A Chromosome-Level Genome Assembly of the Parasitic Wasp

Chelonus formosanus Sonan 1932 (Hymenoptera: Braconidae)

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Abstract

Chelonus formosanus Sonan 1932 (Hymenoptera: Braconidae) is a wasp capable of parasitizing a variety of lepidopteran pests at the “egg-larval” stage which distributes throughout Taiwan, Guangdong, Zhejiang, and Hainan provinces of China. This wasp has been successfully used to control pests such as Spodoptera litura Fabricius, 1775, Spodoptera frugiperda (JE Smith, 1797), Spodoptera exigua (Hübner, 1808), and Helicoverpa armigera (Hübner, 1808). So far, there is only one genome assembled from the Chelonus genus [Chelonus insularis (Cresson, 1865)] and it is fragmented with 455 scaffolds. Here, we report a chromosome-level genome assembly of C. formosanus, which was sequenced using PacBio, Illumina, and Hi-C technologies. The long reads were 35.4 Gb (~150× coverage) with an average length of 15.23 kb. The size of the genome assembly was 139.59 Mb. More than 99.46% of the assembled sequences were anchored to seven pseudochromosomes (138.84 Mb). The Benchmarking University Single-Copy Orthologs (BUSCO) assessment results showed 99.0% of the 1,367 genes (insect_odb10 database) were completely present. We annotated 11,242 protein-coding genes including 98.6% of BUSCO complete genes that were recovered. There were 58 gene families found with significant expansion including allelopathic families (odorant receptors and ionotropic receptors), which may play a crucial role in efficiently locating a wide range of hosts. This high-quality genome assembly and annotation could provide a highly valuable resource of parasitic wasp for the biological control of Lepidoptera pest.

Key words: parasitic wasp, genome, comparative genomics, gene family evolution.

Introduction

The cosmopolitan genus Chelonus Panzer, 1806, harbors near 360 known species, which are ovo-larval endoparasitoids of Lepidoptera (Zhang et al. 2006). Chelonus normally regulate the metamorphic process to kill host larva during their final instar (Jones 1985; Zhang et al. 2006). The genus Chelonus have numerous associations with many Spodoptera species which include some important agricultural pests in the world (Jones 1985). So far, we found only one whole-genome assembly sequenced from the Chelonus genus (Chelonus

Significance

Chelonus formosanus is an important natural predator of agricultural pests and can parasitize a variety of lepidopteran pest species. However, at present, its genetic data are extremely limited. In order to understand the genetic background of C. formosanus more comprehensively, we sequenced its whole genome, which provided a highly valuable resource for understanding its parasitic potential and evolution.
**Results and Discussion**

**Genome Assembly**

We obtained a total of 35.4 Gb PacBio long (~150 × coverage) and 36.43 Gb Illumina short reads. The average length and N50 length of the long reads were 15.23 and 17.94 kb, respectively. The kmer analysis predicted genome size being 139.59 Mb, and it also indicated no significant heterozygosity and approximately 18 Mb (12.95%) repetitive sequences of the genome (supplementary fig. S2, Supplementary Material online). The genome assembly size, GC content, and Benchmarking University Single-Copy Orthologs (BUSCO) assessment results are comparable to the genome assembly of the closely related species *C. insularis* ([table 1](#table1)). However, our genome assembly was more complete with smaller number of scaffolds and more contiguous with fewer gaps compared with *C. insularis*. The mapping-back rates from the Illumina DNA and RNA sequences as well as the PacBio raw reads were 98.29%, 96.92%, and 96.02%, respectively. Overall, our *C. formosanus* genome scaffolds have recovered most of sequencing reads and is suitable for further analysis. According to the long-range linked reads from Hi-C data, we assigned 138.84 Mb of the assembly to the seven pseudochromosomes (supplementary fig. S2, Supplementary Material online). A chromosomal synteny analysis between the *C. formosanus* and *Aphidius gifuensis* Ashmaed, 1906 chromosomes showed limited level of conservation. We also noticed that there was no indication of conservation between the chromosome one of *C. formosanus* and the *A. gifuensis* genome (fig. 1a).

To investigate this further, more chromosomal conservation analysis with other closely related species may be applied when the chromosome-level genome assemblies of those species became available.

**Genome Annotation**

There were 22.25% of the assembly annotated as repetitive sequences. This is over 4% larger than the prediction form the kmer analysis, which is likely due to a better capture of repetitive sequences from long-read assembly. Except for the unclassified repeats (8.62%), DNA elements were the most abundant repeat type (6.19%) in the assembly, followed by the long-terminal repeat elements (LTR; 3.42%), simple repeats (1.79%), and long-interspersed nuclear elements (0.91%) (supplementary table S1, Supplementary Material online). We also predicted 324 noncoding RNAs (ncRNAs) including 38 micro-RNAs (miRNAs), 69 ribosomal RNAs (rRNAs), 28 small nuclear RNAs (snRNAs), 108 transfer RNAs (tRNAs), and 44 others (supplementary table S2, Supplementary Material online). The annotated snRNAs include 14 spliceosomal RNAs (U1, U2, U4, U5, U6, U11), three minor spliceosomal RNAs (U4atac, U6atac, U12), six C/D box snoRNAs, and four H/ACA box snoRNA.

A total of 4.42-Gb RNA-Seq reads were imported into the gene prediction program MAKER as biological evidence for the protein-coding gene prediction. The MAKER process predicted 11,242 protein-coding gene models, which was comparable to that of *C. insularis* (11,574). The average gene and protein-coding region lengths were 4,350 and 1,593 bp, respectively. The average exon length and the average number of exons per gene were 355.13 bp and 5.73, respectively. The average intron length was 522.85 bp. The BUSCO assessment result showed 98.6% complete genes were captured, and 0.4% and 1.0% of the genes were fragmented and missing, respectively. There were 9,019 (80.23%) protein-coding genes identified with protein domains, which were then assigned with 7,822 GO terms, 6,157 KEGG KO terms, 2,020 enzyme codes, 3,576 KEGG pathways, 2,364 Reactome pathways, and 9,071 COG categories (supplementary figs. S3 and S4, Supplementary Material online).

**Table 1**

| Genome Assembly Statistics of *Chelonus formosanus* Compared with *Chelonus insularis* |
|---------------------------------|---------------------------------|
| **Chelonus formosanus** | **Chelonus insularis** |
| Genome assembly | 139.590 | 135.730 |
| Assembly size (Mb) | 26/106 | 455/457 |
| Number of scaffolds/contigs | 24.95/15.194 | 4.699/4.699 |
| Longest scaffold/contig (Mb) | 24.15/5.591 | 1.163/1.163 |
| N50 scaffold/contig length (Mb) | 30.36 | 30.53 |
| GC (%) | 0.006 | 0.043 |
| Gaps (%) | BUSCO completeness (%) | 99.0 | 99.1 |

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Phylogenomics and Gene Family Evolution

A total of 184,525 (94.9%) genes obtained from 16 species were clustered into 13,537 gene families. There are 4,640 gene families with all the species sequences present including 1,702 single-copy and 2,938 multicopy gene families. Among the 11,242 annotated C. formosanus genes, 10,556 were clustered into 8,937 families (fig. 1b). There were 68 genes present in 48 families that are specific to C. formosanus (fig. 1b). A phylogenetic tree with the bootstrap support value of 100/100 was reconstructed based on the 1,581 genes after 121 single-copy genes were removed (fig. 1b). The topology of this phylogeny shows consistency with the previous phylogenetic tree constructed by Peters et al. (2017). For example, parasitoid Apocrita belongs to a monophyletic group and forms a sister group to Orussoidea (Orussus abietinus), and Aculeata, Chalcidoidea, and Ichonoidea are all monophyletic groups (Gauthier et al. 2021). As expected, our analysis showed C. formosanus was closely clustered with C. insularis (fig. 1b) and our calculation indicated the two species diverged approximately 6 Ma.

We identified 355 expanded (58 significantly expanded) and 383 contracted (28 significantly contracted) gene families from the C. formosanus annotated gene models (fig. 1c). The top 15 significantly expanded gene families with numbers of genes in each family shown above each bar.

**Fig. 1.**—Chromosomal synteny, phylogeny, and gene family evolution of Chelonus formosanus. (a) Chromosomal synteny between Chelonus formosanus (CfChr) and Aphidius gifuensis (AgChr) genomes. (b) In the phylogeny, the node values on the tree represent the number of expanded, contracted, and rapidly evolving families for each clade or species. Statistics of orthology inference result: “1:1:1” indicates single-copy orthologs; “N:N:N” indicates multicopy orthologs; “Braconidae” indicates orthologs are specific to Braconidae; “Others” indicates unclassified orthologs; “Unassigned” indicates orthologs that cannot be assigned into any of the orthogroups. (c) The top 15 significantly expanded gene families with numbers of genes in each family shown above each bar.
Materials and Methods

Sample Collection and Sequencing

*Chelonus formosanus* specimens were collected in June 2020 within the Guiyinyang Economic Development Zone, Haikou City, Hainan Province, China (20.0521°N, 110.2067°E) and then reared with *Spodoptera frugiperda* for more than five generations under the laboratory conditions of 26 ± 1 °C, 70 ± 5% RH, and a photoperiod of 14:10 (L:D) h. Male adults were used for genome sequencing: four individuals for Illumina, 20 for PacBio, and five for Hi-C. Four males were used for RNA-Seq (supplementary fig. S1, Supplementary Material online).

The high-quality DNA was extracted using the QIAGEN DNeasy Blood & Tissue kit. For the PacBio sequencing, a 20-kb insert size library was constructed using the SMRTbell Template Prep Kit 2.0. For Illumina DNA sequencing, a library with an insert size of 350 bp was constructed using the TruSeq DNA PCR-free kit. The Hi-C library construction (restriction enzyme: Mbol) was performed by Frasergen Co., Ltd (Wuhan, China). RNA was extracted using the TRIzolTM Reagent kit and the RNA libraries were constructed using the TruSeq RNA v2 kit. The Illumina and PacBio sequencing were performed on NovaSeq 6000 and PacBio Sequel II, respectively, at the Beijing Berry Genomics Co., Ltd (Beijing, China).

Genome Assembly

The quality control of the Illumina short reads was performed using BBTools suite v38.49 (Bushnell 2014): the script “Clumpify.sh” was used to remove duplicated sequences; “bbduk.sh” was used for trimming sites with base quality scores below 20 (≥Q20) and poly-A/CG/C ends with their lengths being more than 10 bp, filtering sequences with lengths below 15 bp, and correcting bases according to the sequence overlap regions (qtrim=minqm trimq=20 minlen=15 ecco=10mna trimpolya=10 trimpolyg=10 trimpolyc=10). The k-mer frequency was calculated using the “khist.sh” script from the BBTools suite (kmer: 21). A genomic survey based on the k-mer distribution frequency was performed using Genomescope v2.0 (Vurture et al. 2017) with the parameters of “-k 21 -p 2 -m 10,000.”

The long-read assembler Flye v2.8.1 (Kolmgorov et al. 2019) was used to generate a preliminary genome assembly with the parameters “-i 2 -m 1,000” (two rounds of long-read polishing with a minimum overlap length of 1,000 bases between sequences). The Illumina reads were then aligned to the preliminary assembly using Minimap2 v2.17 (Li 2018) with default parameters and the alignments were used for the two consecutive rounds of short-read polishing with NextPolish v1.3.0 (Hu et al. 2020). The haplotigs and overlaps in the genome assembly were filtered based on the read depth using Purgedups v1.0.1 (Guan et al. 2020) with the minimum alignment score of 70 (-a 70). The remaining contigs were then assigned to pseudochromosomes based on the contact information from the read alignments of the Hi-C data: first, the raw Hi-C reads were quality-assessed and then removed unusable reads using Juicer v1.6.2 (Durand et al. 2016); second, the pseudochromosomal assignment was performed using 3D-DNA v180922 (Dudchenko et al. 2017); third, the assignment errors were corrected using Juicebox v1.11.08 based on the Hi-C contact maps (Durand et al. 2016). The contaminant sequences were removed using BLAST+ (BlastN) v2.7.1 (Camacho et al. 2009) based on the homologous search against the NCBI nucleotide (nt; downloaded on 31st of December 2020) and UniVec databases. Genome completeness was assessed using BUSCO v3.0.2 pipeline (Waterhouse et al. 2018) by searching against the database of insect_odb10 database (n = 1,367).

To construct a chromosomal synteny between the *C. formosanus* and *A. gifuensis* pseudochromosomes, a BlastP-like alignment method was performed using Mmseq2 v11-e1a1c with default parameters for aligning protein sequences. The generated “all.blast” file and the integrated “all.gff” file were imported in MCScanX to perform collinearity analysis. A circus plot was created using Ttools v1.0692 (Chen et al. 2020).

Genome Annotation

The genome assembly was annotated for repetitive sequences, protein-coding genes, and ncRNAs. To annotate repeats, a de novo repeat library was constructed using RepeatModeler v2.0.1 (Flynn et al. 2020) with the LTR search process (LTRStruct). It was then combined with Dfam3.3 (Hubley et al. 2016) and the RepBase-20181026 databases (Bao et al. 2015) to form a custom library, which was used as input for RepeatMasker v4.0.9 (Smit et al. 2013–2015) to search for repeats and generate a repeat-masked assembly. To annotate ncRNAs, the rRNAs, snRNAs, and miRNAs were identified based on the alignment with the Rfam library using Infernal v1.1.2 ( Nawrocki and Eddy 2013), and the tRNAs were predicted using trNAScan-SE v2.0.6 (Chan and Lowe 2019) and then filtered low-confidence sequences using the “EukHighConfidenceFilter” script.

Protein-coding genes were predicted using MAKER v.3.01.03 (Holt and Yandell 2011), with three supporting evidence files produced from other programs: 1) Ab initio predicted genes generated from BRAKER v2.1.5 (Hoff et al. 2016), which trains Augustus v3.3.2 (Stanke et al. 2004) and GeneMark-ES/ET/EP 4.483.6olc (Lomsadze et al. 2005) based on the RNA-Seq alignments generated from HISAT2 v2.2.0 (Kim et al. 2019) and the OrthoDB10 v1 protein database (Kriventseva et al. 2019) to improve prediction accuracy; 2) Transcript sequences assembled using StringTie v2.1.4 (Kovaka et al. 2019) from the RNA-Seq alignments generated.
by HISAT2; 3) Protein sequences of closely related species [Drosophila melanogaster Meigen, 1830, Tribolium castaneum (Herbst, 1797), Apis mellifera L., 1761, Bombyx terrestris (L. 1758), Nasonia vitripennis (Walker, 1836), and Bombyx mori L., 1758] downloaded from NCBI. Gene functions were annotated using Diamond v0.9.24 (Buchfink et al. 2015) with the sensitive mode (–more-sensitive –e 1e-5) to search against UniProtKB and using InterProScan 5.41–78.0 (Finn et al. 2017) to search against Pfam (El-Gebali et al. 2019), Smart (Letunic and Bork 2018), Gene3D (Lewis et al. 2018), Superfamily (Wilson et al. 2009), and CDD (Marchler-Bauer et al. 2017) databases. The eggnog-mapper v2.0.1 (Huerta-Cepas et al. 2017) was also used to search against the eggnog v5.0 (Huerta-Cepas et al. 2019) database to predict conserved sequences and domains, GO terms, and protein pathways (KEGG, Reactome).

Phylogenomics and Gene Family Evolution
There were 16 species selected for the orthology inference using OrthoFinder, which used Diamond v2.3.8 for rapid protein sequence aligning (Emms and Kelly 2019). These species included Rhopalosiphum maidis (Fitch, 1856) from Heteroptera, Tribolium castaneum (Herbst, 1797) from Coleoptera, Drosophila melanogaster Meigen, 1830 from Diptera, Athalia rosae (Linnaeus, 1758) and Orussus abietinus (Scopoli, 1763) from Symphyta, Apis mellifera, B. terrestris, Oecophora biroi (Forel, 1907) and Vespa mandarina Smith, 1852 from Aculeata, and seven parasitic wasp species [Aphidius gifuensis Ashmaed, 1906, Belonocnema treatae Mayr, 1881, C. formosanus, C. insularis, Fopius arisanus (Sonan, 1932), N. vitripennis, and Trichogramma pretiosum Riley, 1879]. A species phylogeny was constructed using 1,702 single-copy orthologs as following: first, regions of homologous sequences were aligned using MAFFT v7.394 (Katoh and Standley 2013) with the option of L-INS-I; second, the unreliable regions from the alignments were trimmed using BMGE v1.12 (Criscuolo and Gribaldo 2010) with the parameter of “-m BLOSUM90-h 0.4.”; third, the modified alignments were combined to a supermatrix using FASCoCAT-G v1.04. Phylogenetic tree construction was performed using IQ-TREE v2.0-rc1 (Minh et al. 2020) with the parameters of “-syntest-remove-bad -syntest -pval 0.10” to remove sequences that did not meet the substitution, reversible, or homogeneous hypotheses. To reduce computational resources, the model type was limited to LG (-m MFP –rtree 10). The bootstrap values were calculated using ultrafast bootstrap and the SH-aLRT algorithm (-B 1000 –aLRT 1000). The clock dating of species divergence was performed using MCMCtree (clock = 2, BDparas = 1 1 0 1, kappa_gamma = 6 2, alpha_gamma = 1 1, rgene_gamma = 2 20 1, sigma2_gamma = 1 10 1) from the PAML v4.9 package (Yang 2007). The evidence for the fossil calibration points was obtained from the PDDB database (https://www.paleobiodb.org/navigator/, last accessed July 11, 2021): Trichoptera, (3.114–3.146 Ma); Hymenoptera, (2.056–2.12 Ma); Aculeata, (1.402–1.45 Ma); Chalcidoidea, (0.935–0.996 Ma); and Ichneumonoidea, (1.402–1.45 Ma). The estimation of gene family expansion and contraction was performed using CAFÉ v4.2.1 (Han et al. 2013) with the p parameter of 0.01. The R package “clustering profiler” v3.10.1 (Yu et al. 2012) with default parameters was used to analyze and visualize the enriched GO ontology terms and KEGG pathways of the significantly expanded gene families.

Supplementary Material
Supplementary data are available at Genome Biology and Evolution online.

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Data Availability
Genome assembly and raw sequencing data have been deposited at the NCBI under the accessions JAEFX000000000 and SRR13334670–SRR13334673, respectively. Genome annotations are available at the Figshare under the link: https://figshare.com/articles/online_resource/A_Draft_Genome_Assembly_of_Cheilonus_formosanus_Sonan_Hymenoptera_Braconidae_Using_PacBio_Sequencing/16417497.

Literature Cited
Bao W, Kojima KK, Kohany O. 2015. RepBase Update, a database of repetitive elements in eukaryotic genomes. Mol DNA. 6:11.
Buchfink B, Xie C, Huson DH. 2015. Fast and sensitive protein alignment using DIAMOND. Nat Methods. 12(1):59–60.
Bushnell B. 2014. BBtools. Available from: https://sourceforge.net/projects/bbmap/.
Camacho C, et al. 2009. BLAST+: architecture and applications. BMC Bioinformatics 10(1):421.
Chan PP, Lowe TM. 2019. tRNAscan-SE: searching for tRNA genes in genomic sequences. Methods Mol Biol. 1962:1–14.
Chen C, et al. 2020. TBtools: an integrative toolkit developed for interactive analyses of big biological data. Mol Plant. 13(8):1194–1202.
Criscuolo A, Gribaldo S. 2010. BMGE (Block Mapping and Gathering with Entropy): a new software for selection of phylogenetic informative regions from multiple sequence alignments. BMC Evol Biol. 10(1):210.
