Genome and transcriptome assemblies of the kuruma shrimp, *Marsupenaeus japonicus*

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Abstract

The kuruma shrimp *Marsupenaeus japonicus* (order Decapoda, family Penaeidae) is an economically important crustacean that occurs in shallow, warm seas across the Indo-Pacific. Here, using a combination of Illumina and Oxford Nanopore Technologies platforms, we produced a draft genome assembly of *M. japonicus* (1.70 Gbp; 18,210 scaffolds; scaffold N50 = 234.9 kbp; 34.38% GC, 93.4% BUSCO completeness) and a complete mitochondrial genome sequence (15,969 bp). As with other penaeid shrimp genomes, the *M. japonicus* genome is extremely rich in simple repeats, which occupies 27.4% of the assembly. A total of 26,381 protein-coding gene models (94.7% BUSCO completeness) were predicted, of which 18,005 genes (68.2%) were assigned functional description by at least one method. We also produced an Illumina-based transcriptome shotgun assembly (40,991 entries; 93.0% BUSCO completeness) and a PacBio Iso-Seq transcriptome assembly (25,415 entries; 67.5% BUSCO completeness). We envision that the *M. japonicus* genome and transcriptome assemblies will serve as useful resources for the basic research, fisheries management, and breeding programs of *M. japonicus*.

Keywords: penaeidae; penaeid shrimp; kuruma prawn; *Penaeus japonicus*; *Marsupenaeus japonicus*; genome sequencing; genome annotation; transcriptome assembly

Introduction

The kuruma shrimp *Marsupenaeus japonicus* (order Decapoda, family Penaeidae) is an economically important crustacean that occurs across the Indo-West Pacific. In Japan, *M. japonicus* is especially highly prized as a seafood delicacy and has been a major fisheries and aquaculture target since the early 20th century. Artificial spawning and hatching of *M. japonicus* was achieved in 1933 (Hudinaga 1942), and an industrial-scale larval rearing technology was established by the 1960s (Liao 1985). Today, *M. japonicus* farming is practiced in several other countries in Asia and Europe, with China being the largest producer yielding 55,228 tons in 2018 (accessed April 2021; http://www.fao.org/fishery/statistics/).

Genomic resources of an organism provide powerful tools for investigating its basic biology, including economically important aspects such as growth, reproduction, and immunity. Genomic information is also important for developing molecular markers used in resource management and selective breeding programs. High-quality genome assemblies of three economically important shrimps (*Litopenaeus vannamei* [Zhang et al. 2019]; *Penaeus monodon* [Uengwetwanit et al. 2021]; *Fenneropenaeus chinensis* [Yuan et al. 2021]) have been recently reported. In contrast, the genomic resources of *M. japonicus* are severely limited. Although a previous study reported an Illumina-based draft genome assembly of *M. japonicus* (Yuan et al. 2018), this assembly is severely fragmented (2,434,740 scaffolds; scaffold N50: 912 bp), reflecting the repetitiveness and complexity of penaeid shrimp genomes.

Here, we present a high-quality draft genome assembly of *M. japonicus* generated using a combination of Illumina and Oxford Nanopore Technologies platforms. We also generated a transcriptome shotgun assembly and a PacBio Iso-Seq transcriptome assembly, which will complement the draft genome by capturing full-length transcript structures and genes missing from the genome assembly. We envision that the *M. japonicus* genome and transcriptome assemblies will serve as a useful resource for the basic research, fisheries management, and breeding programs of *M. japonicus*.

Materials and methods

General sequencing, assembly, and annotation strategy

We first generated an initial assembly using Illumina paired-end reads, and the assembly was scaffolded using Illumina paired-end reads. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs licence (http://creativecommons.org/licenses/by-nc-nd/4.0/), which permits non-commercial reproduction and distribution of the work, in any medium, provided the original work is not altered or transformed in any way, and that the work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
and mate-pair reads. We mapped RNA-seq reads onto the genome and took forward the mapped reads to generate a transcriptome shotgun assembly. The genome was further scaffolded using the transcriptome shotgun assembly, Iso-Seq cDNA sequences, and the RNA-seq reads. We improved this Illumina-based primary assembly by gap-filling using ONT long reads and scaffolding using ONT long reads, Illumina mate-pair reads, transcriptome shotgun assembly, and PacBio Iso-Seq cDNA sequences. See Supplementary Figure S1 for a summary of the assembly workflow.

To annotate protein-coding genes from the finished assembly, we combined multiple lines of evidence (de novo assembled transcripts, Iso-Seq cDNA sequences, RNA-seq reads, protein alignments, and ab initio predictions) into a consensus gene model set. We discarded low-quality predictions with no detectable similarity to known proteins. To maximize the recovery of gene models from transcript evidence, we again mapped the transcriptome shotgun assembly and Iso-Seq cDNA sequences to the genome. The gene models were again merged and filtered by homology search, deriving the final protein-coding gene model set. tRNA, rRNA, and other noncoding RNA genes were also predicted and included in the annotation file. Supplementary Figure S2 summarizes the annotation workflow.

**Marsupenaeus japonicus samples**

For genome sequencing, we purchased a male *M. japonicus* (Ginoza2017, BioSample: SAMD00276454, 30 g body weight) from a commercial farm in Ginoza, Okinawa Prefecture, Japan, and kept it frozen at −80°C until use. For transcriptome assembly, we sequenced a total of 49 RNA samples, including 37 adult samples covering 18 different tissues and 12 larval samples covering nauplii, zoea, and mysis stages. The *M. japonicus* larvae were obtained from a hatchery on Kumejima Island, Okinawa Prefecture. See Supplementary Table S1 for a summary of the samples used in this study.

**Illumina genomic DNA library preparation and sequencing**

For Illumina sequencing, we prepared genomic DNA from muscle and sperm. We used the muscle DNA to construct a paired-end library and the mate-pair libraries, whereas the sperm DNA was used to prepare a paired-end library. We extracted genomic DNA using the phenol-chloroform-isoamyl alcohol extraction, followed by ethanol precipitation. The DNA preparations were resuspended in Tris-EDTA buffer and were frozen at −30°C until use. For pair-end sequencing, genomic DNA from muscle and sperm were sheared using a Covaris instrument, and paired-end libraries (600-bp nominal insert) were prepared using the KAPA Hyper Prep Kit Illumina platforms (KK8504, Roche). For mate-pair sequencing, genomic DNA from muscle was size-fractionated using SageELF (0.75% 1–18 kb, SafeScience, ELD7510), and eleven mate-pair libraries (2, 3, 4, 5, 6, 7, 8, 10, 12, 14, and 15 kb nominal insert sizes) were prepared using the Nextera Mate Pair Library Prep Kit (FC-132-1001, Illumina). For polishing the assembly, we error-corrected the trimmed raw subread bam files using ccs v3.4.0, and the resulting ccs bam files were processed with lmp v1.10.0 to remove primer sequences. This was followed by Iso-Seq3 refine, cluster, and polish functions in Iso-Seq3 v3.1.2, generating the final assembly. Before submission to DDBJ/ENA/NCBI, we manually removed duplicated entries and trimmed primer sequences remaining in the assembly.

**PacBio Iso-Seq sequencing and analysis**

For PacBio Iso-Seq sequencing, we extracted total RNA using RNeasy Plus (Takara, Japan), followed by isopropanol precipitation. The RNA was resuspended in nuclease-free water and was stored at −80°C until use. Paired-end RNA-seq libraries, with an empirical fragment size of 600–700 bp, were prepared using the KAPA mRNA HyperPrep Kit Illumina Platforms (KK8581, Roche) and were sequenced on the Hiseq4000 platform using the Hiseq3000/4000 PE Cluster Kit cBot (PE-410-1001, Illumina) and HiSeq 3000/4000 SBS Kit (300 Cycles, FC-410-1003, Illumina).

**Genome size estimation and ploidy analysis**

We processed the raw Illumina paired-end reads using Fastp v0.20.1 with relaxed parameters. For genome size estimation and ploidy analysis, we combined the trimmed paired-end reads from both libraries. We used KMC v3.1.1 (Kokot et al. 2017) to tally the occurrence of 17, 21, 23, 25, 27, 29, 31, 33, and 35-mers. The resulting k-mer histograms were used for genome size estimation using GenomeScope 2.0 (Ranallo-Benavidez et al. 2020). We adopted the k-mer size of $k = 23$, which yielded the highest model fit (Supplementary Table S4). We used the same 23-mer index for ploidy analysis using SmudgePlot v0.2.3dev (Ranallo-Benavidez et al. 2020).

For polishing the assembly, we error-corrected the trimmed reads using Tadpole in BBtools v38.86 (Bushnell et al. 2017).

**De novo assembly of Illumina genomic DNA libraries**

The Illumina paired-end reads from the sperm library were trimmed with fastp v0.19.4 (Chen et al. 2018); here, we applied stringent parameters (length_required 140, qualified_quality_phred 20, unqualified_percent_limit 10, n_base_limit = 0,
The raw mate-pair reads were processed using Trimmomatic v0.36 (Bolger et al. 2014) and NextClip v1.3.1 (Leggett et al. 2014). The filtered mate-pair reads were used for the scaffolding and misassembly detection of the initial assembly.

The filtered paired-end reads from the sperm library were de novo assembled using SPAdes v3.13.0 (Nurk et al. 2013). For producing a homologous reference genome assembly, the initial assembly (“SPAdes” in Supplementary Table S5) was processed by Redundans v0.14a (Pryszzcz and Gabaldón 2016) using paired-end reads from the muscle library (preprocessed by Fastp v0.19.6; Supplementary Table S3) and the filtered mate-pair reads from all libraries (“Redundans” in Supplementary Table S5). Misassembled sequences were broken by REAPR v1.0.18 (Hunt et al. 2013) using the filtered mate-pair reads from 04 to 15 k libraries, generating respective broken assemblies for each mate-pair library (“REAPR_04k” to “REAPR_15k” in Supplementary Table S5).

**Transcriptome shotgun assembly**

The raw reads were processed using Fastp v0.12.6. The trimmed reads were aligned to the “REAPR_06k” assembly using HISAT2 v2.1.0. We processed the alignments with SAMtools (Li et al. 2009) and BEDTools bamtofastq v2.25.0 (Quinlan and Hall 2010) to extract paired-end reads that mapped to the genome. The retained paired-end reads were de novo assembled using Trinity v2.8.6. In addition, the in silico-normalized reads generated by Trinity were de novo assembled using maSPAdes v3.13.0 (Bushmanova et al. 2019) and Trans-ABYSS v2.0.1 (Robertson et al. 2010). We used a series of different k-mers (31-, 41-, 51-, 61-, 71-, 81-, 91-, 111-, and 121-mers) for Trans-ABYSS. The assemblies were concatenated into a single file, and coding sequences were predicted using TransDecoder v5.5.0 (https://github.com/TransDecoder/TransDecoder/). The predicted coding sequences labeled as “complete” by TransDecoder were extracted and clustered using cd-hit-est v4.8.1 (Li and Godzik 2006), with the following settings: -c 0.98 -G 0 -aS 0.25 -d 0 -T 0 -low_complexity_filter] with an aim to reduce the complexity of the dataset input to the assembler (Supplementary Table S3).

The predicted cod-mers (61-, 71-, 81-, 91-, 101-, 111-, and 121-mers) for Trans-ABySS. The predicted cod-mers were retained for generating the final, nonredundant transcriptome library. The transcripts harboring the longest CDS in each cluster were retained. The transcripts harboring the longest CDS in each cluster were retained. The transcriptome shotgun assembly and Iso-Seq cDNA sequences (aligned by BLAT v36). Each scaffolding was followed by gap-filling by TGS-GapCloser v1.1.1 and polishing by NtEdit v1.3.2. For the iterative scaffolding, misassembled sequences were broken by REAPR v1.0.18 using 10-kb mate-pair reads, and haplotigs were removed by purge_haplotigs v1.1.1 (Roach et al. 2018) using ONT long reads. This was followed by a second round of iterative scaffolding using LRScf v1.1.11 (Qin et al. 2019), using ONT long reads; BESST v2.2.8 (Sahlin et al. 2014), using Illumina mate-pair reads from all libraries; L_RNA_Scaffolder, using the transcriptome shotgun assembly and Iso-Seq cDNA sequences (aligned by BLAT v36). Each scaffolding was followed by gap-filling by TGS-GapCloser v1.1.1 and polishing by NtEdit v1.3.2. After the iterative scaffolding, misassembled sequences were broken by REAPR v1.0.18 using 10-kb mate-pair reads, and haplotigs were removed by purge_haplotigs v1.1.1 (Roach et al. 2018) using ONT long reads. This was followed by a second round of iterative scaffolding using LRScf, BESST, and L_RNA_Scaffolder, each scaffolding run was followed by gap-filling by TGS-GapCloser and polishing by NtEdit and a purge_haplotigs run. The third round of iterative scaffolding consisted of scaffolding by LRScf v1.1.11 and BESST v2.2.8, with each scaffolding run followed by gap-filling by TGS-GapCloser v1.1.1 and polishing by NtEdit v1.3.2. Finally, the assembly was polished by HyPo v1.0.3 (Kundu et al. 2019) using the trimmed, error-corrected paired-end reads from the muscle library.

The polished assembly was further curated. Mitogenomic contamination was identified using BLASTN (-perc_identity 90) by querying the M. japonicus mitogenome sequence (NCBI Reference Sequence: NC_007010.1) (Yamauchi et al. 2004), and two possible mitogenomic sequences showing abnormally high coverage were removed. Next, scaffolds matching at least one of the following criteria were removed: shorter than 2 kb; GC content over 70% or below 10% (which might represent bacterial contamination or assembly artifacts); gap content above 60%. Gaps occurring at scaffold ends were trimmed to expose the first contig; similarly, if a scaffold ends with a short contig (up to 100 bp) but is followed by a gap, the contig and gap were trimmed to expose the next contig. We mapped the trimmed, error-corrected paired-end reads from the muscle and sperm libraries against the assembly and calculated the read coverage (two libraries combined) of each scaffold using SAMTools coverage. Scaffolds with coverage below 30 or mean depth below 20 were discarded. To rescue genic regions, we queried the discarded scaffolds against the transcriptome shotgun assembly and recovered one scaffold, which had been discarded due to a high gap content. This scaffold was broken into contigs and returned to the assembly. We scanned for bacterial ribosomal DNA sequences by Barrnap v0.9 (https://github.com/teemann/barrnap) and queried the positive hits against the
NCBI nonredundant nucleotide database using BLASTN (Accessed February 12, 2021). All the hits were found to be eukaryotic sequences and therefore were retained in the assembly. The completeness of the finished assembly was assessed by BUSCO v4.1.4 (Simão et al. 2015) using the arthropoda_odb10 dataset.

Mitogenome assembly and annotation
ONT long reads longer than 5 kb (Supplementary Table S3) were de novo assembled by Flye v2.8-b1674 (Kolmogorov et al. 2019), and a circular mitogenome contig was recovered by querying the reference M. japonicus mitogenome (NCBI Reference Sequence: NC_007010.1) against the assembly using BLASTN. The contig was rearranged, using SeqKit (Shen et al. 2016), to start at the same position (the first base of the tRNA-Ile gene) as the reference. The contig was then polished by Pilon v1.23 (Walker et al. 2014) fed with a Minimap2 alignment of error-corrected paired-end reads from the muscle library. We manually annotated the mitogenome based on the reference M. japonicus mitogenome (NCBI Reference Sequence: NC_007010.1).

Repeat identification
To characterize repeat elements in the M. japonicus genome, we generated a custom repeat library using RepeatScout v1.0.5 (Benson 1999), its associated scripts (build_lmer_table, filter-stage-1.prl, and filter-stage-2.prl), and RepeatMasker v4.1.1 (http://www.repeatmasker.org). The Mj_TUMSAT_v1.0 assembly was broken into contigs using split.scaffolds.to.contigs.pl script, and 18,434 contigs (998,779,913 bp) were subsampled using SeqKit. 14-mers occurring in the subsampled contigs were tallied using build_lmer_table script. The resulting 14-mer table was used as input for RepeatScout v1.0.5, and the output was parsed with filter-stage-1.prl script to retain candidate repeat elements. The candidate repeat elements were queried against the subsampled contigs with RepeatMasker v4.1.1, using RMBLAST v2.1.0 as the search engine. Entries occurring no more than 3 times were discarded using filter-stage-2.prl script. The final repeat library was annotated using RepeatClassifier v2.0, a supplementary tool of RepeatModeler2 (Flynn et al. 2020) bundled in the Dfam TE Tools Container v1.2 (https://github.com/Dfam-consortium/TETools). The annotated repeat library was used to characterize the repeat contents of the finished assembly using RepeatMasker with RMBLAST as the search engine.

Prediction and functional annotation of protein-coding genes
To predict the protein-coding genes from the M. japonicus genome assembly, we combined cDNA alignments, RNA-seq read alignments, protein alignments, and ab initio predictions.

We newly generated a transcriptome shotgun assembly to maximize cDNA alignment-based genome annotation. The raw RNA-seq reads were trimmed using Fastp v0.20.1, and the trimmed reads were de novo assembled using Trinity v2.11.0 (Grabherr et al. 2011). The assembly (Trinity transcripts) consists of 1,810,475 contigs (N50: 762, Max: 38,355, min: 163). The Trinity transcripts and Iso-Seq cDNA sequences were used as input of DNA alignment-based gene prediction by PASA v2.4.1 (Haas et al. 2003), using BLAT v36 as aligner and TransDecoder v5.5.0 for CDS prediction.

To generate a genome-guided transcriptome assembly, we mapped the trimmed RNA-seq reads using HISAT2 v2.2.1 and reconstructed the transcripts using StringTie v2.1.4 (Pertea et al. 2015).

The RNA-seq alignments (generated by HISAT2) were used as the hints for ab initio gene prediction on the soft-masked genome by BRAKER2 v2.1.5 (Fetina et al. 2021), which internally runs GeneMark-ES v4.62.lic (Borodovsky and Lomsadze 2011), and Augustus v3.3.3 (Stanke et al. 2006).

To prepare protein-to-genome alignments, we downloaded 694 entries of NCBI Identical Protein Groups associated with M. japonicus (NCBI: txid27405; downloaded December 17, 2020). The proteins were queried to the genome, using TBLASTN, to identify

Figure 1 Characterization of the M. japonicus genome. (A) GenomeScope2.0 analysis using a 23-mer table built from trimmed Illumina paired-end reads (384 Gb). The M. japonicus genome size was estimated to be 1.93 Gb, with repetitive fraction occupying 1.40 Gb (72.5%) and a heterozygosity of 0.72%.

(B) SmudgePlot ploidy analysis using the same 23-mer table with (A). No evidence of whole genome duplication was observed.
Table 1 Assembly statistics of *M. japonicus* and other penaeid shrimp genomes

| Species          | M. japonicus | P. monodon | L. vannamei | F. chinensis |
|------------------|--------------|------------|-------------|--------------|
| Estimated genome size (Gb) | 1.93−2.25b   | 2.09−2.45c | 1.74d−1.88d | 1.74−1.88d   |
| Assembly         | Mj_TUMSAT_v1.0 | NSTDAPmon_1 | ASM378908v1 | ASPM19025.1  |
| Accession No.    | GCA_017512705.1 | GCA_015228065.1 | R63913_1 | GCA_016920825.1 |
| Reference        | This study  | Uengwetwanit et al. (2020) | Van Quyen et al. (2019) | Yuan et al. (2021) |
| Used platforms   | Illumina, ONT | Illumina, PacBio, Illumina, Hi-C | Illumina, ONT | PacBio, Illumina, Hi-C |
| Contig statistics |              |            |             |              |
| Contig number    | 31,741       | 2,937,051  | 1,185,601   | 50,608       |
| Total length     | 1,669,100,725| 1,624,778,290| 2,000,783,471| 1,618,035,433|
| Contig N50       | 113,129      | 682        | 45,084      | 1781         |
| Largest contig   | 862,579      | 16,221     | 1,387,722   | 67,204       |
| Scaffold statistics |            |            |             |              |
| Scaffold number  | 18,210       | 2,434,740  | 1,660,270,162| 1,583,041,647|
| Total length     | 1,704,394,957| 1,660,747,767| 1,632,388,725| 1,581,041,647|
| Scaffold N50     | 234,949      | 912        | 44,862,054  | 605,555      |
| Largest scaffold | 1,343,847    | 29,048     | 65,869,259  | 3,458,385    |
| Gap content (%)  | 2.11         | 2.14       | 1.73        | 2.74         |
| GC (%)           | 34.48        | 39.26      | 38.93       | 36.69        |
| Assembly size/Estimated genome size (%) | 76−88 | 74−87 | 84−86 | 57−58 |
| BUSCO genome completeness | 946 (93.4%) | 143 (14.2%) | 848 (83.7%) | 890 (87.8%) |
| Complete and single-copy | 918 (90.6%) | 113 (11.2%) | 830 (81.9%) | 835 (82.4%) |
| Complete and duplicated | 28 (2.8%) | 30 (3.0%) | 18 (1.8%) | 55 (5.4%) |
| Fragmented       | 29 (2.9%)    | 355 (35.0%)| 48 (4.7%)  | 46 (4.5%)    |
| Missing          | 38 (3.7%)    | 515 (50.8%)| 117 (11.6%)| 68 (6.7%)    |

a Estimated genome size obtained from this study.
b Estimated genome size calculated by converting the values from Swathi et al. (2018) with the following formula: 0.978 Gb/pg DNA (Dolezel et al. 2003).
c Estimated genome size calculated by Swathi et al. (2018) with the following formula: 0.978 Gb/pg DNA (Dolezel et al. 2003).
d Estimated genome size by Yuan et al. (2021).
We then ran Exonerate (Slater and Birney 2005) for each protein-scaffold pair and merged the results into a single file. Transcript evidences (PASA assemblies and StringTie predicted transcripts), protein evidences (TransDecoder CDS predicted on PASA assemblies and Exonerate protein alignments), and ab initio predictions (generated by BRAKER2) were merged using EVidenceModeler v1.1.1 (Haas et al. 2008) to derive a consensus gene model set.

We sought to filter out low-quality predicted gene models with no detectably similarity to known protein sequences. To this end, we first picked up the longest isoform from each locus using gffread v0.12.3 (Pertea and Pertea 2020), GenomeTools v1.6.1 (Gremme et al. 2013), and custom scripts. The nonredundant proteins were queried, using MMSeqs2 v12.113e3, against the UniRef50 protein database (49,410,134 entries; downloaded December 20, 2020) and the accessioned protein products annotated on the RefSeq genome assemblies of *P. monodon* (GCF_015228065.1; 32,900 entries; downloaded December 1, 2020) and *L. vannamei* (GCF_003789085.1; 33,273 entries; downloaded December 1, 2020). Gene models no significant hits (e-value cutoff: 1e−10) were regarded as low-quality and discarded. We noticed that the gene coding for penaeidin-II (an antimicrobial peptide; GenBank Accession no. AMH87234) was missing from the gene model set, and the gene was manually added based on an Exonerate EST-to-genome alignment.

Transcript evidences (PASA assemblies and StringTie predicted transcripts), protein evidences (TransDecoder CDS predicted on PASA assemblies and Exonerate protein alignments), and ab initio predictions (generated by BRAKER2) were merged using EVidenceModeler v1.1.1 (Haas et al. 2008) to derive a consensus gene model set.

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EVidenceModeler integrates predicted gene models from multiple sources to produce high-quality CDS predictions, but the
resulting gene models do not contain UTR information. To rescue the UTR features, the gene models were subjected to two rounds of PASA updates with the input of the Trinity transcripts, transcriptome shotgun assembly, and Iso-Seq cDNA sequences. We sought to maximize the recovery of genes that could be predicted by transcript evidence. The shotgun transcriptome assembly and Iso-Seq cDNA sequences were aligned to the genome by Minimap2, and isoforms were collapsed by cDNA_Cupcake v24.3.0 (https://github.com/Magdoll/cDNA_Cupcake). The collapsed transcripts were structurally annotated into gene models by SQANTI3 v3.3 (https://github.com/ConesaLab/SQANTI3) (Tardaguila et al. 2018), and the gene models from all the sources were merged by TAMA (https://github.com/GenomeRIK/tama) (Kuo et al. 2020). ORFs were predicted by TransDecoder v5.5.0; here, we included homology search (queried using MMSeqs2 against UniRef50, GCF_015228065.1_NSTDA_Pmon_1_protein.faa, and GCF_003789085.1_ASM378908v1_protein.faa; e-value cutoff: 1e−10) as ORF retention criteria. The TransDecoder-predicted proteins were again queried using MMSeqs2 against UniRef50, GCF_015228065.1_NSTDA_Pmon_1_protein.faa, and GCF_003789085.1_ASM378908v1_protein.faa, and proteins with no significant hits (e-value cutoff: 1e−03) were discarded. We additionally discarded a total of six protein-coding genes on scaffold_01936 (GenBank Accession no. BOPN01001936), because they overlapped with a nuclear ribosomal RNA gene cluster coding for 18S, 5.8S, and 28S rRNAs (predicted by cmscan; see “Noncoding RNA annotation”).

To assign functional annotation to the _M. japonicus_ protein-coding genes, we used the Automated Assignment of Human Readable Descriptions (AHRD) pipeline v3.3.3 (https://github.com/groupschoof/AHRD) and KEGG Automatic Annotation Server (KAAS) v2.1 (accessed June 2021; https://www.genome.jp/kaas-bin/kaas_main) (Moriya et al. 2007).

| Table 2 | Transcriptome assembly statistics of _M. japonicus_, _P. monodon_, and _L. vannamei_ |
| Species | M. japonicas | P. monodon | L. vannamei |
| Accession no. | ICRK00000000.1 | ICRJ00000000.1 | GGLH00000000.1 | GGUK00000000.1 |
| Reference | This study | Huerlimann et al. (2018) | Illumina | Zeng et al. (2018) |
| Used platforms | Illumina | PacBio | Illumina | PacBio |
| Transcript number | 40,991 | 25,415 | 236,085 | 47,260 |
| Total length (bp) | 91,091,355 | 64,458,168 | 225,712,489 | 162,735,114 |
| N50 (bp) | 38,693 | 2926 | 1431 | 3978 |
| Maximum length (bp) | 45,83 | 42.18 | 44.25 | 44.76 |
| GC (%) | Complete | 942 (93.0%) | 684 (67.5%) | 965 (95.3%) |
| BUSCO transcriptome completeness | Complete and single-copy | 893 (88.2%) | 364 (35.9%) | 487 (48.1%) |
| (BUSCO v4.1.4, arthropoda_odb10) | Complete and duplicated | 49 (4.8%) | 320 (31.6%) | 478 (47.2%) |
| Fragmented | 8 (0.8%) | 21 (2.1%) | 9 (0.9%) | 31 (3.1%) |
| Missing | 63 (6.2%) | 308 (30.4%) | 39 (3.8%) | 206 (20.2%) |

| Table 3 | The _M. japonicus_ genome annotation statistics |
| Number of predicted protein-coding genes | 26,381 |
| Maximum coding exon count | 170 |
| Median coding exon count | 5 |
| Average coding exon count | 6.51 |
| Median coding exon length (bp) | 151.0 |
| Average coding exon length (bp) | 230.7 |
| Median coding intron length (bp) | 555 |
| Average coding intron length (bp) | 2539 |
| BUSCO completeness of the predicted protein-coding genes | Complete |
| (BUSCO v4.1.4, arthropoda_odb10) | Complete and single-copy |
| Complete and duplicated | 931 (91.9%) |
| Fragmented | 28 (2.8%) |
| Missing | 19 (1.9%) |
| rRNA | 35 (3.4%) |
| tRNA | 81 |
| snRNA | 2314 |
| scRNA | 133 |
| snoRNA | 83 |
| Nuclear RNase P | 69 |
| Guide RNA | 6 |
| Number of predicted noncoding RNA genes | 1 |
| rRNA | 81 |
| tRNA | 2314 |
| snRNA | 133 |
| scRNA | 83 |
| snoRNA | 69 |
| Nuclear RNase P | 6 |
| Guide RNA | 1 |
Noncoding RNA gene prediction
tRNA genes were predicted using tRNAscan-SE 2.0 (Chan et al. 2021). Predicted tRNA genes labeled as “pseudo” or overlapping with protein-coding regions were discarded. Other noncoding RNAs were predicted by Infernal cmscan v1.1.4 (Nawrocki and Eddy 2013), with the Rfam 14.5 database as reference. We used infernal-tblout2gff.pl script (https://github.com/nawrockie/jiffy-infernal-hmmer-scripts/blob/master/infernal-tblout2gff.pl) to convert cmscan output to GFF format. We first discarded spurious or ambiguous hits (e.g., tRNAs, bacterial RNA families, histone H3 UTR, microRNAs, and so on.) and extracted the genomic coordinates matching the retained RNA families. The sequences from each RNA family were clustered by cd-hit-est v4.8.1 (-c 0.98 -s 0.95), and the nonredundant sequences were aligned by MAFFT v7.481 (Katoh et al. 2019) (accessed June 2021; https://mafft.cbrc.jp/alignment/server/) to convert cmscan output to GFF format. The resulting alignments were manually inspected to remove sequences of aberrant length. The remaining predicted RNA genes were projected to the final annotation file.

Construction of GFF3 annotation file
The predicted protein-coding, tRNA, and other noncoding RNA genes were integrated into a GFF3-format annotation file using custom scripts. Functional annotations of the protein-coding genes were incorporated as attributes (column 9) of the corresponding CDS features. We used GenomeTools v1.6.1 (Gremme et al. 2013) for sorting gff3 features.

Structural annotation of Iso-Seq cDNA sequences
The Iso-Seq cDNA sequences were aligned to the genome by Minimap2 and collapsed into nonredundant isoforms by cDNA_Cupcake v2.4.0. The nonredundant isoforms were structurally annotated by SQANTIS v3.3 with the final M. japonicus genome annotation as reference.

Comparison with other arthropod genome assemblies
We downloaded a total of 18 arthropod genome assemblies from the NCBI Genome database (The International Silkworm Genome Consortium 2008; Eyun et al. 2017; Gutekunst et al. 2018; Pouyton et al. 2018; Yuan et al. 2018, 2021; Becking et al. 2019; Chebbi et al. 2019; Lee et al. 2019; Wallberg et al. 2019; Zhang et al. 2019; Kim et al. 2019a; Tan et al. 2020; Van Quyen et al. 2020; Larkin et al. 2021). The BUSCO scores were calculated by BUSCO v4.1.4 (Simão et al. 2015) using the arthropoda_odh10 dataset. A BUSCO analysis plot was drawn using an R script generated by generate_plot.py (https://gitlab.com/ezlab/busco/-/blob/master/scripts/generate_plot.py) (Simão et al. 2015).

Simple repeat contents of 16 representative arthropod genomes were estimated using RepeatMasker v4.1.1, with the following options: -nopt -no_is -norma -gff -xsmall -a.

Comparison of protein-coding gene metrics in three penaeid shrimp genomes
We compared the protein-coding gene metrics (exon counts, exon lengths, intron lengths, and gene lengths, all excluding UTRs) of M. japonicus, L. vannamei, and P. monodon. We downloaded the NCBI RefSeq genome annotations for P. monodon (GenBank GCF_015228065.1_NSTDA_Pmon_1_genomic.gff; downloaded December 1, 2020) and L. vannamei (GenBank GCF_003789085.1_ASM378908v1_genomic.gff; downloaded December 1, 2020). We picked up the longest transcript from each locus using gffread v0.12.3, GenomeTools v1.6.1, and custom scripts. The results were visualized using ggplot2 (Wickham 2016).

Orthologous gene clustering analysis
The nonredundant proteomes of three penaeid shrimps (generated in “Comparison of protein-coding gene metrics in three penaeid shrimp genomes”) were used for ortholog analysis using OrthoFinder2 v2.5.1 (Emms and Kelly 2019). We used VennDiagram (Chen and Boutros 2011) for visualization.

Results and discussion
Marsupenaeus japonicus genome sequencing, characterization, and assembly
We sequenced the M. japonicus genome using Illumina HiSeq4000 and ONT GridION platforms, generating a total of 723 Gb (564 Gb Illumina paired-end reads, 144 Gb Illumina mate-pair reads, and 14.6 Gb ONT long reads; Supplementary Table S2). Collectively, these sequencing data cover 375 times the estimated haploid genome size (1.93 Gb) of M. japonicus.

GenomeScope 2.0 analysis using Illumina paired-end reads (k = 23; model fit = 92.7%; Supplementary Table S4) yielded an estimated genome size of 1.93 Gb (Figure 1A), which is slightly smaller than the previously reported value [2.30 pg by Swathi et al. (2018), which equals 2.25 Gb assuming 0.978 Gb/pg (Doležel et al. 2003)]. The estimated repeat content of 72.5% and heterozygosity of 0.72% indicates that M. japonicus genome is highly repetitive and heterozygous (Figure 1A). SmudgePlot analysis detected no evidence of polyploidization (Figure 1B).

Error correction of Illumina paired-end reads by Tadpole reduced low-coverage k-mers and increased the coverage of homo- and heterozygous k-mer peaks, suggesting that error correction effectively purged erroneous k-mers from the data set (Supplementary Figure S3). However, this also introduced a drop of the heterozygous peak relative to the homozygous peak, which would lead to an underestimation of heterozygosity rate. We thus used the uncorrected reads for genome size estimation and ploidy analysis and used the error-correcting reads for assembly polishing.

To obtain the M. japonicus draft genome assembly, we first assembled filtered Illumina paired-end reads from the sperm library (93.6 Gb) and scaffolded the contigs using illumina mate-pair reads and transcriptome data. The resulting assembly (“Illumina primary assembly” in Supplementary Table S5) consisted of 22,425 scaffolds, with the N50 of 209,993 bp and gap rate of 34.0%. The Illumina primary assembly was further improved using ONT long reads, illumina mate-pair reads, and transcriptome data, followed by manual curation.
The final assembly (Mj_TUMSAT_v1.0) contains 18,210 scaffolds with a total length of 1,704,994,957 bp and 34.48% GC content (Table 1). Assuming a genome size of 1.93 Gb, Mj_TUMSAT_v1.0 covers 88% of the M. japonicus genome; however, 95.28% of muscle and 95.60% of sperm Illumina paired-end reads were successfully mapped back to the assembly. The BUSCO completeness (arthropoda_odb10 dataset; n = 1066) was 93.4%, indicating that the completeness of the genome assembly is high. Mj_TUMSAT_v1.0 significantly improved contiguity and BUSCO score over the previously published M. japonicus genome assembly [Mjap_WGS_v1.0 (Yuan et al. 2018) in Table 1]. Although Mj_TUMSAT_v1.0 had a lower scaffold N50 than PacBio long read-based genome assemblies of other species, the contig N50 was comparable, and the BUSCO score was higher (Table 1). The BUSCO score of Mj_TUMSAT_v1.0 was comparable to other representative arthropod genome assemblies, both in terms of completeness and low redundancy (Figure 2).

We also assembled a complete M. japonicus mitogenome (GenBank Accession No. LC635856, Figure 3) from ONT long reads. The M. japonicus Ginoza2017 mitogenome is 15,969 bp and encodes 13 protein-coding genes, 22 tRNA genes, and two rRNA genes. Pairwise BLASTN alignment showed that the M. japonicus Ginoza2017 mitogenome is 99.51% identical to the reference M. japonicus genome (NCBI Reference Sequence: NC_007010.1).

In summary, Mj_TUMSAT_v1.0 significantly improves the genomic resources of M. japonicus with a draft genome assembly with decent contiguity and high completeness.

Repeat identification

To characterize the repeat landscape of the M. japonicus genome, we constructed a M. japonicus-specific repeat library (4335 entries; total: 1,865,850 bp: maximum: 11,554 bp; average: 430.4 bp; minimum: 51 bp; see Supplementary Table S6 for a summary of annotated repeat elements; repeat library available as Supplementary File Mj_TUMSAT_v1.0.repeats.fna). RepeatMasker analysis using the M. japonicus-specific repeat library masked 54% of Mj_TUMSAT_v1.0, indicating that the kuruma shrimp is repeat-rich (Supplementary Table S7). A majority of the interspersed elements were unclassified (occupying 14.81% of the genome),

![Figure 4](image-url) Protein-coding gene metrics of three penaeid shrimp genomes. The distributions of (A) gene lengths, (B) exon counts, (C) exon lengths, and (D) intron lengths were calculated for the protein-coding genes (excluding UTRs) of M. japonicus (26,381 genes), P. monodon (24,079 genes), and L. vannamei (24,974 genes).
followed by retroelements (7.45%) and DNA transposons (0.06%). Simple repeats covered 27.44% of the assembly. We found similar levels of simple repeats in L. vannamei (34.34%), P. monodon (21.64%), and F. chinesis (31.00%), consistent with previous reports (Zhang et al. 2019; Uengwetwanit et al. 2021; Yuan et al. 2021) (Supplementary Table S8). Among the other arthropods analyzed, only pill bugs (Armadillidium vulgare, 26.37%, A. nasatum, 20.29%) harbored similar proportions of simple repeats (Supplementary Table S8). Overall, the M. japonicus genome, similar to other penaeid shrimps, is repeat-rich and is characterized by an unusually high content of simple repeats.

**Marsupenaeus japonicus transcriptome sequencing and assembly**

To obtain a comprehensive transcript dataset and aid genome scaffolding, we sequenced 49 RNA samples using Illumina HiSeq4000 platform and obtained a total of 346 Gb sequencing reads (Supplementary Table S2). To eliminate possible contamination, we de novo assembled the reads that successfully mapped back to the M. japonicus genome (average mapping rate: 87.25%; max: 94.26%; min: 79.22%). The transcriptome shotgun assembly consisted of 40,991 transcripts with a BUSCO completeness of 93.0% (Table 2). This BUSCO completeness was close to or above those of other penaeid shrimps (Huerlimann 2019; Yuan 2018) (Table 2).

We also sequenced four samples using the PacBio Iso-Seq platform, generating 25,415 transcripts with a BUSCO completeness of 67.5% (Table 2). We mapped the Iso-Seq cDNA sequences to the genome, collapsed redundant isoforms, and structurally annotated the mapped transcripts using SQANTI3 pipeline. A total of 7486 unique genes (6487 annotated, 999 novel) and 16,160 isoforms were mapped to the genome. The full SQANTI3 report and nonredundant, genome-corrected Iso-Seq cDNA sequences are provided as Supplementary Files (IsoSeq_sqanti_report.pdf, IsoSeq_aligned.nr.corrected.fna).

**Genome annotation and ortholog inference**

A total of 26,381 protein-coding, 2314 tRNA, 81 rRNA, and 292 other noncoding RNA genes were predicted on the M. japonicus genome (Table 3; Supplementary File Mj_TUMSAT_v1.0.gff3). An average M. japonicus protein-coding gene (excluding untranslated regions) spanned 15,490 bp containing 6.51 exons, with an average exon length of 230.7 bp and average intron length of 2539 bp (Table 3, Figure 4). The distributions of gene lengths, exon counts, exon lengths, and intron lengths were similar to those of L. vannamei and P. monodon (Figure 4). The translated M. japonicus predicted gene models yielded 94.7% BUSCO completeness (arthropoda_odb10 dataset; n = 1066) (Table 3), closely matching the completeness of the genome mode. This indicates that the predicted gene models capture most of the coding capacity of the M. japonicus genome. The number of predicted M. japonicus protein-coding genes was slightly larger than those in the NCBI RefSeq annotations for other penaeid shrimp genomes (L. vannamei: 24,974 genes; P. monodon: 24,079 genes).

We compared the proteomes of M. japonicus, L. vannamei, and P. monodon using Orthofinder2 (Figure 5). A total of 17,239 orthologs were identified, of which 15,370 orthogroups (89.2%) contained M. japonicus genes. 20,683 M. japonicus genes (78.4%) were clustered in orthogroups shared with at least one other shrimp species. Eight thousand and twenty-seven genes were conserved as single-copy genes across the three species. These observations indicate that most M. japonicus predicted genes have putative orthologs in other penaeid shrimp genomes and further substantiates the quality of the predicted M. japonicus gene models.

**Conclusions**

Here, we present a high-quality draft genome assembly and two transcriptome assemblies of the kuruma shrimp M. japonicus, an economically important crustacean. The 1.70 Gb draft genome assembly covers 88% of the estimated genome size (1.93 Gb) and is characterized by a high BUSCO score (93.4% complete). We predicted a total of 26,381 protein-coding gene models with a BUSCO completeness of 94.7%. The two transcriptome datasets complement the genome assembly by capturing complex transcript structures. We expect that our datasets will serve as a valuable resource for basic research, fisheries management, and breeding programs for M. japonicus as well as comparative genomics of penaeid shrimps and other crustaceans.

**Data availability**

The assembled sequences presented in this study are available in DDBJ/ENA/NCBI under the following accession numbers: genome assembly: GCA_017312705.1; mitogenome: LC635856.1; transcriptome shotgun assembly: ICRK00000000.1; PacBio Iso-Seq assembly: ICRJ00000000.1. The raw reads are available in DDBJ/ENA/NCBI under the accession numbers DRA011460, DRA011716, and DRA011525.

The supplementary figures, supplementary tables, and supplementary files (custom scripts: Mj_TUMSAT_v1.0.html and Mj_TUMSAT_v1.0.md; GFF3 annotation file: Mj_TUMSAT_v1.0.gff3; predicted transcripts: Mj_TUMSAT_v1.0.transcripts.fna; predicted CDS: Mj_TUMSAT_v1.0.cds.fna; translated CDS: Mj_TUMSAT_v1.0.translated.fna) are available on the Dryad Digital Repository (https://doi.org/10.5061/dryad.1234567890).
SQANTI3 report for the Iso-Seq cDNA sequences: IsoSeq_sqanti_report.pdf, genome-corrected, nonredundant Iso-Seq cDNAs: IsoSeq aligned.nr.corrected.fna) are available on figshare: https://doi.org/10.25387/g3.15043521.

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Conflicts of interest
The authors declare that there is no conflict of interest.

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