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## 2 septentrionalis) using Oxford Nanopore PromethION sequencing and Hi-C technology

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- 19 Abstract
- 20 The greenfin horse-faced filefish, *Thamnaconus septentrionalis*, is a valuable commercial
- 21 fish species that is widely distributed in the Indo-West Pacific Ocean. It has characteristic blue-

green fins, rough skin and spine-like first dorsal fin. T. septentrionalis is of a conservation 22 concern as a result of sharply population decline, and it is an important marine aquaculture fish 23 species in China. The genomic resources of this filefish are lacking and no reference genome 24 has been released. In this study, the first chromosome-level genome of T. septentrionalis was 25 constructed using Nanopore sequencing and Hi-C technology. A total of 50.95 Gb polished 26 Nanopore sequence were generated and were assembled to 474.31 Mb genome, accounting for 27 96.45% of the estimated genome size of this filefish. The assembled genome contained only 28 242 contigs, and the achieved contig N50 was 22.46 Mb, reaching a surprising high level among 29 all the sequenced fish species. Hi-C scaffolding of the genome resulted in 20 pseudo-30 chromosomes containing 99.44% of the total assembled sequences. The genome contained 31 67.35 Mb repeat sequences, accounting for 14.2% of the assembly. A total of 22,067 protein-32 coding genes were predicted, of which 94.82% were successfully annotated with putative 33 functions. Furthermore, a phylogenetic tree was constructed using 1,872 single-copy gene 34 families and 67 unique gene families were identified in the filefish genome. This high quality 35 assembled genome will be a valuable genomic resource for understanding the biological 36 characteristics and for facilitating breeding of *T. septentrionalis*. 37

38 Key words

39 Filefish, genome assembly, Oxford Nanopore sequencing, Hi-C

40 **1 Introduction** 

The greenfin horse-faced filefish (*Thamnaconus septentrionalis*; hereafter "filefish")
belongs to the family Monacanthidae (Tetraodontiformes) and has characteristic blue-green fins,

rough skin and spine-like first dorsal fin (Figure 1)(Su & Li, 2002). It is widely distributed in 43 the Indo-West Pacific Ocean, ranging from the Korean Peninsula, Japan and China Sea to East 44 Africa. Filefish is a temperate demersal species inhabiting a depth range of 50-120 m, and 45 feeding on planktons such as copepods, ostracods, and amphipods, as well as mollusks and 46 benthic organisms(Su & Li, 2002). It goes through annual long-distance seasonal migrations 47 and has diurnal vertical migration habits during wintering and spawning(Lin, Gan, Zheng, & 48 Guan, 1984; Su & Li, 2002). Due to a high protein content and good taste, filefish is an 49 important commercial species in China, Korea and Japan. An interesting feature of filefish is 50 its rough skin, whose roughness is actually attributed to the covered dense small scales. These 51 scales are difficult to remove, and people have to peel off the skin before eating. Given this, 52 53 filefish is also called "skinned fish" in China.

The wild resource of filefish has declined dramatically since 1990 due to overfishing, and 54 the annual catch in the East China Sea was only 3,842 tons in 1994(Chen, Li, & Hu, 2000). 55 Since then, researchers have attempted to explore the methods to properly culture filefish. 56 Several key technologies including fertilized eggs collection, sperm cryopreservation, larval 57 rearing, tank and cage culturing have been studied, and this species is cultivated commercially 58 in China, Korea and Japan(Guan et al., 2013; Kang et al., 2004; Li, Jiang, Xu, & Liu, 2002; Liu 59 et al., 2017; Mizuno, Shimizu-Yamaguchi, Miura, & Miura, 2012). The current main challenge 60 of filefish cultivation is the high mortality of fish fry during artificial breeding. A better 61 62 understanding of the underlying genomic-level characteristics will provide significant information to break through the bottleneck and benefit the cultivation industry of this filefish. 63

However, the available genetic information of filefish is scarce. At present, only limited genetic
studies regarding microsatellite loci isolation and population structure are available for this
filefish (An et al., 2011; An, Lee, Park, & Jung, 2013; Bian et al., 2018; Xu, Chen, & Tian,
2010; Xu, Tian, Liao, & Chen, 2009).

Spectacular improvements in high-throughput sequencing technology, especially the 68 single-molecule sequencing methods, have remarkably reduced the sequencing costs, making 69 a genome project affordable for individual labs. Oxford Nanopore sequencing technology is 70 currently the most powerful method for rapid generation of long-read sequences and has the 71 potential to offer relatively low-cost genome sequencing of non-model animals. It directly 72 detects the input DNA without PCR amplification or synthesis, so the length of sequenced DNA 73 can be very long. The longest read generated by Nanopore sequencing has been up to 2,272,580 74 bases(Payne, Holmes, Rakyan, & Loose, 2018). Nanopore sequencing has been used in several 75 76 fish species to construct high-quality genome assembly or to improve the completeness of previous genome drafts(Austin et al., 2017; Ge et al., 2019; Jansen et al., 2017; Kadobianskyi, 77 Schulze, Schuelke, & Judkewitz, 2019; Tan et al., 2018). In the case of red spotted grouper 78 (Epinephelus akaara), a chromosome-level reference genome with a contig N50 length of 5.25 79 Mb was constructed by taking advantage of Nanopore sequencing and Hi-C technology(Ge et 80 al., 2019). In clown anemonefsh (Amphiprion ocellaris), a hybrid Illumina/Nanopore method 81 generated much longer scaffolds than Illumina-only approach with an 18-fold increase in N50 82 83 length and increased the genome completeness by an additional 16%(Tan et al., 2018).

84 In this study, the first chromosome-level genome of filefish was constructed using

Nanopore sequencing and Hi-C technology. This genomic data will benefit a comprehensive conservation study of filefish along the China and Korea coast to implement better protection of wild populations, and allow us to screen for genetic variations correlated with fast-growth and disease-resistance traits of filefish in the future.

# 89 2 Materials and methods

# 90 2.1 Sample and DNA extraction

A single female fish (~325 g) was collected on August 2018 from the Tianyuan Fisheries Co., Ltd (Yantai, China).The muscle tissue below the dorsal fin was taken and stored in the liquid nitrogen until DNA extraction. Genomic DNA was extracted using CTAB (Cetyltrimethylammonium bromide) method. The quality and concentration of the extracted genomic DNA was checked using 1% agarose gel electrophoresis and a Qubit fluorimeter (Invitrogen, Carlsbad, CA, USA). This high-quality DNA was used for subsequent Nanopore and Illumina sequencing.

# 98 2.2 Library construction and genome sequencing

To generate Oxford Nanopore long reads, approximately 15 µg of genomic DNA was sizeselected (30–80 kb) with a BluePippin (Sage Science, Beverly, MA, USA), and processed according to the Ligation Sequencing Kit 1D (SQK-LSK109) protocol. Briefly, DNA fragments were repaired using the NEBNext FFPE Repair Mix (New England Biolabs). After endreparation and 3'-adenylation with the NEBNext End repair/dA-tailing Module reagents (New England Biolabs), the Oxford Nanopore sequencing adapters were ligated using NEBNext Quick Ligation Module (E6056) (New England Biolabs). The final library was sequenced on 3 different R9.4 flow cells using the PromethION DNA sequencer (Oxford Nanopore, Oxford,
UK) for 48 hours. The MinKNOW software (version 2.0) was used to conduct base calling of
raw signal data and convert the fast5 files into fastq files. These raw data was then filtered to
remove short reads (<5 kb) and the reads with low-quality bases and adapter sequences.</li>

110 Illumina sequencing libraries were prepared to carry out genome size estimation, 111 correction of genome assembly, and assembly evaluation. The paired-end (PE) libraries with 112 insert sizes of 300 bp were constructed according to the Illumina standard protocol (San Diego, 113 CA, USA) and subjected to PE ( $2 \times 150$  bp) sequencing on an Illumina HiSeq X Ten platform 114 (Illumina, San Diego, CA, USA). After discarding the reads with low-quality bases, adapter 115 sequences, and duplicated sequences, the clean reads were used for subsequent analysis.

## 116 **2.3 Genome size estimation and genome assembly**

117 A k-mer depth frequency distribution analysis of the Illumina data was conducted to 118 estimate the genome size, heterozygosity, and content of repetitive sequences of the filefish. 119 The k-mer analysis was carried out using "kmer freq stat" software (developed by Biomarker 120 Technologies Corporation, Beijing, China). Genome size (G) was estimated based on the 121 following formula: G = k-mer number/average k-mer depth, where k-mer number = total k-122 mers—abnormal k-mers (with too low or too high frequency).

For genome assembly, Canu (version 1.5) (Koren et al., 2017)was conducted for initial
read correction, and the assembly was performed by Wtdbg (https://github.com/ruanjue/wtdbg).
The consensus assembly was generated by 2 rounds of Racon (version 1.32)(Vaser, Sović,
Nagaranjan, & Šikić, 2017), and 3 rounds of Pilon (version 1.21)(Walker et al., 2014) polishing

127 using the Illumina reads with default settings.

## 128 **2.4 Hi-C library construction and sequencing**

For Hi-C sequencing, the muscle tissue of filefish was used for library preparation 129 according to Rao et al., 2014). Briefly, the tissue cells were fixed with formaldehyde 130 and restriction endonuclease Hind III was used to digest DNA. The 5' overhang of the fragments 131 were repaired and labeled using biotinylated nucleotides, followed by ligation in a small volume. 132 After reversal of crosslinks, ligated DNA was purified and sheared to a length of 300-700 bp. 133 The DNA fragments with interaction relationship were captured with streptavidin beads and 134 prepared for Illumina sequencing. The final Hi-C libraries were sequenced on an Illumina 135 HiSeq X Ten platform (Illumina, San Diego, CA, USA) to obtain  $2 \times 150$  bp paired-end reads. 136 137 To assess the quality of Hi-C data, the plot of insert fragments length frequency was first made to detect the quality of Illumina sequencing. Second, we used BWA-MEM (version 0.7.10-r789) 138 (Li & Durbin, 2009)to align the PE clean reads to the draft genome assembly. In the end, HiC-139 Pro (Servant et al., 2015) (version 2.10.0) was performed to find the valid reads from unique 140 mapped read pairs. 141

## 142 **2.5** Chromosomal-level genome assembly using Hi-C data

We first performed a preassembly for error correction of contigs by breaking the contigs into segments of 500 kb on average and mapping the Hi-C data to these segments using BWA-MEM (version 0.7.10-r789)(Li & Durbin, 2009). The corrected contigs and valid reads of Hi-C were used to perform chromosomal-level genome assembly using LACHESIS(Burton et al., with the following parameters: CLUSTER MIN RE SITES=22; 148 CLUSTER\_MAX\_LINK\_DENSITY=2; CLUSTER\_NONINFORMATIVE\_RATIO=2;
149 ORDER\_MIN\_N\_RES\_IN\_TRUNK=10; ORDER\_MIN\_N\_RES\_IN\_SHREDS=10. To
150 evaluate the quality of the chromosomal-level genome assembly, a genome-wide Hi-C heatmap
151 was generated by ggplot2 in R package.

# 152 **2.6 Assessment of the genome assemblies**

To assess the genome assembly completeness and accuracy, we first aligned the Illumina reads to the filefish assembly using BWA-MEM (version 0.7.10-r789)(Li & Durbin, 2009). Furthermore, CEGMA (version 2.5) (Parra, Bradnam, & Korf, 2007)was conducted to find core eukaryotic genes (CEGs) in the genome with parameter set as identity>70%. Finally, the completeness of the genome assembly was also evaluated by using BUSCO (version 2.0)(Simao, Waterhouse, Ioannidis, Kriventseva, & Zdobnov, 2015) search the genome against the actinopterygii database, which consisted of 4584 orthologs.

## 160 **2.7 Repeat annotation, gene prediction and gene annotation**

We first used MITE-Hunter(Han & Wessler, 2010), LTR-FINDER (version 1.05)(Xu & 161 Wang, 2007), RepeatScout (version 1.0.5)(Price, Jones, & Pevzner, 2005) and PILER(Edgar & 162 Myers, 2005) to construct a *de novo* repeat library for filefish with default settings. These 163 predicted repeats were classified using PASTEClassifer (version 1.0)(Hoede et al., 2014), and 164 then integrated with Repbase (19.06)(Bao, Kojima, & Kohany, 2015) to build a new repeat 165 library for final repeat annotation. In the end, RepeatMasker (version 4.0.6)(Tarailo-Graovac & 166 167 Chen, 2009) was performed to detect repetitive sequences in the filefish genome with the following parameters: "-nolow -no is -norna -engine wublast". 168

169	Ab initio-based, homolog-based, and RNA-sequencing (RNA-seq)-based methods were
170	conducted in combination to detect the protein-coding genes in filefish genome assembly.
171	Genscan(Burge & Karlin, 1997), Augustus (version 2.4)(Stanke & Waack, 2003),
172	GlimmerHMM (version 3.0.4)(Majoros, Pertea, & Salzberg, 2004), GeneID (version
173	1.4)(Blanco, Parra, & Guigó, 2007), and SNAP (version 2006-07-28)(Korf, 2004) were used
174	for <i>ab initio</i> -based gene prediction in filefish genome assembly. For the homolog-based
175	method, tiger pufferfish (Takifugu rubripes), spotted green pufferfish (Tetraodon nigroviridis)
176	and zebrafish (Danio rerio) were chosen to conduct gene annotation using GeMoMa (version
177	1.3.1)(Keilwagen et al., 2016). For the RNA-seq-based method, a mixture of 10 tissues
178	(including brain, eye, gill, heart, liver, intestine, spleen, ovary, kidney and muscle) of a
179	female and the testis of a male filefish was used to construct Illumina sequencing library and
180	subjected to PE (2 $\times$ 150 bp) sequencing on an Illumina HiSeq X Ten platform (Illumina, San
181	Diego, CA, USA). After discarding the reads with low-quality bases, adapter sequences, and
182	duplicated sequences, the retained high-quality clean reads were first assembled by Hisat
183	(version 2.0.4)(Kim, Langmead, & Salzberg, 2015) and Stringtie (version 1.2.3)(Pertea et al.,
184	2015), and then the gene prediction was performed using TransDecoder
185	(http://transdecoder.github.io) (version 2.0), GeneMarkS-T (version 5.1)(Tang, Lomsadze, &
186	Borodovsky, 2015), and PASA (version 2.0.2)(Haas et al., 2003). EVM (version 1.1.1)(Haas
187	et al., 2008) was performed to integrate the prediction results obtained from three methods.
188	We then added the genes that were supported by homolog and RNA-seq analysis after-manual
189	evaluation.

190	To functionally annotate the predicted genes, they were aligned to the Non-redundant
191	protein sequences (NR), eukaryotic orthologous groups of proteins (KOG)(Tatusov et al.,
192	2003), Kyoto Encyclopedia of Genes and Genomes (KEGG) (Kanehisa & Goto, 2000)and
193	TrEMBL(Boeckmann et al., 2003) databases using BLAST (version 2.2.31)(Altschul, Gish,
194	Miller, Myers, & Lipman, 1990) with an e-value cutoff of 1E-5. Gene ontology (GO)
195	(Consortium, 2004)annotation was performed with Blast2GO (version 4.1)(Conesa et al.,
196	2005). For non-coding RNA prediction, we first used tRNAscan-SE (version 1.3.1)(Lowe &
197	Eddy, 1997) to annotate transfer RNAs (tRNAs). Furthermore, Infenal (version 1.1)(Nawrocki
198	& Eddy, 2013) was conducted to search for ribosomal RNAs (rRNAs) and microRNAs based
199	on Rfam (version 13.0)(Daub, Eberhardt, Tate, & Burge, 2015) and miRbase (version
200	21.0)(Griffiths-Jones, Grocock, Van Dongen, Bateman, & Enright, 2006) database.
201	2.8 Comparative genomics
202	To resolve the phylogenetic position of the filefish, we first used OrthoMCL (version
203	
200	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data
204	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish
204 205	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish ( <i>Takifugu flavidus</i> ), spotted green pufferfish ( <i>Tetraodon nigroviridis</i> ), red seabream ( <i>Pagrus</i>
204 205 206	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish ( <i>Takifugu flavidus</i> ), spotted green pufferfish ( <i>Tetraodon nigroviridis</i> ), red seabream ( <i>Pagrus</i> <i>major</i> ), medaka ( <i>Oryzias latipes</i> ), large yellow croaker ( <i>Larimichthys crocea</i> ), three-spined
204 205 206 207	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish ( <i>Takifugu flavidus</i> ), spotted green pufferfish ( <i>Tetraodon nigroviridis</i> ), red seabream ( <i>Pagrus</i> <i>major</i> ), medaka ( <i>Oryzias latipes</i> ), large yellow croaker ( <i>Larimichthys crocea</i> ), three-spined stickleback ( <i>Gasterosteus aculeatus</i> ), nile tilapia ( <i>Oreochromis niloticus</i> ), japanese seabass
204 205 206 207 208	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish ( <i>Takifugu flavidus</i> ), spotted green pufferfish ( <i>Tetraodon nigroviridis</i> ), red seabream ( <i>Pagrus</i> <i>major</i> ), medaka ( <i>Oryzias latipes</i> ), large yellow croaker ( <i>Larimichthys crocea</i> ), three-spined stickleback ( <i>Gasterosteus aculeatus</i> ), nile tilapia ( <i>Oreochromis niloticus</i> ), japanese seabass ( <i>Lateolabrax maculatus</i> ), spotted gar ( <i>Lepisosteus oculatus</i> ) and zebrafish ( <i>Danio rerio</i> ). The
204 205 206 207 208 209	2.0.9) (Li, Stoeckert, & Roos, 2003) to detect orthologue groups by retrieving the protein data of eleven teleost species including tiger pufferfish ( <i>Takifugu rubripes</i> ), yellowbelly pufferfish ( <i>Takifugu flavidus</i> ), spotted green pufferfish ( <i>Tetraodon nigroviridis</i> ), red seabream ( <i>Pagrus</i> <i>major</i> ), medaka ( <i>Oryzias latipes</i> ), large yellow croaker ( <i>Larimichthys crocea</i> ), three-spined stickleback ( <i>Gasterosteus aculeatus</i> ), nile tilapia ( <i>Oreochromis niloticus</i> ), japanese seabass ( <i>Lateolabrax maculatus</i> ), spotted gar ( <i>Lepisosteus oculatus</i> ) and zebrafish ( <i>Danio rerio</i> ). The single copy orthologous genes shared by all 12 species were further aligned using MUSCLE

PhyML(Guindon et al., 2010). The divergence time among species was estimated by the
MCMCTree program of the PAML package(Yang, 2007) and CAFÉ(version 4.0) (De Bie,
Cristianini, Demuth, & Hahn, 2006) was used to identified expanded and contracted gene
families.

# 215 **3.Results and discussion**

# 216 **3.1 Initial characterization of the filefish genome**

The k-mer (k = 19 in this case) depth frequency distribution analysis of the 45.97 Gb 217 clean Illumina data was conducted to estimate the genome size, heterozygosity, and repeat 218 content of filefish (Table 1). The k-mer depth of 76 was found to be the highest peak in the 219 plot, and a k-mer number of 37,677,330,713 was used to calculate the genome size of filefish 220 221 (Figure S1). The sequences around k-mer depth of 38 were heterozygous sequences, and kmer depth more than 153 represented repetitive sequences. The filefish genome size was 222 223 estimated to be 491.74 Mb, the heterozygosity was approximately 0.35%, and the content of repetitive sequences and guanine-cytosine were about 16.62% and 46.05%, respectively. 224

#### 225 **3.2 Genome assembly**

A total of 50.95 Gb high quality clean reads, representing a 104-fold coverage of the genome, were generated from PromethION DNA sequencer (Table 1, Table S1-2,). These data was assembled using Wtdbg, followed by Racon and Pilon polishing, which produced a 465.93 Mb genome assembly with a surprising long contig N50 of 22.07 Mb (Table S3). The length of this assembly was close to the genome size estimated by k-mer analysis (491.74 Mb), indicating an appropriate assembly size was obtained from the Nanopore data. Among

the sequenced tetraodontiform species, the genome size of filefish was larger than *Takifugu* 

and *Tetraodon* species, but smaller than *Mola mola*(Aparicio et al., 2002; Gao et al., 2014;

234 Jaillon et al., 2004; Pan et al., 2016) (Table 2).

For Hi-C data, overall 39.44 Gb clean reads were obtained and used for subsequent

analysis (Table 1). To assess the quality of Hi-C data, we first made a plot of insert fragments

237 length frequency, which showed a relatively narrow unimodal length distribution with the

highest peak around 350 bp (Figure S2), indicating efficient purification of streptavidin beads

during library construction. The alignment results revealed that about 89.78% of the Hi-C

read pairs were mapped on the genome, and 78.18% of the read pairs were unique detected on

the assembly (Table S4). Lastly, a total of 47,111,219 valid reads, which accounted for

242 66.95% of the unique mapped reads, were detected by HiC-Pro in the Hi-C dataset (Table S5).

Taken together, our evaluation suggested an overall high quality of the Hi-C data, and only

the valid read pairs were used for subsequent analysis.

Before chromosomal-level genome assembly, an error correction of the initial assembly was performed by BWA-MEM with Hi-C data. The corrected filefish genome assembly was approximately 474.30 Mb with only 242 contigs, the contig N50 reached up to 22.46 Mb, and the longest contig was 32.32 Mb (Table 2, Table S6). The results indicated that high-coverage Nanopore long read-only assembly, followed by multiple iterations of genome polishing using Illumina reads is an effective method to generate high-quality genome assemblies.

A chromosomal-level genome was then assembled using LACHESIS, the results showed that overall 147 contigs spanning 471.65 Mb (99.44% of the assembly) were scaffolded into

253	20 pseudo-chromosomes, and 107 contigs spanning 469.46 Mb (98.98% of the assembly)
254	were successfully ordered and oriented (Table 3). Several of the pseudo-chromosomes were
255	scaffolded with only 2 or 3 contigs, representing a high contiguity of the genome. The final
256	assembled genome was 474.31 Mb with a scaffold N50 length of 23.05 Mb and a longest
257	scaffold of 34.81 Mb (Table 2, Table S6). As far as we know, this assembled genome was one
258	of the most contiguous fish genome assembly with the highest contig N50 when compared
259	with other published fish genomes.
260	To further evaluate the quality of the chromosomal-level genome assembly, a genome-
261	wide Hi-C heatmap was generated. The 20 pseudo-chromosomes could be easily
262	distinguished and the interaction signal strength around the diagonal was much stronger than
263	that of other positions within each pseudo-chromosome, which indicated a high quality of this
264	genome assembly (Figure 2).
265	3.3 Completeness of the assembled genome
266	Illumina reads were aligned to the filefish assembly, and 97.41% of the clean reads can
267	be mapped to the contigs (Table S7). Then the CEGMA analysis identified 442 CEGs,
268	accounting for 96.51% of all 458 CEGs in the program, and 226 CEGs could be detected by
269	using a highly conserved 248 CEGs dataset (Table S8). Lastly, approximately 94.33%
270	(4324/4584) of complete BUSCOs were found in the assembly (Table S9). Overall, the
271	assessment results indicated our filefish genome assembly was complete and of high quality.
272	3.4 Repeat annotation, gene prediction and gene annotation
273	A total of 67.35 Mb of repeat sequences that accounted for 14.2% of the assembly were

274	found in filefish (Table S10). This repeat content was close to the value (16.62%) obtained
275	from k-mer analysis. The predominant repeats type were TIRs (4.35%), LINEs (2.40%) and
276	LARDs (1.65%).

The combination of *Ab initio*-based, homolog-based, and RNA-seq-based methods predicted overall 22,067 protein-coding genes with an average gene length, average exon length, and average intron length of 11,291bp, 230 bp, and 905 bp, respectively (Table 1, Table 4). A total of 20,924 genes, which counted for 94.82% of the predicted genes, were successfully annotated with putative functions (Table 5). The non-coding RNA prediction identified 1,703 tRNAs, 649 rRNAs and 109 microRNAs, respectively (Table S11).

283 **3.5 Comparative genomics** 

Comparison of the filefish genome assembly with other eleven teleost species genomes 284 found a total of 22,665 gene families, of which 5,692 were shared among all eleven species, 285 including 1,872 single-copy orthologous genes (Table S12). Overall 20,261 genes of filefish 286 can be clustered into 15,433 gene families, including 67 unique gene families containing 193 287 genes (Table S12). The phylogenetic tree showed that four tetraodontiform species were 288 clustered together, and the divergence time between filefish and the other three species was 289 around 124.4 million years ago (Mya) (Figure 3). We also found 59 expanded gene families 290 and 98 contracted gene families in filefish compared with the other fish species (Figure S3). A 291 Venn diagram of orthologous gene families among four tetraodontiform species was also 292 293 constructed, and 971 unique gene families containing 6485 genes were identified in the filefish genome (Figure 4). 294

## 295 **4. Conclusion**

In the present study, we assembled the chromosome-level genome of T. septentrionalis, a 296 first reference genome of the genus Thamnaconus. The assembled genome was 474.31 Mb, 297 which is larger than the sequenced *Takifugu* and *Tetraodon* species, but smaller than *Mola* 298 299 mola. With the powerful sequencing ability of Oxford Nanopore technology, the contig N50 of the assembled genome achieved 22.46 Mb ,and the longest contig was 32.32 Mb. To the 300 best of our knowledge, this is the highest contig N50 among all the sequenced fish genomes. 301 This revealed that a combination of high-coverage Nanopore sequencing and Illumina data 302 polishing can effectively produce highly contiguous genome assemblies. The contigs were 303 clustered and ordered onto 20 pseudo-chromosomes with Hi-C data, and several pseudo-304 305 chromosomes were scaffolded with only 2 or 3 contigs. This high-quality genome will lay a strong foundation for a range of breeding, conservation and phylogenetic studies of filefish in 306 307 the future.

308

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# 491 Data Accessibility

- 492 Raw sequencing reads are available on GenBank as BioProject PRJNA565600. Raw
- 493 sequencing data (Nanopore, Illumina, Hi-C and RNA-seq data) have been deposited in SRA
- 494 (Sequence Read Archive) database as SRX6875837, SRX6862879, SRX6875660, and
- 495 SRX6875519.
- 496

# 497 Author Contributions

- 498 S.C., C.S. and Z.L. designed and managed the project. L.B., F.L. and J.G. interpreted the data
- and drafted the manuscript. P.W., S.Z., C.L. and X.L. prepared the materials. Q.C., J.L., K.L.
- and H.C. preformed the DNA extraction, RNA extraction and libraries construction. L.B.,
- 501 F.L., X.L. and C.S. performed the bioinformatic analysis. All authors contributed to the final
- 502 manuscript editing.

	TABLE 1 Statistics of the sequencing data					
Types	Method	Sequencing	Library size	Clean data	Coverage	
		platform	(bp)	(Gb)	$(\times)^{\dagger}$	
Genome	Illumina	Illumina HiSeq X	300	45.97	93.48	
Genome	Nanopore	PromethION	ultra-long	50.95	103.61	
Genome	Hi-C	Illumina HiSeq X	300	39.44	80.20	
Transcriptome	Illumina	Illumina HiSeq X	300	11.31	23.00	

## TABLE 1 Statistics of the sequencing data

<sup>†</sup> The coverage was calculated using an estimated genome size of 491.74 Mb.

#### TABLE 2 Assembly statistics of filefish and other tetraodontiform genomes

Species	T. septentrionalis	Takifugu rubripes <sup>†</sup>	Takifugu flavidus	Tetraodon nigroviridis	Mola mola
Sequencing technology	Oxford Nanopore	PacBio Sequel	PacBio Sequel	Plasmid library + BAC library	Illumina Hiseq 2000
	sequencing			sequencing	
Assembly size (Mb)	474.31	384.13	366.29	342.40	639.45
Number of scaffolds	155	128	867	25773	5552
N50 scaffold size (Mb)	23.05	16.71	15.68	0.73	8.77
Number of contigs	242	530	1111	41566	51826
N50 contig length (Mb)	22.46	3.14	4.36	0.03	0.02

<sup>†</sup> The assembly statistics of other tetraodontiform genomes were from NCBI assembly database. The GenBank assembly accession numbers were as follows: *Takifugu rubripes* (GCA\_901000725.2), *Takifugu flavidus* (GCA\_003711565.2), *Tetraodon nigroviridis* (GCA\_000180735.1), *Mola mola* (GCA\_001698575.1).

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## TABLE 3 Statistics of the pseudo-chromosome assemblies using Hi-C data

Group	Contig number	Contig length (bp)
Group 1	3	34,805,468
Group 2	3	34,142,503
Group 3	3	29,239,029
Group 4	13	27,092,115
Group 5	3	24,789,104
Group 6	7	24,144,372
Group 7	10	23,815,151
Group 8	3	23,107,901
Group 9	11	22,985,309
Group 10	5	23,048,615
Group 11	2	22,982,431
Group 12	6	23,025,906
Group 13	3	22,547,364
Group 14	11	22,005,842
Group 15	16	20,921,416
Group 16	3	20,603,809
Group 17	2	19,738,352
Group 18	5	17,694,734
Group19	13	18,094,054
Group 20	25	16,862,837
Total contigs clustered	147	471,646,312
Total contigs ordered and oriented	107	469,464,378

# TABLE 4 Summary of predicted protein-coding genes in the filefish genome

Method	Software	Species	Number of predicted genes
	Genscan		28,628
	Augustus		44,749
Ab inito	GlimmerHMM		34,576
	GeneID		24,446
	SNAP		58,914
		Takifugu rubripes	19,643
Homology-based	GeMoMa	Tetraodon nigroviridis	21,885
		Danio rerio	19,808
	PASA		30,768
RNA-seq	GeneMarkS-T		47,856
	TransDecoder		78,130
Integration	EVM		22,067

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TABLE 5 Summary of functional annotations for predicted genes

Annotation database	Annotated number of predicted genes	Percentage (%)	
GO	11,257	51.01%	
KEGG	13,714	62.15%	
KOG	14,760	66.89%	
TrEMBL	20,795	94.24%	
NR	20,905	94.73%	
All Annotated	20,924	94.82%	
Predicted Genes	22,067	-	





FIGURE 1 The greenfin horse-faced filefish (Thamnaconus septentrionalis)

COT



534FIGURE 2 The genome-wide Hi-C heatmap of the filefish. LG 1-20 are the abbreviations of Lachesis535Group 1-20, representing the 20 pseudo-chromosomes.



FIGURE 3 Phylogenetic analysis of the filefish with other teleost species. *Lepisosteus oculatus* was
 used as the outgroup. The estimated species divergence time (million years ago) and the 95%
 confidential intervals were labeled at each branch site.

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FIGURE 4 Venn diagram of orthologous gene families among four tetraodontiform species.