Data Descriptor: Embryonic transcriptome sequencing of the ocellate spot skate Okamejei kenojei

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Chondrichthysans (cartilaginous fishes) exhibit highly variable reproductive styles, categorized as viviparity and oviparity. Among these, species with oviparity provide an enormous potential of molecular experimentation with stable sample supply which does not demand the sacrifices of live mothers. Cartilaginous fishes are divided into two subclasses, chimaeras (Holocephali) and elasmobranchs (Elasmobranchii), and the latter consists of two monophyletic groups, Batoidea (rays, skates and torpedoes) and Selachimorpha (sharks). Here we report transcriptome assemblies of the ocellate spot skate Okamejei kenojei, produced by strand-specific RNA-seq of its embryonic tissues. We obtained a total of 325 million illumina short reads from libraries prepared using four different tissue domains and assembled them all together. Our assembly result confirmed the species authenticity and high continuity of contig sequences. Also, assessment of its coverage of pre-selected one-to-one orthologs supported high diversity of transcripts in the assemblies. Our products are expected to provide a basis of comparative molecular studies encompassing other chondrichthyan species with emerging genomic and transcriptomic sequence information.

| Design Type(s)                  | organism development design | transcription profiling by high throughput sequencing design |
|--------------------------------|-----------------------------|----------------------------------------------------------|
| Measurement Type(s)            | transcription profiling assay| Transcriptome assembly                                    |
| Technology Type(s)             | RNA sequencing              |                                                          |
| Factor Type(s)                 | animal body part            |                                                          |
| Sample Characteristic(s)       | Okamejei kenojei            | embryo                                                  |

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Background & Summary

Oviparous (egg-laying) chondrichthyans are distributed into three different selachimorphan (shark) orders, Carcharhiniformes, Heterodontiformes, and Orectolobiformes, and one batoïd (skate) order Rajiformes (summarized in1). The order Rajiformes consists of four families, Anacanthobatidae, Arhynchobatidae, Gergusiellidae, and Rajidae. As of February 2018, molecular sequence data for rajiformes concentrate on species in Rajidae, such as the little skate Leucoraja erinacea, whose genome is being sequenced2, in the head of the list, and it is followed by Amblyraja radiata and Raja clavata. One of the genera in Rajidae is Okamejei, and among more than a dozen of species in the genus Okamejei, we focused on the ocellate spot skate Okamejei kenojei in the present study. Previously, this species was recognized as Raja porosa, Okamejei porosa, or Raja kenojei. The NCBI RefSeq entry for the whole mitochondrial genome sequence of this species Okamejei kenojei (NC_007173.1) was originally registered in GenBank under one of the former species names, Raja porosa (AY525783.1)3. The habitat of O. kenojei is the coasts of Japan, Korea, China, and Taiwan in the Northwest Pacific4. While the skate species with the most abundant sequence information, L. erinacea, inhabits a small area on the east coast of North America, O. kenojei is one of the most promising oviparous skate species available in East Asia for experimentation.

For modern life science studies, genomic and transcriptomic information serves as an indispensable fundamental resource. Especially for a species without abundant molecular sequences, even small-scale transcriptome sequencing with short-read data acquisition can provide a valuable start point, which can be achieved by simple tissue sampling and small financial investment. For efficient transcriptome sequencing data acquisition, a number of technical factors have been considered to optimize sample preparation, sequencing run design, and post-sequencing data analysis. Previously, the authors’ group proposed some of those factors5, and in the present study, we extend those factors by incorporating latest reagents for sample preparation and tools for sequence data analysis.

Methods

Animal sampling and RNA extraction

All animal experiments were conducted in accordance with the Guideline of the Institutional Animal Care and Use Committee (IACUC) of RIKEN Kobe Branch (Approval ID: H16-11). An approximately 8 cm-long fertilized egg of the ocellate spot skate, O. kenojei (Fig. 1a), was purchased from a commercial marine organism supplier in Minami-ise town in Mie Prefecture, Japan, in February 2016. The embryo contained in the eggcase was 5 cm-long, corresponding morphologically to the stage 31 of typical shark embryos5 and was dissected into six pieces labelled as head, gill, trunk, pectoral fins, cloaca, and caudal (Fig. 1b). Total RNAs were extracted from the four parts of the embryo (head, trunk, cloaca, and caudal) using TRIzol reagent (Life Technologies) (Fig. 1c).

Library preparation and sequencing

Using 1 μg of each of the extracted total RNAs, four strand-specific RNA-seq libraries were prepared with KAPA Stranded mRNA-Seq Kit (Kapa Biosystems, cat. No. KK8420) according to its standard protocol unless stated otherwise below. Before the total volume PCR amplification was performed, we performed a preliminary PCR using a 1.5 μl aliquot of 10 μl DNA from the previous step, with KAPA Real-Time Library Amplification Kit (Kapa Biosystems, cat. No. KK2702). This demonstrated that the amplification of the products reached Standard 1 accompanying this kit between three and four PCR cycles, which instructed us to perform the full-volume PCR with three PCR cycles, introducing the minimal amplification (Fig. 1d). The libraries were first sequenced in-house after on-board cluster generation for 127 cycles using 3x HiSeq Rapid SBS Kit v2-HS (50 cycles) (Illumina, cat. No. FC-402-4022) and HiSeq PE Rapid Cluster Kit v2-HS (Illumina, cat. No. PE-402-4002) on a HiSeq 1500 (Illumina) operated by HiSeq Control Software v2.2.58. The output was processed with Illumina RTA v1.18.64 for basecalling and with bcl2fastq v1.8.4 for de-multiplexing. To obtain more reads, we outsourced the sequencing for three of the four above-mentioned libraries on HiSeq 4000. Quality control of the obtained fastq files for individual libraries was performed with FASTQC v0.11.5 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/).

Read trimming and assembly

The obtained sequence reads in the fastq files (Data Citation 1) were processed with the program Trim Galore! v0.3.3 (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) with the options ‘--phred33 --stringency 2 --quality 30 --length 25 --paired’. The reads after adaptor trimming were assembled with the program Trinity v2.5.18 with the options ‘--trimmomatic --min_kmer_cov 2 --SS_lib_type RF’. Among the resultant contig sequences, those matching PhiX, mitochondrial DNA, and genome and transcriptome assemblies of the species sequenced in the same HiSeq run in the BLASTN v2.2.30*b results executed with the options ‘perc_identity 95’ were removed. We designated the resultant sequence set as the redundant nuclear transcriptome assembly (Data Citation 2). In this assembly, protein-coding regions were predicted with the program Transdecoder v5.0.210 following its user documentation, which employed the protein-level similarity to SWISSPROT11 and PFAM12 by the programs BLASTP v2.2.30+ and Hmmer v3.1b213, respectively (Data Citation 3). The obtained amino acid sequences were processed by CD-HIT v4.714 with its default parameters to reduce redundancy in it.
Annotation of the putative peptide sequences were performed with similarity searches using BLASTP v2.2.30+ towards RefSeq Protein sequences for the human (GRCh38, 113,620 sequences - as of July 12, 2018) and the Callorhinchus milii (NCBI RefSeq, 28,600 sequences - as of July 12, 2018).

The whole post-sequencing procedure is outlined in Fig. 2.

**Code availability**

No custom computer code was employed in this study.

**Data Records**

The approximately 325 million raw sequence reads from four different portions of a *O. kenojei* embryo were released in the NCBI Sequence Read Archive (Table 1 and Data Citation 1). The nuclear transcriptome assembly using all the obtained reads consisted of 1,081,614 sequences (Data Citation 2), which could include spurious intergenic transcripts and alternative splicing variants. Mapping of the reads employed in the assembly to the assembled transcript contig sequences (Data Citation 2) yielded the mapping rates of as high as 90.2 to 93.1%. The nuclear protein-coding transcriptome assembly consisted of 167,783 sequences available both in the nucleotide and amino acid sequences (Data Citation 3). Out of those putative protein-coding sequences, 88,376 (52.7%) were predicted to have complete ORFs.
with start and stop codons. The amino acid sequence dataset after removing the redundancy consists of 79,083 components (Data Citation 4). Transcriptome assembly was also performed for the individual libraries for caudal, cloaca, trunk, and head samples, which resulted in 498,477, 377,609, 448,394, and 342,765 contigs, respectively (Data Citation 5–8). These data are available as multifasta files at FigShare.

We also provide a table containing best BLASTP hits to the human and C. milii RefSeq Peptide sequences as a reference for cross-species annotation (Data Citation 9).

### Technical Validation

#### Contiguity of RNA-seq reads

Next, we validated the contiguity of the transcript sequences by short read assembly. For this purpose, we focused on known transcript sequences of O. kenojei available at NCBI Nucleotide (https://www.ncbi.nlm.nih.gov/nuccore) that are longer than 1 Kbp and derived from the nuclear genome (with the filter ‘Okamejei kenojei[Organism] AND 1000:1000000[slen] NOT mitochondrial’, accessed on March 15, 2018). With this criterion, we have identified five sequence entries (AB371645.1, AB295474.1, etc.).

### Table 1. Sequencing statistics. All the total RNA samples exhibited the RIN of 10, and all the resultant libraries were amplified with as few as three PCR cycles.

| Library ID | Tissue      | Peak of library molecule length distribution (nt) | Sequencing configuration | # of raw read pairs obtained | # of qualified read pairs | SRA ID               |
|------------|-------------|-------------------------------------------------|--------------------------|-----------------------------|---------------------------|-----------------------|
| Ok1        | Caudal      | 272                                             | HiSeq 1500 paired-end 127 nt | 81,561,105                  | 80,108,371                | SRR6866827            |
| Ok2        | Cloaca      | 257                                             | HiSeq 1500 paired-end 101 nt | 29,204,559                  | 28,792,861                | SRR6866830            |
| Ok3        | Trunk       | 275                                             | HiSeq 1500, paired-end127 nt | 33,239,929                  | 32,302,602                | SRR6866826            |
| Ok4        | Head        | 269                                             | HiSeq 1500 paired-end 127 nt | 39,350,049                  | 38,390,345                | SRR6866824            |

### Table 2. Long contig sequence matches to existing database entries.

| NCBI ID       | Gene | Length (nt) | Transcript ID | Length (nt) | % Identity | % Gap |
|---------------|------|-------------|---------------|-------------|------------|-------|
| AB371645.1    | ptx-l| 1,416       | DN146223_c3_g1_i4 | 1,077       | 99         | 0     |
| AB295474.1    | ptx  | 1,540       | DN136073_c1_g3_i2 | 584         | 98         | 0     |
| AB195842.1    | SkCOL1A1 | 4,866     | DN146743_c0_g1_i7 | 3,558       | 99         | 0     |
| AB201248.1    | COL2A1 | 1,802      | DN128141_c7_g2_i3 | 4,913       | 99         | 0     |
| AB201247.1    | COL5/11A1 | 6,451     | DN133476_c9_g3_i7 | 2,630       | 98         | 1     |

### Table 3. Sequence datasets produced in this study. "See the existing literature" for the definitions of ‘complete’, ‘fragmented’ and ‘missing’ in ortholog detection by BUSCO. CVGs consists of 233 orthologs in total, while Vertebrata BUSCO has 2,586 orthologs.

| Dataset                              | Number of sequences | Number of core genes * | CVG (Vertebrata BUSCO) | N50 contig length (bp) | Data Records   |
|--------------------------------------|---------------------|------------------------|------------------------|------------------------|----------------|
| Transcriptome assembly - all libraries | 1,081,614           | 218 (2407)             | 232 (2524)             | 1 (62)                 | 992            | Data Citation 2 |
| Transcriptome assembly - caudal       | 498,477             | 226 (2423)             | 229 (2496)             | 4 (90)                 | 1,679          | Data Citation 5 |
| Transcriptome assembly - cloaca       | 377,609             | 224 (2371)             | 230 (2476)             | 3 (110)                | 1,744          | Data Citation 6 |
| Transcriptome assembly - trunk        | 448,394             | 225 (2389)             | 232 (2489)             | 1 (97)                 | 1,560          | Data Citation 7 |
| Transcriptome assembly - head         | 342,765             | 221 (2361)             | 228 (2454)             | 5 (132)                | 1,808          | Data Citation 8 |
| Protein-coding assembly - all libraries | 167,783             | 218 (2401)             | 232 (2514)             | 1 (72)                 | 1,782          | Data Citation 3 |
| Non-redundant peptides - all libraries | 79,083              | 219 (2400)             | 233 (2514)             | 0 (72)                 | N/A            | Data Citation 4 |

*See the existing literature for the definitions of ‘complete’, ‘fragmented’ and ‘missing’ in ortholog detection by BUSCO. CVGs consists of 233 orthologs in total, while Vertebrata BUSCO has 2,586 orthologs.
AB195842.1, AB201248.1, and AB201247.1), and for each of those sequences, a transcript contig in our assembly showed no less than 98% similarity (no more than 1% gaps) (Table 2). These high similarities authenticate that the species used for this study was *O. kenojei*.

### Transcript diversity measured by one-to-one ortholog coverage

We also employed a completeness assessment on the web server gVolante in which the ortholog search pipeline BUSCO is implemented. This method evaluates the coverage of one-to-one protein-coding orthologs selected in advance. For this purpose, we used the ortholog set CVG introduced for more accurate assessment for vertebrate sequence sets than performed with other ortholog sets, as well as the Vertebrata ortholog set released together with BUSCO. Although the assessment results showed a slight fluctuation depending on the type of input data (nucleotide or amino acid sequences), out of the 233 components of CVG, our resultant data were shown to always contain at least 218 full-length ortholog sequences (‘complete’) and 232 partial sequences (‘fragmented’) (Table 3). Our assessment with the BUSCO’s Vertebrata ortholog set also produced comparable scores, mostly >92% in percentages. These high scores ascertain the high coverage of protein-coding genes in our resultant transcript sequence data set.

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### Data Citations

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### Author Contributions

S.K. and M.K. conceived and supervised the project. F.M., K.T. and C.T. performed experiments. O.N. and S.K. processed the data. S.K. drafted the manuscript.

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