

Data in Brief

The draft genome of *Corchorus olitorius* cv. JRO-524 (Navin)

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ABSTRACT

Here, we present the draft genome (377.3 Mbp) of *Corchorus olitorius* cv. JRO-524 (Navin), which is a leading dark jute variety developed from a cross between African (cv. Sudan Green) and indigenous (cv. JRO-632) types. We predicted from the draft genome a total of 57,087 protein-coding genes with annotated functions. We identified a large number of 1765 disease resistance-like and defense response genes in the jute genome. The annotated genes showed the highest sequence similarities with that of *Theobroma cacao* followed by *Gossypium raimondii*. Seven chromosome-scale genetically anchored pseudomolecules were constructed with a total size of 8.53 Mbp and used for synteny analyses with the cocoa and cotton genomes. Like other plant species, *gypsy* and *copia* retrotransposons were the most abundant classes of repeat elements in jute. The raw data of our study are available in SRA database of NCBI with accession number SRX1506532. The genome sequence has been deposited at DDBJ/EMBL/GenBank under the accession LLWS00000000, and the version described in this paper will be the first version (LLWS01000000).

Specifications

Organism/cell line/tissue	Dark jute (<i>Corchorus olitorius</i> cv. JRO-524)/leaves
Sex	Hermaphrodite
Sequence or array type	Illumina MiSeq
Data format	Raw and processed
Experimental factors	The draft genome sequence of <i>Corchorus olitorius</i> cv. JRO-524 (Navin)
Experimental features	DNA was extracted from seedling leaves of <i>C. olitorius</i> cv. JRO-524, and shotgun libraries were prepared followed by paired-end sequencing on an Illumina MiSeq platform, generating 2 × 250 bp overlapping reads. The cleaned sequence reads were merged with PANDASeq and assembled <i>de novo</i> using Newbler software.

Genes were predicted by FGENESH and annotated using BLASTx against the NCBI non-redundant protein database. We used SyMap for pairwise synteny mapping and ALLMAPS to integrate our draft genome with a RAD-SNP-based genetic map of *C. olitorius*.

Consent	N/A
Sample source location	Barrackpore, Kolkata, India (22°46'2.7372" N 88°23'18.0384" E)

1. Direct link to deposited data

<http://www.ncbi.nlm.nih.gov/bioproject/PRJNA278717> for *Corchorus olitorius* cv. JRO-524 (<http://www.ncbi.nlm.nih.gov/sra/SRX1506532>). (<https://www.ncbi.nlm.nih.gov/biosample/SAMN04160039>).

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2. Introduction

Corchorus olitorius L. ($2n = 2 \times = 14$; Malvaceae s. l.), commonly known as dark jute or jute mallow, is an important ligno-cellulosic bast fibre crop, with > 80% acreage of jute growing areas of the world. Grown in tropical lowland areas, it produces one of the strongest vegetable fibres and is only next to cotton in terms of production [1]. Though it is ideally suited for transplanted paddy-based crop rotation and makes softer and stronger fibre than its other cultivated counterpart *C. capsularis* (white jute), there are several biological constraints that limit its diversified uses in textile industry [2]. Besides yield enhancement, there is an urgent need to develop dark jute varieties with quality fibre in terms of fibre fineness and tensile strength including low-lignin content using genomics-assisted breeding approaches. Recently, the draft genome sequence of *C. olitorius* cv. O-4 has been released by Bangladesh [3]. However, the variety sequenced by Bangladesh is a pure line selection from a local landrace [4]. Since *C. olitorius* originated in Africa [5] and reached India together with many African crops in prehistory [6], it is of potential interest to decode one of its genomes that represents an admixture of both African and Indian gene pools. In this study, we sequenced a leading Indian variety JRO-524 (Navin), which was developed from a cross between African (cv. Sudan Green from Sudan) and indigenous (cv. JRO-632; a local selection) types. Our results provide new insights into the *C. olitorius* genome, and its availability would not only facilitate jute research and development, but also foster the application of translational genomics in jute improvement.

3. Experimental design, material and methods

3.1. Plant material and DNA isolation

Seeds of *C. olitorius* cv. JRO-524 were germinated in petri dishes and leaves were collected from 10-day-old seedlings. Twenty leaves collected from ten seedlings were pooled and used for DNA extraction using the GenElute™ Plant Genomic DNA Miniprep Kit (Sigma-Aldrich Co., St. Louis, USA).

3.2. Genome sequencing, de-novo assembly and annotation

DNA was fragmented using the Covaris AFA™ system (Covaris, Inc., Woburn, USA) with a median fragment size of 544 bp, and shotgun libraries were prepared using the Illumina TruSeq DNA PCR-Free Sample Preparation Kit (Illumina, San Diego, USA). Paired-end sequencing was performed on two flow cells of an Illumina MiSeq (2×250 bp) platform. The sequence reads were quality-checked using FASTQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>). Following adapter trimming, the poor-quality bases were removed using Trimmomatic v0.36 [7]. The genome size was evaluated using the K-mer Analysis Toolkit (KAT) [8]. High-quality reads were merged using PANDASeq v2.7 [9], and then assembled *de novo* using Newbler v. 2.6 with default parameters (Roche Inc. Germany). We used the FGENESH gene prediction pipeline from the software package Molquest v4.5 (<http://www.softberry.com>) for the *in silico* prediction of genes. The predicted genes were annotated using BLASTX ($E < 10^{-6}$) search against the NCBI non-redundant (nr) protein database.

3.3. Synteny mapping and pseudomolecule construction

SyMap v3.4 [10] was used for pairwise synteny mapping with cocoa (*Theobroma cacao*) and diploid cotton (*Gossypium raimondii*) that showed the highest sequence similarities with our assembled *C. olitorius* genome during the BLAST similarity search. For the construction of seven chromosome-scale pseudomolecules, we used ALLMAPS [11] to integrate the genome assembly with a RAD-SNP-based genetic map of *C. olitorius* [12].

3.4. Identification of disease resistance-like and defense response genes

The disease resistance-like (R-like) and defense response (DR) genes were manually categorized using different keywords/phrases that represent R-like and DR genes into five main classes as follows: (i) NBS-LRR (matching with NBS-LRR, but not with LZ-NBS-LRR and LRR, CC-NBS-LRR, *Pib*, *Pita*, *Rp 1-d8*, *Lr10*, *Mla 1* and rust resistance), (ii) LZ-NBS-LRR (matching with LZ-NBS-LRR, but not with NBS-LRR, CC-NBS-LRR, LRR and RPM1), (iii) LRR-TM (matching with *Xa21*, serine/threonine kinases and *Cf2/Cf5* resistance), (iv) LRR (matching with disease resistance, viral resistance, *Yr10*, LRR, but not with NBS-LRR, CC-NBS-LRR, LZ-NBS-LRR), and (v) defense response genes (matching with glucanases, chitinases and thaumatin like genes) [13]. We mapped these R-like and DR genes to an integrated RAD-SNP-based genetic map of jute [12].

3.5. Repeat elements and SSR identification

All assembled contigs were screened for the presence of simple sequence repeats (SSRs) using MISA (<http://pgrc.ipk-gatersleben.de/misa/>). The assembled contigs were analyzed to identify repeat sequences using RepeatModeler and RepeatMasker with Repbase library v22.01 [14].

4. Data description

Illumina MiSeq sequencing generated 52,507,986 overlapping 2×250 bp paired-end raw reads (~15.65 Gbp sequence) that were processed to yield 24,996,514 merged high-quality reads with an average read length of 450 bp (~12.9 Gbp) and a $31.32 \times$ coverage of the K-mer based estimated 415 Mbp genome of *C. olitorius* cv. JRO-524. The longer merged reads from Illumina MiSeq platform facilitated economical *de-novo* assembly of jute genome into 52,373 contigs (377.3 Mbp) covering 90.8% of the estimated genome size. The mean contig size was 7206 bp, while the N50 size was 16,573 bp (Table 1). The raw sequence data are available in NCBI SRA database with accession number SRX1506532, and the assembled genome sequence has been deposited at DDBJ/EMBL/GenBank with the accession number LLWS00000000 vide BioProject PRJNA278717 and BioSample SAMN04160039. We predicted 76,881 gene models, with an average and the largest gene size of 1.3 kbp and 37 kbp, respectively. In total 59,531 (77.4%) of the predicted genes were annotated using BLASTx, while 17,350 genes (22.6%) remained non-annotated and were thus unique to *C. olitorius* cv. JRO-524 genome. Of these, 57,087 were protein-coding genes with annotated functions. The predicted genes showed the highest sequence similarity with that of *T. cacao* (37.45%), followed by *G. raimondii* (9.68%). Using a restriction site-associated DNA (RAD)-SNP linkage map, we have shown earlier that *C. olitorius* has the maximum syntenic relationship with cocoa followed by diploid cotton [12]. Recently, Islam et al. [3] have also reported the same

Table 1
Summary statistics of *de novo*-assembled draft genome of *C. olitorius* cv. JRO-524.

Index	Statistics
Raw reads	52,507,986
High-quality merged reads	24,996,514
Number of assembled contigs	52,373
Size of assembled contigs (bp)	377,376,943
Longest contig (bp)	177,749
Shortest contig (bp)	500
Number of contigs > 1 kb	41,086
Number of contigs > 10 kb	11,958
Number of contigs > 100 kb	38
Mean contig size (bp)	7206
Contig N50 (bp)	16,573

Table 2

Summary of seven chromosome-scale pseudomolecules of *C. olitorius* cv. JRO-524. The assembled genome was integrated with a RAD-SNP-based genetic map of *C. olitorius* [12] and anchored contigs were joined together with 50 Ns to generate the chromosome-scale pseudomolecules.

Chromosome	No. of RAD-SNP markers in genetic map	No. of mapped RAD-SNP markers in genome	No. of anchored contigs	Size of anchored contigs (bp)
Chr1	139	139	76	2,336,828
Chr2	119	119	65	1,979,308
Chr3	114	114	69	2,035,515
Chr4	48	47	38	742,950
Chr5	32	32	17	582,942
Chr6	29	29	6	400,300
Chr7	22	21	17	441,461
Total	503	501	288	8,519,304

pattern of syntenic relationship for *C. olitorius*. In the present study, 501 (99.6%) of the published RAD-SNP markers were mapped to 288 contigs (8.53 Mbp) of the draft genome (Table 2).

Further, we annotated 1765 genes with disease resistance (R-like) and defense response (DR) functions. Of the total R-like and DR genes, 831 (47.1%) belong to LRR-TM, 440 (25%) to NBS-LRR, 352 (19.9%) to LRR and 44 (2.49%) to LZ-NBS-LRR categories. Further, we identified 87 (4.9%) DR genes and categorized them into three sub-categories of chitinases (40 genes), glucanases (28 genes) and thaumatin-like proteins (19 genes).

In the genome of *C. olitorius* cv. JRO-524, 51.9% of the repeat elements were masked, which was much higher than that reported for its closest related published genome of *T. cacao* (25.7%) [15], but less than that its second-closest related species of *G. raimondii* (57.0%) [16]. Expectedly, our assembled jute genome was characterized by much higher proportion of retro-transposons (45.7%) than DNA transposons (5.5%). The most dominant classes of transposable elements (TEs) were identified as *gypsy* (34.3%) and *copla* (5.7%) that belongs to the LTR superfamily. Earlier, Begum et al. [17] have also predicted high number of LTR retro-transposons in jute. Further, we identified a total of 185,698 genomic SSRs, with mononucleotide repeats being the most abundant class (76.0%), followed by di- (16.0%), tri- (5.7%), tetra- (0.8%), penta- (0.2%) and hexa-nucleotide (0.2%) repeats.

Using genetically anchored contigs seven chromosome-scale pseu-

domolecules were constructed with a mean size of 1,219,051 bp and N50 of 2,038,915 bp (Table 2). Chromosome1 was the longest, while chromosome 6 was the shortest pseudomolecule. Comparative analysis of seven genetically anchored jute chromosomes with 10 chromosomes of *T. cacao* [15] revealed significant syntenic relationship between the two species, however, collinearity was not conserved (Fig. 1). Jute chromosomes 1, 4 and 7 showed synteny with cocoa chromosomes 9, 5 and 2, respectively, whereas chromosome 2 shared synteny with cocoa chromosomes 3 and 10 and chromosome 3 with cocoa chromosomes 4 and 2. However, jute chromosomes 5 and 6 shared synteny with a single cocoa chromosome 1. Similarly, comparative analysis of jute and diploid cotton species *G. raimondii* [16] revealed synteny of jute chromosomes 6 and 7 with cotton chromosomes 4 and 13, respectively (Fig. 1), with chromosomes 1, 2 and 3 showing matches with multiple chromosomes of cotton, viz., (1, 4, 9 and 10), (3, 4, 8 and 11) and (5, 6, 7 and 12), respectively. Thus comparative analysis with a small fraction (8.53 Mbp) of genetically anchored jute genome revealed chromosomal level synteny of jute with both cocoa and cotton genomes.

5. Conclusions

To our knowledge, the work presented here is the first whole genome sequence for a *C. olitorius* genotype derived from an African jute. *C. olitorius* cv. Sudan Green, one of the parents of cv. JRO-524, was primarily used to transfer premature flowering resistance (in early sowing) to indigenous types [18]. Thus an in-depth comparison of the present sequence with the recently published draft genome [3], would provide new insights that could help understand the mechanisms underlying premature flowering *vis-à-vis* photoperiodic control of bast fibre development in jute. This would allow breeding of high-yielding varieties with durable premature flowering resistance, which has been recently observed to be breaking down when dark jute crops are sown early under long-day conditions, possibly due to climate change.

Conflict of interest

The authors declare that they have no conflict of interests.

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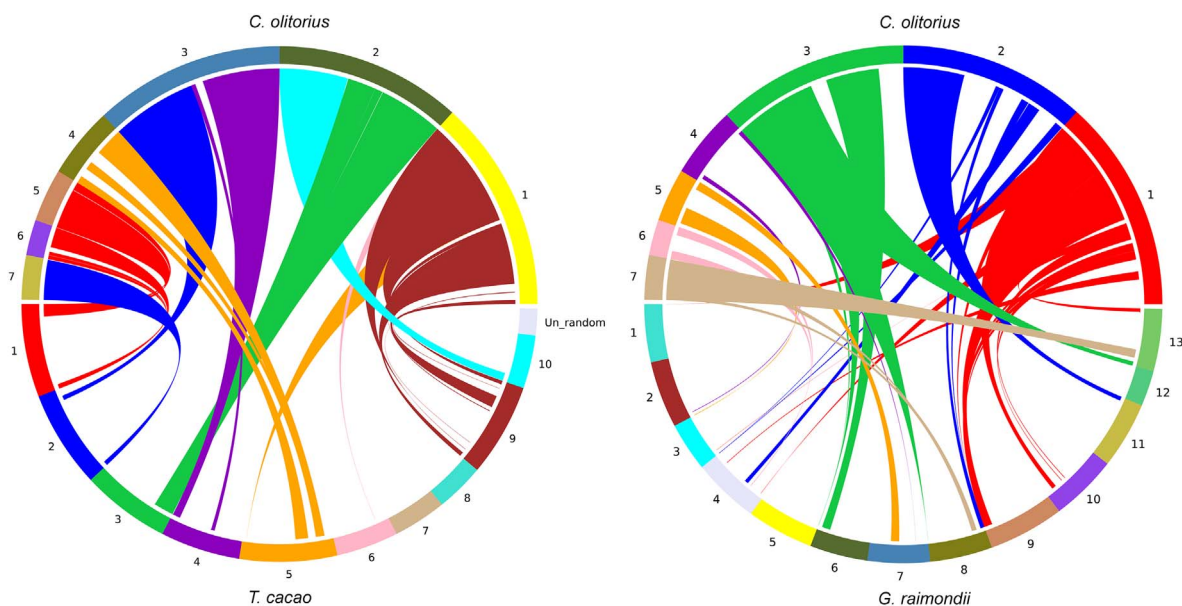


Fig. 1. Genomic syntenic relationships of *C. olitorius* ($2n = 2x = 14$) with *T. cacao* ($2n = 2x = 20$) and *G. raimondii* ($2n = 2x = 26$).

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