MaGenDB: a functional genomics hub for Malvaceae plants

Dehe Wang Δ1,2,†, Weiliang Fan Δ1,†, Xiaolong Guo Δ1,2,†, Kai Wu Δ1,2, Siyu Zhou 3, Zonggui Chen 4, Danyang Li 1, Kun Wang Δ1,†, Yuxian Zhu Δ1, 4,* and Yu Zhou Δ1,2,4,*

1College of Life Sciences, Wuhan University, Wuhan 430072, China, 2State Key Laboratory of Virology, Wuhan University, Wuhan 430072, China, 3College of Information Science and Engineering, Hunan University, Changsha 410082, China and 4Institute for Advanced Studies, Wuhan University, Wuhan 430072, China

Received August 15, 2019; Revised October 08, 2019; Editorial Decision October 09, 2019; Accepted October 10, 2019

ABSTRACT

Malvaceae is a family of flowering plants containing many economically important plant species including cotton, cacao and durian. Recently, the genomes of several Malvaceae species have been decoded, and many omics data were generated for individual species. However, no integrative database of multiple species, enabling users to jointly compare and analyse relevant data, is available for Malvaceae. Thus, we developed a user-friendly database named MaGenDB (http://magen.whu.edu.cn) as a functional genomics hub for the plant community. We collected the genomes of 13 Malvaceae species, and comprehensively annotated genes from different perspectives including functional RNA/protein element, gene ontology, KEGG orthology, and gene family. We processed 374 sets of diverse omics data with the ENCODE pipelines and integrated them into a customised genome browser, and designed multiple dynamic charts to present gene/RNA/protein-level knowledge such as dynamic expression profiles and functional elements. We also implemented a smart search system for efficiently mining genes. In addition, we constructed a functional comparison system to help comparative analysis between genes on multiple features in one species or across closely related species. This database and associated tools will allow users to quickly retrieve large-scale functional information for biological discovery.

INTRODUCTION

The Malvaceae is a family of flowering plants containing 243 genera with 4225 known species, including many economically important crop plants (1). These include cotton (Gossypium spp.), a principal natural fibre source for the textile industry; cacao (Theobroma spp.), yielding millions of tonnes of cocoa for making chocolate; durian (Durio spp.), a major Southeast Asian food known as the ‘king of fruits’; and okra (Abelmoschus spp.), which is a nutritious vegetable. These species exhibit wide phenotypic diversity in life forms (from herbaceous to woody), flowers and seeds. The underlying functional genomic elements need to be deciphered to enhance their traits in productivity and quality.

In recent years, genome sequencing data have rapidly accumulated for Malvaceae species, including Theobroma cacao (2), four species of cotton (3–9), Durio zibethinus (10), and Bombax ceiba (11). Many transcriptomic, epigenomic and proteomic studies have uncovered the gene expression profiles and genomic features in individual species. However, an integrative functional genomic database of multiple species, enabling users to jointly examine and utilize relevant data, is missing for the Malvaceae, probably due to difficulties in collecting, analyzing, storing, and visualizing large-scale data. The previously published CottonGen (12), CottonFGD (13) and ccNET (14) databases are useful, but are cotton-specific and lack functions for comparative genomics.

To provide a functional genomics hub for the Malvaceae and the plant community, we developed a user-friendly database named the MaGenDB (http://magen.whu.edu.cn). We collected the genomes of 13 species which coverage all genera in Malvaceae with whole genome sequence in NCBI genome database, and comprehensively analysed the functional features of genes including various RNA/protein elements, gene ontology (GO), KEGG orthology (KO), enzyme characterization, protein 3D structure and gene families. To date, we processed 374 sets of diverse omics data with standard pipelines and integrated them into MaGenDB, including ChIP-seq, DNase-seq, BS-
Gene functional annotation

All functional elements in Malvaceae genomes were thoroughly annotated with a unified and standard procedure (Figure 1B). For genomes without gene annotations, gene models were reconstructed using StringTie (16) with default parameters from RNA-seq data, except the IGIA gene annotation for *Gossypium arboreum* was recently assembled from integrative multi-omeric RNA-seq data (17). The tRNA and various kinds of ncRNA genes were predicted using tRNAscan-SE 2.0 (18) and rnam_scan with RFam models (19), respectively. The tandem repeats were identified using TRF program (20). The protein-coding potential scores were computed using CPAT (21). Three local BLAST databases were built, including the NCBI Nucleotide (NT), non-redundant protein, and the Arabidopsis Information Resource (TAIR) databases (22), to which the DNA and protein sequences of all gene models were compared using blastn and blastp programs (23) with E-value cutoff 1e-6. The GO terms and EC numbers for genes were assigned using Blast2GO (24). The KO identifiers were annotated using the KAAS web server (25), and homologous genes were identified using the eggNOG web server (26). Potential transcription-factor (TF) families of protein-coding genes were mapped to the PlantTFDB database (27).

VARIOUS functional protein domains and RNA elements were also identified. InterPro, Pfam, and conserved protein domains were predicted using InterProScan (28) and InterPro database (29), PfamScan (30) and Pfam database (31), and NCBI conserved domains (CDD) database (32), respectively. Signal peptides, transmembrane helices and disorder regions were predicted using signalP (33), TMHMM (34) and IUPred2A (35), respectively. The RNA G-quadruplexes (RG4) were predicted using QGRS (36) and miRNA target sites were predicted by psRNATarget web server (37). Protein 3D structures were modelled using the SWISS-MODEL web server (38).

**MATERIALS AND METHODS**

All data sources, data processing, and web interface features are summarized in Figure 1. The MaGenDB consists of four parts: data collection, gene functional annotation and omics data processing, database construction and web interface and toolkit development. The key steps, tools, and generated data are summarized in Supplementary Figure S1 and described below.

**Data sources**

All species and genome assemblies used in this study were collected from public databases (Supplementary Table S1). The genome-wide association study (GWAS) and single nucleotide polymorphism (SNP) data for genomes were manually extracted from the NCBI PubMed database. Diverse types of omics data—including RNA-seq, small RNA-seq (smRNA-seq), CAGE-seq, polyA-seq, ChIP-seq, DNase-seq, Bisulfite-seq (BS-seq) and PacBio long-read sequencing—were downloaded from the NCBI SRA database (www.ncbi.nlm.nih.gov/sra). Mass spectrometry data were downloaded from the ProteomeXchange (15) database (Figure 1A). All metadata of these omics data are summarised in Supplementary Table S2.

**Gene functional annotation**

All functional elements in Malvaceae genomes were thoroughly annotated with a unified and standard procedure (Figure 1B). For genomes without gene annotations, gene models were reconstructed using StringTie (16) with default parameters from RNA-seq data, except the IGIA gene annotation for *Gossypium arboreum* was recently assembled from integrative multi-omeric RNA-seq data (17). The tRNA and various kinds of ncRNA genes were predicted using tRNAscan-SE 2.0 (18) and rnam_scan with RFam models (19), respectively. The tandem repeats were identified using TRF program (20). The protein-coding potential scores were computed using CPAT (21). Three local BLAST databases were built, including the NCBI Nucleotide (NT), non-redundant protein, and the Arabidopsis Information Resource (TAIR) databases (22), to which the DNA and protein sequences of all gene models were compared using blastn and blastp programs (23) with E-value cutoff 1e-6. The GO terms and EC numbers for genes were assigned using Blast2GO (24). The KO identifiers were annotated using the KAAS web server (25), and homologous genes were identified using the eggNOG web server (26). Potential transcription-factor (TF) families of protein-coding genes were mapped to the PlantTFDB database (27).

VARIOUS functional protein domains and RNA elements were also identified. InterPro, Pfam, and conserved protein domains were predicted using InterProScan (28) and InterPro database (29), PfamScan (30) and Pfam database (31), and NCBI conserved domains (CDD) database (32), respectively. Signal peptides, transmembrane helices and disorder regions were predicted using signalP (33), TMHMM (34) and IUPred2A (35), respectively. The RNA G-quadruplexes (RG4) were predicted using QGRS (36) and miRNA target sites were predicted by psRNATarget web server (37). Protein 3D structures were modelled using the SWISS-MODEL web server (38).

**Comparative genome analysis**

The plant genomes for comparative analysis with Malvaceae were downloaded from PLAZA (39). Gene synteny clusters between any two genomes in MaGenDB were predicted using MCScanX (40). Multiple sequence alignments between collinear genes were built using Clustal Omega (41).

Protein–protein interactions (PPI) were extracted from the STRING database (42) for two Malvaceae genomes (*Gossypium raimondii* and *Theobroma cacao*) and *Arabidopsis thaliana*. A similar but stricter strategy to that used in STRING, transferring PPI score by gene collinearity, was used to predict the PPI network for other genomes in MaGenDB which were highly collinear. For two proteins A and B, the ‘combined score’ from one genome was calculated as the average interaction scores between collinear genes pairs of A and B. The ‘combined score’ from multiple genomes were averaged as the predicted interaction strength.

**Omits data analysis**

High-throughput sequencing data passing the fastQC quality-control (www.bioinformatics.babraham.ac.uk/projects/fastqc/) were used in this study. The raw reads were filtered to remove the adaptors and low-quality bases using cutadapt (43). For the omics data of RNA-seq, smRNA-seq, ChIP-seq, DNase-seq and BS-seq, the data processing followed the recommendations of the ENCODE pipelines (44). Briefly, RNA-seq and smRNA-seq reads were mapped to the genome using STAR (45). The ChIP-seq and DNase-seq reads were mapped by BWA (46), and the BS-seq reads were processed by Bismark (47). The gene expression quantification in FPKM (Fragments Per Kb of exon per Million mapped fragments) was computed by StringTie (16). The peaks for ChIP-seq and DNase-seq were identified by MACS2 (48) and Hotspot2 (newer version of Hotspot) (49), respectively.

For CAGE-seq and polyA-seq, the 5′ random barcodes were extracted and trimmed before mapping. The potential rRNAs and PCR duplicate reads were removed, and the cleaned reads were mapped to the genome using STAR in End-to-End mode.

For PacBio data, the long subreads were corrected with RNA-seq data using proovread (50) to reduce the sequencing errors. The corrected reads were aligned to the genome
Figure 1. Schematic of MaGenDB. (A) Data sources included 13 species from five subfamilies in Malvaceae and associated gene models, GWAS, eight types of sequencing data, and MS data. (B) Summary of data processing and obtained biological knowledge. (C) All data are stored in a MySQL relationship database with additional indexes. Django and React frameworks are used for interactive queries between frontend and backend. (D) Overview of the web interface and usage of MaGenDB. Main functions or data are listed under the first-level menu. The dashed lines indicate the linkages between different pages.

using STARlong with the parameters recommended by the PacBio official documentation.

The raw mass spectra data were converted to mgf format using ProteoWizard (51) before peptide searching. The programs SearchGUI (52) and peptideShaker (53) were used to match the MS/MS fragmentation spectra to peptide sequences with default parameters. The genomic positions of matched peptides were recovered using custom Python scripts.

Database construction

All pre-processed data were integrated into the MaGenDB MySQL database, in which different tables were associated together by the functional element, ontology, and genomic position (Figure 1C). Django framework was used to provide query and computation supports from the database backend.

Web interface and toolkit development

A user-friendly web interface was developed by React framework (Figure 1D), and multiple customised dynamic charts were implemented using BizCharts library. All genomic features and omics data were visualised using JBrowse (54) and its plugins. The BLAST server was driven by Sequenceserver (55). The PCR primer design was im-
Table 1. Summary of MaGenDB data

| Data                                      | Statistics                      |
|-------------------------------------------|---------------------------------|
| Subfamily/Genus/Species                   | 5/9/13                          |
| Transcripts/Proteins                      | 759 700/728 366                 |
| Types of omics data                      | 9                               |
| Omics data samples                        | 374                             |
| GO/KO/EC annotation                       | 494 364/254 551/85 157          |
| Gene/TF families                          | 656 516/280 976                 |
| BLAST annotation (NT/NR/TAIR)            | 63 313 713/307 542 675/635 804  |
| Types of functional elements              | 7                               |
| Functional elements                       | 24 855 369                      |
| Synteny gene pairs                        | 28 976 317                      |
| Protein-protein interactions              | 170 660 615                     |
| Protein 3D structures                     | 98 831                          |

Table 2. Query fields and examples in smart search system

| Query field                        | Query content                                         | Query example                        |
|------------------------------------|-------------------------------------------------------|--------------------------------------|
| Name                               | Locus/Transcript/Protein name                         | GhRDL1                               |
| Name                               | Locus/Transcript/Protein ID                           | Ga14g01129.F1                        |
| Name                               | TAIR homolog gene symbol                              | ACX3                                 |
| GO                                 | GO accession ID                                       | GO:0000032                           |
| KO                                 | KO accession ID                                       | K11091                               |
| KO                                 | KEGG ontology name                                    | ATPF1B, atpD                         |
| EC                                 | Enzyme Commission ID                                  | 1.17.7.2                             |
| EC                                 | Enzyme Commission name                                | Very-long-chain 3-oxoacyl-CoA synthase|
| KEGG pathway                       | KEGG pathway ID                                       | cAMP signaling pathway               |
| KEGG pathway                       | KEGG pathway name                                     | M00087                               |
| KEGG model                         | KEGG model ID                                         | Fe-S protein                         |
| KEGG model                         | KEGG model name                                       | H00254                               |
| KEGG disease                       | KEGG disease ID                                       | Lysosomal acid lipase deficiency     |
| KEGG disease                       | KEGG disease name                                     | IPR001736                            |
| InterPro domain                    | InterPro ID                                           | cd01254                              |
| InterPro domain                    | InterPro domain name                                  | PF00614.22                           |
| Pfam domain                        | Pfam ID                                               | PLDc                                 |
| Pfam domain                        | Pfam domain name                                      | 197200                               |
| CDD domain                         | NCBI CDD ID                                           |                                       |
| TF family                          | TF family name                                        | C2H2                                 |
| eggNOG gene family                 | eggNOG root name                                      | E0Y06593                             |
| eggNOG gene family                 | eggNOG hit gene                                       | COG1502                              |

implemented by Primer3 (56) with customization to accommodate support for alternative splicing events. The GO enrichment analysis was provided using R package topGO. Protein 3D structure was presented using NGL viewer (57). GenomeSyn viewer was developed to visualize gene synteny clusters between two genomes. All programs and packages used in this study are listed in Supplementary Table S3.

RESULTS

Overview of MaGenDB

MaGenDB (http://magen.whu.edu.cn) is a family-level functional genomics hub for Malvaceae plants, which contains 367 available deep-sequencing data of eight types for 13 species (Figure 1A). For each genome, the database provides multiple functional annotations, DNA variations, chromatin and RNA polymerase II states, and RNA landscape including gene expression across different tissues (Figure 1B). The GO/KO/EC annotations, gene family, and TF family for 759 700 transcripts are stored in MaGenDB. About 370 million BLAST annotations to the NT, NCBI non-redundant protein database, and TAIR are saved in MaGenDB. More than 24 million functional elements are annotated, including InterPro domains, Pfam domains, CDD, signal peptides, transmembrane helices, protein disorder regions and RG4. The database contains 170 660 615 PPIs and 98 831 protein 3D structures from prediction (Table 1). The coding-potential scores for all transcripts and the proteins with mass spectrometry evidence were recorded. For comparative genome analysis, 51 other plant genomes were collected (Supplementary Table S4) and a total of 28 976 317 gene syntenies were annotated (Table 1).

The MaGenDB is aimed to be a user-friendly comparative functional genomics database, in which multiple dynamic charts and hyperlinks are generated. The web pages were well-designed, and several useful tools were developed in MaGenDB for researchers (Figure 1D). A detailed user guide was provided to help use MaGenDB efficiently. The database was designed to also be compatible with tablet and phone devices. All data stored in the MaGenDB are freely accessible and downloadable for academic purposes.

Smart search system

For users to quickly find the data of interest, a smart search system was designed for genes, transcripts, and proteins. The supported queries include the gene symbol; the identifiers of gene, transcript, and protein; the name and iden-
For example, the gene name ‘Pgam5’, TAIR gene symbol, or gene ID ‘Ga01g00214.F3’ can be used to query the same gene. Fuzzy matching is supported, for example, ‘cAMP’ instead of ‘cAMP signalling pathway’. Additionally, the search box can trigger the appearance of a list of suitable queries that is filtered as the user types.

**GeneWiki**

GeneWiki is the page where MaGenDB displays all functional annotations for gene, transcript, and protein (Figure 2), such as gene structure, ontology annotation, gene expression, mass spectra evidence, functional domains, gene synteny and predicted protein 3D structure. Users land in this page from the ‘Search’ page or other hyperlinks. Detailed data can be viewed as organised sections by clicking the drop-down buttons (Figure 2A).

For a gene locus, the information of gene symbol, taxonomy, genomic position and gene type are shown in the ‘Summary’ section. Users can view the taxonomy, jump to the genome browser, or download DNA sequences via hyperlinks. All unique transcripts and proteins are listed in the ‘Transcript information’ and ‘ORF information’ sections, respectively (Figure 2A).

The ‘Functional annotation’ section contains the NT annotation and *Arabidopsis thaliana* homolog gene predicted by BLAST. The annotation of eggNOG gene family and TF genes from PlantTFDB are arranged in the sections ‘Gene family’ and ‘TF family’. The GO, KEGG ontology annotation, pathway, model and disease annotation, and enzyme commission annotation are listed in the ‘Ontology annotation’ section (Figure 2A).

The mass spectra evidence from the different sources is shown using dynamic charts and tables (Figure 2B). The detailed information includes data source, matched pep-
Figure 3. Genome browser and functional genomics tools in MaGenDB. (A) Genome browser view of processed omics data for a gene example. For any genome, the available data are organised in track groups and can be dynamically selected/unselected. (B) Overview of genomics tools provided in MaGenDB. (C) The interface of BLAST service for the different databases. (D) Example of genome map viewer of multiple genes. (E) Primer design page considering alternative splicing (AS) events.

tide, and peptide-spectrum match (PSM). The elements can be viewed at multiple linear scales including gene locus, mRNA transcript, and amino acid sequence of protein.

The ‘Gene expression’ section shows the expression profiles in different tissues, with error bars indicating the standard deviation among different datasets (Figure 2C). The expression values in FPKM of each data set are shown in the bottom table.

The ‘Functional element’ section includes InterPro domain, NCBI CDD domain, Pfam domain, signal peptide, transmembrane helices, disorder region, and RG4, displayed by dynamic charts in different colours (Figure 2D). The protein network interacting with the query gene is shown in an interactive chart (Figure 2E). The combined scores of interactions are shown and can be used to filter the network. The information for a specific gene appears when clicking the corresponding node. The 3D structure model of protein predicted using SWISS-MODEL is displayed as an interactive figure in the ‘Protein 3D structure’ section (Figure 2F). The sequence alignment between the protein and its template can be dynamically explored.

Genome browser and multi-omics data integration

All gene models, processed omics data, annotated functional elements in MaGenDB are visualised using customised JBrowse with suitable default settings (Figure 3A). A set of plugins were configured to enhance the functional-
Figure 4. Comparative genomics tools in MaGenDB. (A) Management of gene list of interest for comparative analysis. (B) Gene expression heatmap across different tissues for collinear genes to a query gene from the GeneWiki page. (C) Genomic position mapping of collinear genes. (D) Comparison of functional domains between two proteins. Grey box marks the missing region in one vs the other. (E) An interactive view of multiple alignments of collinear genes from the gene structure perspective. (F) A circular view of the gene synteny clusters between \textit{G. arboreum} Chr06 with \textit{G. hirsutum} chromosomes.

characteristics (60). The MaGenDB can automatically generate gene expression heatmaps for all genes in the synteny block of the query gene, as shown in a fiber and ovule early stage specific gene example, of which the collinear genes have same expression pattern (Figure 4B). Furthermore, the chromosomal locations of collinear gene clusters can be visualized interactively (Figure 4C).

To intuitively evaluate the effects on proteins of alternative RNA processing, we developed the gene comparator tool to compare biological annotations and functional structures between two gene loci, transcripts, or proteins. An example is shown in Figure 4D, in which the protein Ga05g02643.F1 lost almost two-thirds of the amino acid sequence compared to the protein evm.model.Ga05G2709 due to alternative transcription start sites, and so the Ga05g02643.F1 protein is missing a significant amount of InterPro domains. The Gene Synteny viewer was developed to compare gene structures between collinear genes. The
multi-alignments are dynamically presented, which facilitates discovery of conserved elements, as shown in Figure 4E. The gene synteny clusters between two genomes can be dynamically presented with GenomeSyn viewer as circular plot. As shown in Figure 4F, the Chr06 from diploid *G. arboresum* and the two homologous chromosomes from tetraploid *G. hirsutum* have the largest synteny blocks, indicating good quality of our results.

**DISCUSSION**

The MaGenDB fills the gap for an important plant family, integrates large-scale diverse omics data, implements comprehensive data visualization methods and constructs a new functional comparison system. A total of 374 processed omics data of nine techniques, 18 types of annotation and >24 million functional elements are stored in MaGenDB and presented in a user-friendly way using well-designed custom dynamic charts. The large set of curated data sets and several powerful tools provide comparative functional genomics resources and services. Thus, MaGenDB will be useful for plant and evolution scientists in both experimental and computational directions. The differences between MaGenDB and other cotton-specific databases are also summarized in Supplementary Table S5. In the future, we will continuously update MaGenDB as new Malvaceae genomes and omics data become available, and will add more annotation and functionalities to the database including whole genome alignments in the context of comparative genome analysis, and other types of omics data such as Hi-C, ChIA-PET and CLIP-seq data.

**DATA AVAILABILITY**

The MaGenDB database can be accessed through the web server at http://magen.whu.edu.cn. The codes for the omics data analysis are available in the GitHub repository at https://github.com/zhouyulab/magendb.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**ACKNOWLEDGEMENTS**

Part of the computation in this work was done on the supercomputing system in the Supercomputing Center of Wuhan University.

**FUNDING**

Innovation Team Program from Wuhan University [2042017kf0233 to Y.Z., Y.X.Z., K.W.]; National Natural Science Foundation of China [91640115 and 31670827 to Y.Z., 31690090 to Y.X.Z., 31770310 to K.W.]; National Key R&D Program of China [2017YFA0504400 to Y.Z.]; Funding for open access charge: Innovation Team Program from Wuhan University [2042017kf0233].

Conflict of interest statement. None declared.

**REFERENCES**

1. Group, T. The Angiosperm Phylogeny (2016) An update of the Angiosperm Phylogeny Group classification for the orders and families of flowering plants. *APG IV*. *Bot. J. Linn. Soc.*, 181, 1–20.
2. Argout, X., Salse, J., Aury, J.-M., Guittinan, M., Droc, G., Gouzy, J., Allegr, M., Chaparro, C., Legare, T., Maximova, S. et al. (2011) The genome of Theobroma cacao. *Nat. Genet.*, 43, 101–108.
3. Paterson, A.H., Wendel, J.F., Gundlach, H., Guo, H., Jenkins, J., Jin, D., Llewellyn, D., Showmaker, K.C., Shu, S., Udall, J. et al. (2012) Repeated polyploidization of Gossypium genomes and the evolution of spinnable cotton fibres. *Nature*, 492, 423–427.
4. Wang, K., Wang, Z., Li, F., Ye, W., Wang, J., Song, G., Yue, Z., Cong, L., Shang, H., Zhu, S. et al. (2012) The draft genome of a diploid cotton *Gossypium raimondii*. *Nat. Genet.*, 44, 1098–1103.
5. Li, F., Fan, G., Wang, K., Sun, F., Yuan, Y., Song, G., Li, Q., Ma, Z., Lu, C., Zou, C. et al. (2014) Genome sequence of the cultivated cotton *Gossypium arboresum*. *Nat. Genet.*, 46, 567–572.
6. Li, F., Fan, G., Lu, C., Xiao, G., Zou, C., Kohel, R.J., Ma, Z., Shang, H., Ma, X., Wu, J. et al. (2015) Genome sequence of cultivated Upland cotton (*Gossypium hirsutum* TM-1) improves insights into genome evolution. *Nat. Biotechnol.*, 33, 524–530.
7. Du, X., Huang, G., He, S., Yang, Z., Sun, G., Ma, X., Li, N., Zhang, X., Sun, J., Liu, M. et al. (2018) Resequencing of 243 diploid cotton accessions based on an updated A genome identifies the genetic basis of key agronomic traits. *Nat. Genet.*, 50, 796–802.
8. Wang, M., Tu, L., Yuan, D., Zhu, D., Shen, C., Li, J., Liu, F., Pei, L., Wang, P., Zhao, G. et al. (2019) Reference genome sequences of two cultivated allotetraploid cottons, *Gossypium hirsutum* and *Gossypium barbadense*. *Nat. Genet.*, 51, 224–229.
9. Hu, Y., Chen, J., Fan, L., Zhang, Z., Ma, W., Niu, Y., Ju, L., Deng, J., Zhao, T., Lian, J. et al. (2019) *Gossypium barbadense* and *Gossypium hirsutum* genome sequences provide insights into the origin and evolution of allotetraploid cotton. *Nat. Genet.*, 51, 739–748.
10. Teh, B.T., Lim, K., Yong, C.H., Ng, C.C.Y., Rao, S.R., Rajasegaran, V., Lim, W.K., Ong, C.K., Chan, K., Cheng, V.Y. et al. (2017) The draft genome of tropical fruit durian (*Durio zibethinus*). *Nat. Genet.*, 49, 1633–1641.
11. Gao, Y., Wang, H., Liu, C., Chu, H., Dai, D., Song, S., Yu, L., Han, L., Fu, Y., Tian, B. et al. (2018) De novo genome assembly of the red silk cotton tree (*Bombax ceiba*). *GigaScience*, 7, giy051.
12. Yu, J., Jung, S., Cheng, C.-H., Ficklin, S.P., Lee, T., Zheng, P., Jones, D., Percy, R.G. and Main, D. (2014) CottonGen: a genomics, genetics and breeding database for cotton research. *Nucleic Acids Res.*, 42, D1229–D1236.
13. Zhu, T., Liang, C., Meng, Z., Sun, G., Meng, Z., Guo, S. and Zhang, R. (2017) CottonFGD: an integrated functional genomics database for cotton. *BMC Plant Biol.*, 17, 101.
14. You, Q., Xu, W., Zhang, K., Zhang, L., Yi, X., Yao, D., Wang, C., Zhang, X., Zhao, X., Provart, N.J. et al. (2017) ccNET: Database of co-expression networks with functional modules for diploid and polyploid *Gossypium*. *Nucleic Acids Res.*, 45, D1090–D1099.
15. Deutsch, E.W., Csdoras, A., Sun, Z., Jarnuczak, A., Perez-Riverol, Y., Terenti, T., Campbell, D.S., Bernal-Llinares, M., Okuda, S., Kawano, S. et al. (2017) The ProteomeXchange consortium in 2017: supporting the cultural change in proteomics public data deposition. *Nucleic Acids Res.*, 45, D1100–D1106.
16. Pertea, M., Kim, D., Pertea, G.M., Leek, J.T. and Salzberg, S.L. (2016) Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. *Nat. Genet.*, 48, 1650–1657.
17. Wang, K., Wang, D., Zheng, X., Qin, A., Zhou, J., Guo, B., Chen, Y., Wen, X., Ye, W., Zhou, Y. et al. (2019) Multi-strategic RNA-seq analysis reveals a high-resolution transcriptional landscape in cotton. *Nat. Commun.*, 10, 4714.
18. Lowe, T.M. and Chan, P.P. (2016) RNASeq-SE On-line: integrating search and context for analysis of transfer RNA genes. *Nucleic Acids Res.*, 44, W54–W57.
19. Kavalari, I., Argasinska, J., Quinones-Olvera, N., Navrocki, E.P., Rivas, E., Eddy, S.R., Bateman, A., Finn, R.D. and Petrov, A.I. (2018) Rfam 13.0: shifting to a genome-centric resource for non-coding RNA families. *Nucleic Acids Res.*, 46, D335–D342.
20. Benson, G. (1999) Tandem repeats finder: a program to analyze DNA sequences. *Nucleic Acids Res.*, 27, 573–580.
