genome, with an average transcript size of 2,533 bp (as determined by GLEAN) and a mean of 4.6 exons per gene (**Fig. 1b**, **Table 1** and **Supplementary Table 9**). The genome encoded 431 micro-RNAs (miRNAs), 10,464 rRNAs, 2,289 tRNAs and 7,619 small nuclear RNAs (snRNAs) (**Table 1**). Among the annotated genes, 85.64% encoded proteins that showed homology to proteins in the TrEMBL database, and 68.71% were identified in InterPro (**Supplementary Table 10**). Over 96% of predicted coding sequences were supported by transcriptome sequencing data (**Fig. 1c** and **Supplementary Table 9**), which indicated high accuracy of *G. arboreum* gene predictions from the genome sequence. Orthologous clustering of the *G. arboreum* proteome with 3 closely related plant genomes identified 11,699 gene families in common, with 739 gene families that were present specifically in *G. arboreum* (**Supplementary Fig. 7**).

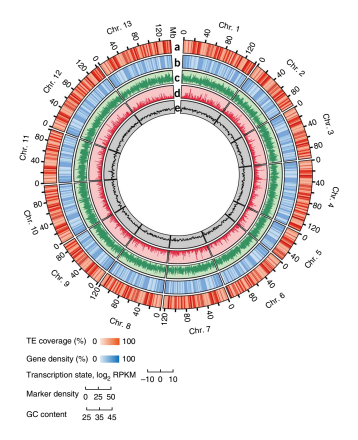


Figure 2 Phylogenetic and evolutionary analysis of the *Gossypium* genomes. (a) Phylogenetic analysis indicated that *G. arboreum* and *G. raimondii* separated ~5 million years ago (mya). (b) Fourfold-degenerate analyses suggested that the *Gossypium* genomes might have undergone two WGD events and one speciation event. (c) Large numbers of syntenic blocks were found when comparing either the *G. raimondii* (*Gr*) or *G. arboreum* (*Ga*) genome with the genome of *T. cacao* (*Tc*). Numbered rectangles represent the chromosomes.

Genome evolution

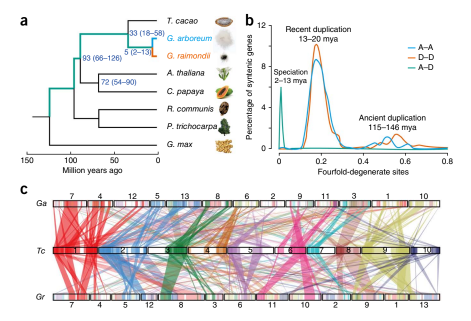
The current descendants of the two diploid cotton species that hybridized to create all cultivated allopolyploid cotton species, *G. arboreum* (A₂) and *G. raimondii* (D₅), exhibit a twofold difference in genome size^{4,26}. Molecular phylogenetic analyses²⁷ suggested a divergence time for *G. arboreum* and *G. raimondii* of 2–13 million years ago,

with their common ancestor having diverged from *T. cacao* 18–58 million years ago (**Fig. 2a**).

We scanned the G. arboreum genome for syntenic gene blocks identified on the basis of protein-coding sequence similarity. An age distribution was calculated for all duplicate gene pairs on the basis of the number of transversions at fourfold-degenerate sites. Using 1,917 paralogous gene pairs of similar age and excluding tandem or local duplications, a large peak centered around 0.17 synonymous transversions per site and a second peak at about 0.54 synonymous transversions per site were observed in both the G. arboreum and G. raimondii genomes, indicating that a recent and an ancient whole-genome duplication (WGD) event occurred in cotton (Fig. 2b). Both of these WGD events in G. arboreum coincided with those in G. raimondii, which were estimated to have occurred at 13-20 and 115-146 million years ago^{5,28}. The ancient duplication event corresponds to the ancient hexaploidization event that is shared among the eudicots^{29,30}. Using a whole-genome alignment approach, we found that both the G. raimondii and G. arboreum genomes showed close collinear relationships with the genome of T. cacao. There were 209 and 295 collinear blocks covering 82% and 66% of the G. raimondii and G. arboreum genomes, respectively (Fig. 2c and Supplementary Table 11). When the G. raimondii assembly reported by Paterson et al.⁶ was used for the analysis, a slightly lower number of collinear blocks with higher genome coverage were identified (Supplementary Table 11). Approximately 50% of the T. cacao genome was aligned with two segments in each of the cotton genomes (Fig. 2c), undoubtedly suggesting the existence of a Gossypium-specific WGD event after speciation from the *T. cacao* lineage.

Orthology maps and retrotransposition analysis

Whole-genome alignment also identified 68,863 orthologous hits (anchors) on the 2 cotton genomes, including 33,229 genes for *G. arboreum* and 34,204 genes for *G. raimondii*. These anchors spanned 17.1% of the A genome and 35.0% of the D genome (**Supplementary Table 12**). When the *G. raimondii* assembly by Paterson *et al.*⁶ was used for the analysis, we observed a total of 63,810 anchors that spanned 23% of the A genome and 48% of the D genome, indicating an improvement in data quality for the later assembly in this respect (**Supplementary Table 12**). Using these anchors, we found that the 2 cotton genomes shared 780 syntenic blocks that covered 73% and



88% of the assembled chromosomes in *G. arboreum* and *G. raimondii*, respectively (**Supplementary Table 13**). Chromosomes 1, 4–6 and 9–13 were highly collinear in the two cotton species, whereas large-scale rearrangements were observed on chromosomes 2 and 3 of *G. raimondii* and deletions and insertions were observed on chromosomes 7 and 8 of *G. arboreum* (**Fig. 3a,b**).

Expansions of transposable element (TE) families have been reported in the genome of G. raimondii⁵. In G. arboreum, two major clusters of retrotransposition activity were found at 0-0.5 and 3.5-4.5 million years ago, with two additional minor sets of retrotransposition activity present around 1.0 and 7.0-8.0 million years ago (Fig. 3c). As a result, the difference in LTR numbers between these two cotton species showed two separate peaks at ~0-0.5 and ~3.5 million years ago (Fig. 3c). Data analysis indicated that the G. arboreum genome tended to harbor more LTRs inserted during the last 0.5 million years, whereas extremely low LTR activity was observed in the G. raimondii genome over the same period of time. Thus, LTR activities substantially contributed to the twofold increase in the size of the G. arboreum genome, as previously suggested³¹. We confirmed our assertion by analyzing microsyntenic blocks on chromosome 7 of both G. arboreum and G. raimondii that were 3.5 and 1.5 Mb in length, respectively (Fig. 3d and Supplementary Fig. 8). The gene loci in these syntenic blocks were highly collinear, with 191 loci in G. raimondii and 216 loci in G. arboreum. However, 4,098 TEs were identified in G. arboreum, whereas only 1,542 TEs were found in G. raimondii (Fig. 3d). Close to 58% of the TEs reported in Figure 3d were Gorge elements in G. arboreum, whereas these elements represented only ~21% of TEs in G. raimondii (Supplementary Table 14).

Analysis of potential *Verticillium* wilt resistance genes

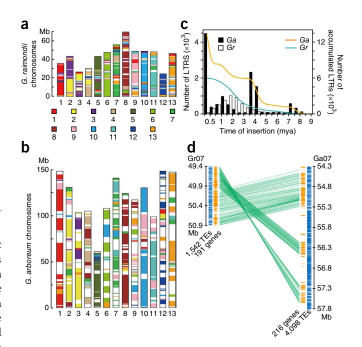
Verticillium wilt is a widespread and destructive cotton disease caused by the soil-borne fungus pathogen *V. dahliae. G. raimondii* is nearly immune to the pathogen, whereas *G. arboreum* is easily susceptible^{14,15}. Genes related to disease resistance (R genes) have critical roles during various stages of disease development in plants. Thus far, most of the cloned R genes encode NBS domains^{21,32}. We compared the *G. arboreum* genome to the *G. raimondii*⁵ and *T. cacao*¹⁹ genome sequences to identify differences in the expression patterns of NBS domain–encoding genes. We identified 391, 280 and 302 such genes in the genomes of *G. raimondii*, *G. arboreum* and *T. cacao*, respectively

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Figure 3 Whole-genome alignment and expansion of the *G. arboreum* genome. (**a**,**b**) Whole-genome alignments of *G. arboreum* on *G. raimondii* (**a**) and of *G. raimondii* on *G. arboreum* (**b**). Chromosomes from both genomes were plotted according to their assembled lengths. Each of the 13 chromosomes in **a** and **b** is designated by color; white blocks represent regions that could not be aligned. Only syntenic blocks of \geq 2 Mb in length are shown. (**c**) Analysis of LTR number and insertion time in *G. arboreum* (*Ga*) and *G. raimondii* (*Gr*). Bars (left *y* axis) indicate the number of LTRs inserted at any specific time point, whereas curves (right *y* axis) represent the accumulated total number of LTRs throughout evolution. (**d**) In-depth study of a microsyntenic block on chromosomes 7 of *G. arboreum* (GaO7) and *G. raimondii*⁵ (GrO7). Yellow bars, gene models; green lines, syntenic gene pairs; blue bars, TEs present at the corresponding chromosomal locations in each genome. The total number of genes and TEs found in each region is shown beneath each schematic.

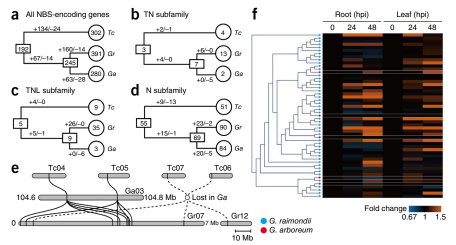
(Fig. 4a). T. cacao is also susceptible to Verticillium wilt. Phylogenetic analysis predicted that there were 192 ancestral NBS-encoding genes in the most recent common ancestor (MRCA) of cotton and cocoa (Fig. 4a and Supplementary Fig. 9). The greatest differences in the numbers of NBS-encoding gene family members in these two cotton species existed in the TN and TNL subfamilies (Fig. 4b,c), whereas the number of genes in the N subfamily remained relatively unchanged (Fig. 4d). Tandem duplications seemed to have a significant role in the expansion of the NBS-encoding gene family in G. raimondii after its divergence from G. arboreum ~5 million years ago, and segmental loss contributed to its contraction in G. arboreum (Fig. 4e). Quantitative RT-PCR (qRT-PCR) analysis of members of the TNL and TN subfamilies showed substantial gene expansion in the G. raimondii genome and indicated that many orthologs that were not present in G. arboreum responded to the pathogen soon after infection. All early pathogen-responsive genes came from the G. raimondii genome (Fig. 4f), suggesting that expansion and contraction in the numbers of NBS-encoding genes in different cotton species may have altered their resistance to V. dahliae. Less than 20% of the NBS-encoding genes from each of the diploid cotton species could be aligned with available Gossypium hirsutum ESTs (Supplementary Table 15a). Similarly, only 9 of the 53 TN and TNL subfamily members reported in Figure 4 could be amplified in the tetraploid genome (Supplementary Table 15b). These results indicate that NBS-encoding genes have evolved rapidly in either diploid or allotetraploid cotton species.

Figure 4 Analysis of NBS-encoding genes and potential molecular targets for resistance to Verticillium wilt. (a) Analysis of gene numbers in all NBS-encoding disease resistance gene families in G. arboreum (Ga), G. raimondii (Gr) and T. cacao (Tc). (b-d) Analysis of gene numbers in the TN subfamily (b), the TNL subfamily (c) and the N subfamily (d). Numbers in circles represent the number of family members in each genome, and gene numbers for the MRCA are shown in rectangles. Numbers with plus and minus signs indicate the numbers of duplicated and deleted genes, respectively. (e) Example of tandem repeat and gene deletion of several TN subfamily members in G. arboreum, G. raimondii and T. cacao. Solid lines link orthologous genes on different chromosomes, and dashed lines represent inferred orthologous relationships plus



Ethylene is a key modulator of cotton fiber cell growth

Ethylene is an important signaling molecule that promotes cotton fiber elongation in G. hirsutum^{10,11,13}. Products of the 1-aminocyclopropane-1-carboxylic acid oxidase (ACO) gene are involved in the last and rate-limiting step in ethylene biosynthesis during cotton fiber development^{10,11}. In G. raimondii ovules collected at 3 days postanthesis (dpa), the levels of ACO1 transcripts were ~1,000-fold higher and the levels of ACO3 transcripts were ~500-fold higher than in G. arboreum (Fig. 5a). These two genes were chosen because they accounted for >90% of all ACO transcripts during early fiber development (Supplementary Table 16). GrACO1 and GrACO3 showed 98% to 99% sequence identity in coding regions, respectively, with their counterparts in the G. arboreum genome (data not shown). Dot plots of promoter regions showed that a deletion of ~130 bp beginning at -470 bp relative to the transcription start site of GaACO1 resulted in loss of a putative MYB-binding site (Fig. 5b). The pivotal role of MYB transcription factors in regulating cotton fiber development and



putative deletions. Sections of *G. arboreum* chromosome 3 (GaO3) and *G. raimondii* chromosome 7 (GrO7) are enlarged for clarity, whereas all other pseudochromosomes are shown according to the scale bar. (f) qRT-PCR analysis of induced gene expression in the TN and TNL subfamilies upon *V. dahliae* infection. hpi, hours post-infection.

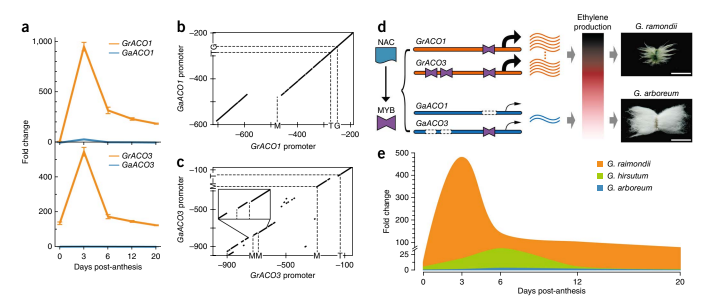


Figure 5 Ethylene modulates the elongation of cotton fiber cells. (a) Comparisons of *ACO1* and *ACO3* transcript levels during fiber cell development in *G. arboreum* and *G. raimondii*. For calculation of fold change in expression, we arbitrarily set the expression level of each *G. arboreum ACO* transcript collected at 0 dpa to 1. Error bars, s.d. for three independent experiments. (b,c) Dot plots showing consensus in the promoter regions of *ACO1* (b) and *ACO3* (c) in the two cotton species. M, putative MYB-binding site; T, putative TATA box; G, putative GATA box. (d) Schematics showing the role of ethylene production in the regulation of cotton fiber elongation. Promoters for *GrACO1* and *GrACO3* (top) are potentially bound to more MYB protein than their counterparts in *G. arboreum* (bottom). Curved lines represent corresponding *ACO* transcripts, and the gradient bar shows the amount of ethylene production from low (white) to high (black). Scale bars, 1.0 cm. (e) Comparison of total *ACO* transcript levels in three cotton species on the basis of qRT-PCR analysis (*n* = 3 biological replicates). As shown in d, *G. raimondii* produces almost no mature fiber and *G. arboreum* produces short fiber of ~1.5 cm in length; *G. hirsutum* produces fiber usually of >3.0 cm in length.

secondary cell wall biosynthesis has previously been proposed^{33,34}. Although GaACO3 and GrACO3 share one common MYB-binding site ~150 bp upstream of the transcriptional start site, two small segments of mismatch located around -750 bp to -800 bp led to loss of two additional MYB-binding sites in *G. arboreum* (Fig. 5c). Very high levels of ACO transcripts in G. raimondii ovules in conjunction with an ethylene burst^{5,10} might force an early fiber senescence phenotype, whereas the inactivation of ACO gene transcription in G. arboreum ovules might be responsible for the short-fiber phenotype in this species (Fig. 5d). When all ACO transcripts were quantified from different cotton species (Fig. 5e; see Supplementary Table 17 for primer sequences), we found that fiber-specific upregulation of ACO expression, as in the case of G. hirsutum, seems to be required for normal fiber growth. Inactivation or overproduction of ACO, as in G. arboreum and G. raimondii, respectively, might suppress fiber development. Further functional characterization of biosynthesis and signaling pathways is required to pin down such a fundamental role for ethylene in the regulation of the growth of cotton fiber cells.

DISCUSSION

Comparative genomic analysis showed that LTR insertions and expansions of LTR families contributed substantially to forming the doublesized *G. arboreum* genome relative to *G. raimondii*. The amount of sequence encompassing LTR-type retrotransposons increased from 348 Mb in *G. raimondii*⁵ to 1,145 Mb (or 67.6%) in *G. arboreum*, whereas the protein-coding capacities of these two species remained largely unchanged (**Table 1** and **Supplementary Table 7**). Typically, data obtained from the whole-genome shotgun approach in plants with genome sizes exceeding a few hundred megabases are quite difficult to assemble satisfactorily because of their high TE content^{35–37}. Here 90.4% of the 1,694-Mb *G. arboreum* genome, which is the largest of all sequenced eudicots^{19–24}, was anchored and oriented. Problems related to high TE content were circumvented by using a high-density genetic map that involved 154 F_2 RAD lines. A previous study showed an association between LTR number and ecogeography in barley, with more LTR copies present in the genomes of plants from sites with greater environmental stress³⁷. All wild *Gossypium* species seem to be adapted to arid and stressful environments. The different number of LTRs in *G. arboreum* and *G. raimondii* does not support the ecogeography hypothesis. There must be other unknown mechanisms and causes for the dramatic increase in the number of LTRs in the domesticated species *G. arboreum*.

Qualitative differences in several NBS-encoding subfamilies and in ACO gene expression between the G. arboreum and G. raimondii genomes were observed. Our data thus identify prime targets in deciphering the molecular mechanisms that control resistance to Verticillium wilt and fiber cell development. We suggest that this G. arboreum genome will be an essential reference for the assembly of tetraploid cotton genomes and for evolutionary studies of Gossypium species. It also provides an essential tool for the identification, isolation and manipulation of important cotton genes conferring agronomic traits for molecular breeding and genetic improvement.

URLs. Cotton Genome Project (CGP), http://cgp.genomics.org.cn/; LASTZ, http://www.bx.psu.edu/miller_lab/; RepeatModeler, http:// www.repeatmasker.org/RepeatModeler.html.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. The *G. arboreum* genome sequence and gene annotation information have been deposited at the Cotton Genome Project (CGP), available at http://cgp.genomics.org.cn/, and all raw sequencing

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data are accessible through the NCBI Sequence Read Archive (SRA) under accession SRA150181.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

EL., K.W., G.F., F.S., Y.Y., G.S., Jun Wang and S.Y. designed the analyses. G.F., F.S., X.X., W.C., X.L., C.G., W.L., C.S. and Junyi Wang performed sequencing, assembly and genome annotation. EL., K.W., Y.Y., G.S., C.L., C.Z., H.S. and W.Y. managed the project. Q.L., C.L., C.Z. and Y.-X.Z. performed genome analysis and physical map integration. C.L., Q.L., C.Z., G.X., W.Y., X.Z. and H.W. prepared DNA and RNA samples and performed PCR analysis. G.S., K.W., Y.Y., C.L., H.S., Z.L., K.L. and G.Z. provided the homozygous seeds. Y.-X.Z., Q.L., C.Z., C.Z. and G.X. performed transcriptome and lineage-specific gene functional analyses. Y.-X.Z., C.L., C.Z., and Q.L. conceived the project and wrote the manuscript. Jun Wang, Z.M., R.J.K., R.G.P. and J.Z.Y. conceived the project and revised the manuscript. S.Y. conceived and directed the project.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Strain selection. We used DNA samples prepared from a specific genetic line (cultivar *Shixiya1*, *SXY1*) of the cultivated diploid cotton *G. arboreum* for sequencing and assembly. This line was brought to near homozygosity after 18 successive generations of self-fertilization. *G. arboreum* cv. *Shixiya1* (*SXY1*) and *G. raimondii* D5-3 (CMD#10) plants were maintained in the National Wild Cotton Nursery in Sanya, Hainan Province, China.

DNA extraction, library construction and sequencing. Fresh young leaves were collected, immediately frozen in liquid nitrogen and stored at -80 °C until DNA extraction. We used the standard phenol-chloroform method for DNA extraction with RNase A and proteinase K treatment to prevent RNA and protein contamination¹⁰. Genomic libraries were prepared following the manufacturer's standard instructions and sequenced on the Illumina HiSeq 2000 platform. To construct paired-end libraries, DNA was fragmented by nebulization with compressed nitrogen gas, and DNA ends were blunted before adding an A base to each 3' end. DNA adaptors with a single T-base 3'-end overhang were ligated to the above products. Ligation products were purified on 0.5%, 1% or 2% agarose gels, which each targeted a specific range of insert sizes. We constructed *G. arboreum* genome sequencing libraries with insert sizes of 180 bp, 250 bp, 350 bp, 500 bp, 800 bp, 2 kb, 5 kb, 10 kb, 20 kb and 40 kb.

Genome assembly. All sequences were assembled with SOAPdenovo software^{16,17}. Unusable reads were filtered out before assembly, including (i) reads that contained sequences with >10% "N" bases, (ii) reads with low-quality data (i.e., \leq 7) for 65% of bases for short insert sizes or 80% of bases for long insert sizes, (iii) reads that contained >10 bp of adaptor sequence, (iv) reads with >10 bp that overlapped between two ends of reads of short insert size and (v) reads with identical sequences at the two ends.

The 193.6 Gb of clean data were assembled to contigs and scaffolds using the de Bruijn graph–based assembler of SOAPdenovo^{16,17} with the following four steps.

Constructing K-mer graphs. SOAPdenovo split reads with an insert size of <1,000 bases into K-mers to use in constructing the de Bruijn graphs. The parameters of this step in the cotton genome assembly were as follows: SOAPdenovo-127mer pregraph –s cotton_0308.lib –K 75 –R –p 24 –D –o GOSakoD>pregraph.log.

Building contigs. SOAPdenovo built contigs using the simplified de Bruijn graphs. The parameters of this step in the cotton genome assembly were as follows: SOAPdenovo-127mer contig –g GOSakoD –R >contig.log.

Linking scaffolds. SOAPdenovo aligned paired-end reads on the contig sequences. We subsequently calculated the total number of paired-end relationships between each pair of contigs and analyzed the rates of conflicting and consistent paired ends. A connecting relationship was regarded as a credible relationship between two contigs when that relationship met the given conditions. We then linked the contigs into scaffolds with N bases in a step-by-step manner from the paired-end reads with the shortest insert size to the paired-end reads with the longest insert size. We determined the orientation of the scaffolds on the basis of the paired-end relationship between reads and prescaffolds while we constructed the scaffolds. A reliable link was built between 2 contigs when there were \geq 3 paired-end reads for short inserts or ≥5 paired-end reads for long inserts. We started with the smallest libraries, using the RANK parameter of the configuration files to control the order of the libraries. The parameters of this step in the cotton genome assembly were as follows: SOAPdenovo-127mer map -s cotton_0308.lib -p 16 -g GOSakoD -k 39 >map.log; SOAPdenovo-127mer scaff -g GOSakoD -F -p 16 >scaff.log.

Filling gaps. We used GapCloser to fill gaps in the scaffolds. This program makes use of reads for local assembly within the gaps by aligning the other ends of paired-end reads onto scaffolds. The parameters were as follows: GapCloser –a ./GOSakoD.scafSeq –b ./cotton_0330.lib –o ./GOSakoD.scafSeq.FG –t 64.

Genetic map construction. Genetic linkage maps were constructed to develop the integrated genome map for anchoring the scaffolds, using 154 individuals from an F2 population of a cross between G. arboreum cultivars SXY1 and AnhuiFuyangZisedahua (AFZ). Young leaves of parents and individuals from the F2 population were collected for DNA extraction using the Plant Genomics DNA kit (Qiagen). Genomic DNA from this mapping population was digested with the restriction endonuclease EcoRI and processed into RAD libraries³⁸. Every 20 cotton individuals were pooled into 1 sequencing library with nucleotide multiplex identifiers (4, 6 and 8 bp), and each sequencing library was barcoded. Approximately 1,600 Mb of 50-bp reads (10 Mb of read data for each progeny on average) was generated on the HiSeq 2000 nextgeneration sequencing platform. The SNP calling process was carried out using the SOAP2+SOAPsnp pipeline³⁹. JoinMap4.0 (ref. 40) was used to conduct the linkage analysis. A χ^2 test was used to determine whether the RAD-based SNP markers corresponded with the expected segregation ratio. Unusable markers (P < 0.01) were filtered out before a genetic map was constructed. ABH-style markers were used to build the main framework of 13 linkage groups, and a logarithm of odds (LOD) score of 7 was initially set as the linkage threshold for linkage group identification. AC-style markers were also included to expand the length of linkage groups by reducing the LOD value to 3. All high-quality markers were used to construct the consensus maps with the F2 population in JoinMap4.0. We calculated the recombination fractions between all pairs of SNP markers in a scaffold and chose the SNP marker that had the minimum recombination fraction in the sum. Scaffold order was determined by tag SNPs, and inner SNPs were then used to orient the scaffolds.

Annotation of transposable elements. We searched the genome for tandem repeats using the software program Tandem Repeats Finder⁴¹. Both homology-based and de novo approaches were used to find TEs. The homologybased approach involved applying commonly used databases of known repetitive sequences along with such programs as RepeatProteinMask and RepeatMasker⁴². To identify DNA-level TEs, we used Repbase⁴³ along with a database of plant repeating sequences and our de novo TE library to find repeats with RepeatMasker. We used four software programs, LTR_FINDER⁴⁴, PILER⁴⁵, RepeatModeler and RepeatScout⁴⁶, to generate the *de novo* repeat library. These programs predict repeats in different ways. LTR_FINDER retrieves full-length LTR retrotransposons, which usually contain ~18 bp of sequence that is complementary to the 3' tail of certain tRNAs. PILER searches for repeats in the genome by aligning the genome with itself. RepeatScout builds a consensus sequence on the basis of lmer using the fit-preferred alignment score. RepeatModeler uses two ab initio repeat prediction programs (RECON and RepeatScout), which identify repeat element boundaries and family relationships among sequences. Intact LTRs were predicted using LTR_STRUC47 Windows-based software, and solo LTRs were predicted on the basis of comparison with intact ancestor LTRs. Single-copy gene families were used to construct the phylogenetic tree, and syntenic blocks were identified by MCscan⁴⁸.

Gene prediction. Homology-based gene prediction. The target locations of homologous proteins were obtained by aligning the protein sequences of *A. thaliana, Carica papaya, Populus trichocarpa, G. raimondii, Populus trichocarpa, T. cacao* and *Vitis vinifera* to the *G. arboreum* genome using TBLASTN with an *E*-value parameter of 1×10^{-5} . Proteins encoded by TEs were filtered out before alignment. We next extracted target sequences from the genome, including a 2,000-bp extension at both ends of the alignment with intron regions, and again aligned the homologous protein sequences to these DNA fragments using GeneWise⁴⁹ (with parameters –trev –sum –genesf). Coding sequences were aligned against the genome using BLAT to generate spliced alignments, and overlapping sequences were filtered out to link the spliced alignments using PASA⁵⁰.

De novo gene prediction. AUGUSTUS (version 2.5.5)⁵¹ and SNAP (version 2006-07-28)⁵² were applied for *de novo* gene prediction using gene model parameters trained by *A. thaliana.* We masked all TEs from the genome before gene prediction and filtered out all short coding regions that were <150 bp in length during the process.

Transcriptome alignment. We used TopHat⁵³ to map raw reads of cDNA sequence to the *G. arboreum* genome sequence to identify potential areas of exons in addition to donor or receptor sites with the following parameters: –p 4-max-intron-length 20,000 –m 1 –r 20-mate-std-dev 20. Cufflinks⁵³ software was used to assemble potential alignments to transcripts, with the following parameters: –I 20,000 –p 4. To obtain complete gene models, we used the fifth-order Markov model to predict ORFs.

Combination of all methods. Data derived from homology-based, *de novo* and transcriptome alignment methods were integrated to generate a consensus gene set by GLEAN⁵⁴ with default parameters.

Phylogenetic analysis. Gene Ontology (GO)⁵⁵ terms for each gene were obtained from its corresponding InterPro entry. We used OrthoMCL⁵⁶ to confirm the genes that were orthologous among four species (*A. thaliana*, *G. arboreum*, *G. raimondii* and *T. cacao*). All-versus-all BLASTP (*E* value < 1×10^{-5}) comparison of all protein sequences for each species was performed, and orthologous genes were clustered by OrthoMCL (inflation parameter: 1.5). We performed multiple alignments of protein sequences with MUSCLE⁵⁷ for each single-copy gene family and converted the protein alignments to coding sequences using a Perl script. Phase-1 sites were extracted from each family and concatenated to one supergene for every species, and Mrbayes 3.1.2 (ref. 58) was used to construct the phylogenetic tree. Branch-specific Ka/Ks ratios (the number of nonsynonymous substitutions per nonsynonymous site, Ka, over the number of synonymous substitutions per synonymous site, Ks) were calculated using codeml in the PAML⁵⁹ software package with the branch model.

Syntenic analysis and whole-genome alignment. A BLASTP search (with an *E*-value cutoff of 1×10^{-5}) was performed to identify paralogous genes. Syntenic blocks (with at least five genes per block) were identified by MCscan⁴⁸. For the alignment results between these, each aligned block represented the orthologous pair derived from the common ancestor, and the sequences that contained the genes were used to show the intergenome relationships with their length information. The fourfold-degenerate value of the blocks was calculated as revised by the HKY model, and whole-genome alignment was carried out by LASTZ (see URLs) between *G. raimondii* and *G. arboreum* after repeat regions were first masked.

LTR analysis. Intact LTRs from the A genome and D genome were predicted using LTR_STRUC 47 Windows-based software. We constructed ancestor subfamilies on the basis of the following two rules: (i) family members had to share the same polypurine tract sequences and primer-binding-site sequences and (ii) the *E* value had to be $< 1 \times 10^{-10}$ among transposable gene sequences. For each family, we aligned all intact LTRs with MUSCLE⁵⁷ and manually corrected the alignments using MEGA5.5 (ref. 60). We then constructed ancestor sequences for this family using the cons program (contained in the EMBOSS package⁶¹). To find more solo LTRs, we used the ancestor sequences for different families as queries for LTR prediction based on the standard prediction pipeline, and we followed the 80-80-80 rule (identity, >0.8; align rate, >80%; alignment length, >80 bp)⁶² to find which family the solo LTRs belonged to. All intact LTRs and solo LTRs were used to calculate the insert time with the formula time = K/r (where K is the distance between all alignment pairs and *r* is the rate of nucleotide substitution). The value was set to 7×10^{-9} , and *K* was calculated with the distmat program implemented in the EMBOSS package⁶¹ with the Kimura two-parameter model.

Analysis of disease resistance- and fiber development-related genes. The largest class of characterized disease resistance (R) genes encodes intracellular proteins that contain a nucleotide-binding site (NBS) and C terminus with leucine-rich repeats (LRRs)⁶³. These genes are rather abundant in plant genomes and have an important role in mediating resistance to pathogens^{19,64}. Resistance-related genes were identified using the HMMER 3.0 (ref. 65) software package. CAFE⁶⁶ was used to predict the expansion and contraction of gene numbers on the basis of the topological gene tree.

ACO genes in *G. raimondii* and *G. arboreum* were obtained on the basis of a homology search with the ACO genes in *G. hirsutum*. Dot plot figures were generated using a word size of 9. All fold-change values were calculated with respect to the *G. arboreum* expression level at 0 dpa. For qRT–PCR analysis, total RNA (~2 µg) was reverse transcribed in a 20-µl reaction mixture using SuperScript II reverse transcriptase (Invitrogen). After the reaction, 1-µl aliquots were used as a template for PCR amplification with gene-specific primers. As an internal control, the *UBQ7* transcript was used to quantify the relative transcript level of each target gene in each tissue type. Three biological replicates were analyzed for all qRT-PCR experiments.

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