- 1 Transcriptome analysis of tetrodotoxin sensing and action of tetrodotoxin in central nervous
- 2 system of tiger puffer Takifugu rubripes juveniles

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#### Abstract

To reveal tetrodotoxin (TTX) sensing and action of TTX in central nervous system (CNS) of tiger puffer *Takifugu rubripes* juveniles, we conducted transcriptome analysis by next-generation sequencing for the olfactory and the brain of non-toxic cultured juveniles which were sensed and administered TTX. Sixty seven million reads from the nasal region (olfactory epithelium and skin) and the brain of each of three individuals of the control, TTX-sensed and TTX-administered juveniles were assembled into 153,958 contigs. A mapping of raw reads from the each sample onto the nucleotide sequences of predicted transcripts in *T. rubripes* genome (FUGU version 4) and the de novo assembled contigs, conducted to investigate their frequency of expression, revealed that the expression of 21 and 81 known genes significantly changed in TTX-sensed and TTX-administered juveniles in comparison with control juveniles, respectively. These genes included those related to feeding regulation and reward system, indicate that TTX ingestion of *T. rubripes* juveniles is controlled at feeding center in brain and *T. rubripes* may sense TTX as a reward, and accumulated TTX will directly act on CNS to adjust TTX ingestion.

- Keywords Takifugu rubripes Tetrodotoxin (TTX) Central nervous system RNA-seq Feeding
- 45 center Reward system

#### Introduction

Marine pufferfish of the genus *Takifugu* contain tetrodotoxin (TTX) which is one type of potent neurotoxin specific to voltage-gated sodium channels of excitable membranes of muscle and nerve tissues [1-3]. Matsumura [4] found that the toxin levels in embryos of grass puffer *Takifugu niphobles* increase from fertilization to hatching and concluded that TTX is produced by pufferfish. Other studies claimed that pufferfish accumulates TTX through food chain [3, 5], that is originally produced by marine bacteria belonging to the genera *Vibrio* and *Shewanella* [6-9]. The hypothesis that TTX in pufferfish is exogenous and is derived via the food chain is now widely accepted, because this hypothesis was supported by the fact that artificially raised tiger puffer *Takifugu rubripes* become nontoxic when fed with non-toxic diets in the environment where the invasion of TTX-bearing organisms was eliminated [10, 11], and such non-toxic *T. rubripes* are attracted to TTX [12, 13] and become toxic when they were fed with TTX-containing diets [14, 15, 16].

Non-toxic fishes can detect TTX at very low levels by gustatory organ [17]. Once non-toxic fishes ingest toxic eggs of pufferfish, they spit out pufferfish eggs immediately [18]. It was also confirmed that non-toxic fishes die even in trace amounts of TTX when administered directly into their bodies [19]. These evidences indicate that non-toxic fishes can recognize and avoid TTX as toxin. In contrast, *T. rubripes* detects TTX by olfactory organ, and actively ingests [13] and then accumulate high amounts of TTX [10]. Recently, several proteins implicated in the toxicity of pufferfish have been reported. Skeletal muscle voltage-gated Na<sup>+</sup> channel in pufferfish gain TTX resistance by amino acid substitutions in the P-loop region of the proteins [20-22]. Pufferfish saxitoxin and tetrodotoxin-binding proteins (PSTBPs) that bind to TTX and paralytic shellfish toxins were isolated from the plasma of panther puffer *Takifugu pardalis* and also found in the other *Takifugu* species [23, 24]. PSTBPs share high sequence homology (47 %) with a tributyltin-binding protein 2 (TBT-bp2) in Japanese flounder *Paralichthys olivaceus* [25], suggesting that PSTBPs originated in TBT-bp2s. These findings suggest that pufferfish become able to ingest TTX without recognizing as toxin through evolutional processes.

Generally liver and ovary of wild T. rubripes adults are strongly toxic [26]. However, in

juvenile stage, TTX is detected not only in liver but also in skin and brain of wild *T. rubripes* [16, 27]. It was further confirmed that TTX was transferred to skin and brain when TTX was administered to cultured non-toxic *T. rubripes* juveniles [27]. Since predation is a major cause of mortality in *T. rubripes* juveniles [28-30], bearing of TTX in skin may be functional as predator defense for the juvenile pufferfish [16]. Therefore, pufferfish utilize TTX for its survival through evolutional processes and alter the recognition of TTX as toxin for taking TTX into their body. Accumulation of TTX in brain [27] suggests that TTX passed through blood-brain barrier and was transferred to the central nervous system (CNS) of *T. rubripes* juveniles. Brain membranes of *T. pardalis* are harder to bind to saxitoxin that has the same Na<sup>+</sup> channel blocking function as TTX than corresponding membranes of rat same as skeletal muscle membranes including TTX-resistant Na<sup>+</sup> channel [20]. Thus, TTX may be functional in brain of pufferfish without blocking Na<sup>+</sup> channel.

Given these evidences, we hypothesized that *T. rubripes* juvenile senses TTX as a pharmacological agent and accumulated TTX is physiologically functional in CNS, and then some changes occur in the expression of genes associated with TTX sensing and action of TTX in CNS. Recently, next-generation sequencing technologies greatly improved the speed and efficiency of transcriptome analysis in many organisms including fishes [31] and the availability of the whole genome sequence of *T. rubripes* allowed us to use this technique. Thus, we conducted transcriptome analysis by next-generation sequencing for the olfactory and the brain of non-toxic cultured *T. rubripes* juveniles which were sensed and administered TTX.

# Materials and methods

# **Experimental fish**

- Non-toxic cultured T. rubripes juveniles (about 5 months old; body length,  $11.0 \pm 0.5$  cm; body weight,
- $37.7 \pm 4.1$  g; n = 150) were purchased from a private hatchery (Tawaki Suisan Corp., Kumamoto,
- 99 Japan) and were transported to Research Center for Marine Invertebrates, National Research Institute

of Fisheries and Environment of Inland Sea, Fisheries Research Agency, Momoshima, Hiroshima, Japan, in July 2014. The fish were fed with the commercial diets (Otohime EP3, Marubeni Nissin Feed Co., Ltd., Tokyo, Japan) in an aerated 5,000-l tank until TTX treatment.

### **Purification of TTX**

TTX was extracted from the ovary of a wild-caught adult *T. rubripes* according to the method of Ikeda et al. [32] with a slight modification. The extract was partially purified with Bio-Gel P-2 column (Bio-Rad Laboratories Inc., Herucles, CA, USA) and the absorbed TTX by the gel was eluted with 0.05 M acetic acid. TTX fraction was subjected to LC/MS analysis on an alliance LC/MS system equipped with a ZSpray MS 2000 detector (Waters, Milford, MA, USA) according to Nakashima et al. [33]. The amount of TTX (nanograms) determined by LC/MS was converted to mouse units (MU) based on the specific toxicity of TTX (220 ng/MU). Purified TTX was dried and frozen at -80°C until use.

### TTX-sensing and TTX-administration treatment to T. rubripes juveniles

Preliminary tests [13, 27] elucidated that non-toxic cultured juveniles were generally attracted to TTX within 30 minutes of starting to smell TTX and intramuscularly administered TTX in the fish was transferred to brain at least 24 hours after administration [unpublished data]. Based on these results, the following methods were established. For TTX-sensing treatment, three non-toxic cultured juveniles were accommodated in an aerated 30 l tank filled with 20-l fresh sand filtered seawater for 30 minutes as control, and three other non-toxic juveniles were sensed to TTX by immersing 200 MU (44 µg) TTX-containing seawater during the same period. For TTX-administration treatment, 0.1 ml of saline (1.35 % NaCl) as control and 150 MU (33 µg) TTX solution dissolved with saline was administered in a single injection into the dorsal muscle of three other non-toxic cultured juveniles using a 1 ml disposable syringe (Terumo, Tokyo, Japan), and the both groups of juveniles were immediately returned to the 90-l tank. Then, all fish were collected at 24 hours after administration.

The control, TTX-sensed and TTX-administered juveniles were anesthetized on ice, and then nasal region (olfactory epithelium and skin) and brain tissues were sampled, and stored in RNA later (Qiagen, Valencia, CA, USA) at -80°C until use.

## RNA extraction and cDNA library construction

Total RNA was extracted from the samples using RNeasy Mini Kit (Qiagen) following the manufacturer's instruction. The RNA samples were treated with DNase I (Takara, Tokyo, Japan) to digest contaminating genomic DNA. mRNA was then isolated from total RNA with Dynabeads® mRNA DIRECT™ Micro Kit (Life Technologies, Carlsbad, CA, USA). mRNA samples were fragmented, reverse transcribed and amplified to make barcoded whole transcriptome libraries using Ion Total RNA-seq Kit v2 (Life Technologies). Yield and size distribution of the fragmented RNA and the amplified cDNA were checked using an Agilent 2200 Tapestation with High Sensitivity RNA ScreenTape® and High Sensitivity D1000 ScreenTape® (Agilent Technologies, Palo Alto, CA, USA) at each step. We have performed a left size selection (< about 100 bp) with SPRIselect (Beckman Coulter, Krefeld, Germany) by using 1.2x volume of SPRI reagent to the nasal region samples. The average sizes of the amplified cDNAs were adjusted to be about 200 bp. Ion OneTouch™ System with Ion PI™ Template OT2 200 Kit v3 (Life Technologies) was used to prepare enriched, template-positive Ion PI™ Ion Sphere Particles.

### Next-generation sequencing and data analysis

The cDNA libraries were sequenced with an Ion Proton<sup>TM</sup> System with an Ion PI<sup>TM</sup> Sequencing 200 Kit v3 (Life Technologies) following the manufacturer's instructions. Sequencing results were imported into CLC Genomic Workbench7.5 (CLC bio, Aarhus, Denmark) as FASTQ files for further analysis. On CLC Genomic Workbench, the raw reads with the quality score less than 0.05 were trimmed using the "Trim Sequences" tool. Reads shorter than 50 bp were discarded. De novo sequence

assembly was carried out on all trimmed reads from all libraries using the Trinity software [34] to generate contigs. Duplicated and highly similar sequences were removed by the software CD-HIT (ver. 4.5.6. option, -c 0.9 [35]). Expression analysis was performed with RNA-seq Analysis Tool of CLC Genomics Workbench for each library using the nucleotide sequences of predicted transcripts in T. rubripes genome (FUGU version 4) cited from the Ensembl database and de novo assembled contigs as references, respectively. Parameters for read mapping were set as follows: Length fraction 0.7, similarity fraction 0.95. Gene expression was represented as RPKM (Reads Per Kilobase of exon model per Million). Cluster analysis based on the RPKM was performed by CLUSTER3.0 (http://bonsai.hgc.jp/~mdehoon/software/cluster/software.htm#ctv) using spearman correlation, and Java TreeView (http://jtreeview.sourceforge.net/) was used to visualize clustering relationship. Differential expression analysis between the control and TTX-sensed or TTX-administered juvenile samples was performed using R version 2.15.2 software (R Development Core Team 2008) package TCC with a false discovery rate (FDR) < 0.05 [36]. The homology searches of contigs detected as differential expression genes (DEGs) were conducted using BLASTX (e value 1e-5) against the NCBI non-redundant protein database. The DEGs assigned as unnamed protein products or uncharacterized proteins were excluded and we called them "known DEGs" in this paper.

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### Results

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# Sequencing and de novo assembly of nasal region and brain tissue transcripts

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Next-generation sequencing was conducted to generate expressed short reads from nasal region (olfactory epithelium and skin) and brain of the control, TTX-sensed and TTX-administered *T. rubripes* juveniles. We obtained 66,971,623 reads (2,192k – 3,325k reads/individual), with total nucleotides of 7,167,786,900 bp (231M – 364M bp/individual) (Table 1). Based on the reads, 153,958 contigs, with an average length of 648 bp were assembled (Table 2).

## Read mapping and gene annotation

The sequence reads were mapped to the nucleotide sequences of predicted transcripts in *T. rubripes* genome (FUGU version 4) cited from the Ensembl database and de novo assembled contigs to calculate the expression values. A hierarchical clustering analysis using the RNA-seq data analyzed by mapping to *T. rubripes* genome revealed that in Nasal region, the smaller and medium clusters tended to be form among samples (control and TTX) and between the trial groups (sensing and administration), respectively, and the larger clusters were formed between the tissues (nasal region and brain). However, the clusters were only formed about tissues by mapping to the constructed contigs (Fig. 1). The expression values were compared between the control and TTX-sensed or TTX-administered *T. rubripes* juveniles. The number of known DEGs detected under TTX-sensing treatment compared to the control were 4 (19.0 % of total number of DEGs) in nasal region and 17 (25.0 %) in brain, respectively (Table 3). In TTX-administration treatment, the number of known DEGs were 38 (37.3 %) in nasal region and 43 (35.0 %) in brain, respectively (Table 3).

### Expressed genes for TTX-sensed or TTX-administered juveniles

The distinctly expressed known DEGs in nasal region of TTX-sensed juveniles showed no high (fold change (FC) > 10) and low (FC < -10) expression levels, while in brain, relatively high and low expression levels were observed in several genes such as those encoding long-chain-fatty-acid--CoA ligase 5-like (FC value of 78.04), hemoglobin embryonic subunit alpha-like (FC value of 61.93) and peptide yy-like (FC value of -14.42) (Table 4, 5). The known DEGs which showed relatively high and low expression levels were also detected under TTX-administration treatment. In nasal region, extracellular superoxide dismutase (FC value of 36.79), envelope polyprotein (FC value of 15.22), receptor (chemosensory) transporter protein 4 (FC value of 12.20), podocalyxin-like (FC value of -24.90), tRNA-splicing endonuclease subunit sen15-like (FC value of -21.11), nuclear fragile x mental retardation-interacting protein 1-like (FC value of -20.19), period homolog 3 (drosophila) (FC value

of -16.71) and integrin alpha-3-like (FC value of -12.78) were detected (Table 6). In brain, potassium voltage-gated channel subfamily b member 2-like (FC value of 11.85), sorbin and sh3 domain-containing protein 2-like (FC value of 10.09) and period homolog 3 (drosophila) (FC value of -10.26) were detected (Table 7). In addition, several known DEGs were detected in both nasal region and brain of *T. rubripes* juvenile, such as those encoding long-chain-fatty-acid--CoA ligase 5-like under TTX-sensing treatment (Table 4, 5) and period homolog 3 (drosophila), envelope polyprotein, period circadian protein homolog 2-like and lipocalin precursor under TTX-administration treatment (Table 6, 7), and vasoactive intestinal peptide (vip) were down-regulated in brain under both TTX-sensing and TTX-administration treatment (Table 5, 7).

#### Discussion

In this study, we compared the gene expression in olfactory and brain among cultured *T. rubripes* juveniles with or without TTX-sensing and TTX-administration by transcriptome analysis using next-generation sequencing. Hierarchical cluster analysis of expressed genes was performed to assess the transcriptional pattern variation. In the case of using the RNA-seq data analyzed by mapping to *T. rubripes* genome revealed that in Nasal region, the smaller clusters tended to be form among samples (control and TTX), but the medium clusters tended to be form between the trial groups (sensing and administration) for each tissue. These results indicate that the gene expression in olfactory and brain of *T. rubripes* juveniles was affected by the operation and was not dramatically changed by TTX treatment. However, a number of DEGs detected under TTX-sensing and TTX-administration treatment compared to the control. Based on these DEGs, the following shows TTX sensing and action of TTX in CNS of *T. rubripes* juveniles.

### TTX sensing of *T. rubripes* juveniles

In nasal region (olfactory epithelium and skin) of TTX-sensed juveniles, mitogen-activated protein 4

kinase 4-like isoform x2 gene that is inhibitor of adipogenesis [37] was highest up-regulated than the fresh seawater-immersed control juveniles. In addition, long-chain-fatty-acid--CoA ligase 5 (ACSL5)-like gene which plays role in triacylglycerol (TAG) synthesis [38, 39] was up-regulated by TTX-sensing. These results and evidences suggest that TTX-sensing affects lipid metabolism in nasal region of *T. rubripes* juveniles. However, the expression of genes related to olfaction did not change by TTX-sensing. Given that cultured *T. rubripes* has not encountered TTX-bearing organisms, *T. rubripes* may instinctively sense TTX.

In brain of TTX-sensed juveniles, ACSL5-like and hemoglobin embryonic subunit alpha-like genes were extremely up-regulated than control fish. ACSL5 that is involved in TAG synthesis [38, 39] was also highly expressed in nasal region of TTX-sensed juveniles, suggesting that TTX-sensing particularly affects lipid metabolism in nervous system. Highly expression of one kind of hemoglobin, which is involved in oxygen transport, suggests that nervous activity is promoted in brain of TTX-sensed *T. rubripes* juveniles. Peptide yy (PYY)-like gene that has an appetite-regulation effect on fish [40-42] was down-regulated by TTX-sensing. In addition, vip peptides-like and TPA\_inf: tachykinin 1 genes which have a function as anorexigenic peptides in fishes [43, 44] were also down-regulated by TTX-sensing. Tachykinins is also related to dopaminergic system in mammals [45, 46]. In addition, Thy-1 membrane glycoprotein gene which may modulate dopamine metabolism in mammals [47] was down-regulated by TTX-sensing. If these evidences are applied to in fishes, some changes might occur in dopaminergic systems of TTX-sensed *T. rubripes* juveniles. Some studies have suggested the involvement of dopaminergic pathways in the central regulation of food intake in fishes [48-50]. Thus, TTX ingestion of *T. rubripes* juveniles is controlled at feeding center in brain and *T. rubripes* juveniles might sense TTX as a reward.

### Action of TTX in CNS of T. rubripes juveniles

In nasal region of TTX-administered juveniles, extracellular superoxide dismutase gene, which protects the living body from oxidative stress, was highest up-regulated than saline-administered

control juveniles. This study demonstrated the up-regulation of receptor (chemosensory) transporter protein 4 (RTP4). RTP family members are probable chaperon protein which facilitates trafficking and functional cell surface expression of some G-protein coupled receptors such as odorant receptor [51] and bitter taste receptor [52], suggesting that RTP4 is expressed in olfactory epithelium by TTXadministration and acts as a transporting protein of TTX sensing receptor. Podocalyxin-like gene, which is known to be expressed in the developing brain of the mouse and plays multiple roles in neural development [53], was lowest down-regulated. In addition, this study demonstrated the up- and downregulation of cyclin-dependent kinase inhibitor 1-like isoform x1 which associates with olfactory epithelium regeneration [54], and immunoglobulin superfamily member 8-like, which facilitates olfactory sensory synapse formation [55], respectively. In fishes, neurogenesis continues throughout life under the influence of environmental experience [56]. Synthesizing these results and evidences, we presume that nerve cell renewal occurs in the olfactory system of T. rubripes under the influence of TTX which exists in the olfactory epithelium. The expression of per genetic group which ticks in the center of cell clock [57] was specifically down-regulated by TTX-administration as following: period circadian protein homolog 1 -like isoform x1, period circadian protein homolog 2-like and period homolog 3 (Drosophila). These results suggest that biological rhythm of T. rubripes juveniles changed by accumulating TTX in their body. The core feedback loop of clock genes accurately ticks every 24 h [57]. Thus, there was another possibility that sampling times of juveniles was related to the clock genes expression.

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In brain of TTX-administered juveniles, potassium voltage-gated channel subfamily b member 2-like gene, which mediates membrane hyperpolarization during trains of action potentials [58, 59], was highest up-regulated than control fish. In addition, the expression of some genes which may be related to release of neurotransmitters changed by TTX-administration as following: clathrin, light chain [60], SRC kinase signaling inhibitor 1-like [61] and synaptotagmin-c-like [62]. SRC kinase signaling inhibitor 1 is involved in the formation and maintenance of synapses during developmental processes of brain [61]. Further, protein phosphatase 1B-like which involves in neurodegeneration [63] was up-regulated by TTX-administration, respectively. There are at least two main forms of neural

plasticity; biochemical switching and structural reorganization [64, 65]. Neural plasticity aids in the adaptation and flexibility demanded by the diverse environment in which fishes inhabit [66]. Non-toxic cultured *T. rubripes* juveniles is inferior in fear response comparing to the toxic wild juveniles, and release experiment into the pond with predators revealed that survival of cultured pufferfish with no TTX was significantly lower than that of toxic wild juveniles [28, 29]. These evidences suggest that *T. rubripes* juveniles utilize TTX to adapt to the environment with action of TTX in CNS. This study demonstrated the down-regulation of lipocalin precursor by TTX-administration in both nasal region and brain of *T. rubripes* juveniles. TBT-bp2 in the blood of *P. olivaceus* belongs to the lipocalin superfamily and shows highly identity to PSTBPs of *T. pardalis* [25]. From the fact that *T. rubripes* also have PSTBPs [24, 67, 68], the expression of lipocalin precursor may change in relation to the accumulation of TTX in their body. Interestingly, in brain of *T. rubripes* juveniles, vip which have a function as anorexigenic peptides in fish [44] were down-regulated by not only TTX-sensing but also TTX-administration. It may interpret that action of TTX to feeding center is not limited to only at the time of TTX ingestion, accumulated TTX also directly acts on CNS and adjust the intake.

In this study, we focused on the gene expression associated with TTX sensing and action of TTX in CNS of *T. rubripes* juveniles, thus did not concern the specificity of the fish to TTX. In the future, we need to use some other alkaloid such as palytoxin that is known for having other kinds of pufferfish [69] to investigate whether gene expression, behavioral and physiological change of *T. rubripes* juveniles are specific to TTX.

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# Tables

Table 1 Overview of the sequencing of cDNA from nasal region (olfactory epithelium and skin) and brain of TTX-sensed and TTX-administered Takifugu rubripes juveniles

	TTX-sensing treatment			TTX-administration treatment				
Items	Nasal region		Brain		Nasal region		Brain	
	Control	TTX	Control	TTX	Control	TTX	Control	TTX
Total number of reads	$2{,}587k\pm94k^a$	$3{,}325k\pm829k$	$2,192k \pm 197k$	$2,551k \pm 117k$	$3,\!073k\pm406k$	$3,\!052k\pm386k$	$2,\!870k\pm439k$	$2,675$ k $\pm 191$ k
Total nucleotide length (bp)	$261M\pm12M$	$355M \pm 77M$	$231M\pm17M$	$269M \pm 9M$	$364M \pm 51M$	$337M \pm 28M$	$299M \pm 36M$	$273M \pm 20M$

 $<sup>^{\</sup>text{a}}$  Results are shown as mean  $\pm$  SD of 3 fish

**Table 2** Summary of de novo assembly of contigs from sequence reads for nasal region (olfactory epithelium and skin) and brain of TTX-sensed and TTX-administered *Takifugu rubripes* juveniles

Items	Number
Total number of reads	66,971,623
Total nucleotide length (bp)	7,167,786,900
Total length of contigs (bp)	99,781,233
Number of contigs	153,958
Longest contig (bp)	17,962
Average length (bp)	648

Table 3 Number of differential expression genes (DEGs) detected in nasal region (olfactory epithelium and skin) and brain of TTX-sensed and TTX-administered Takifugu rubripes juveniles

	Items	TTX-sensing treatment			TTX-administration treatment				
Mapping reference		Nasal region		Brain		Nasal region		Brain	
		Up-regulation	Down-regulation	Up-regulation	Down-regulation	Up-regulation	Down-regulation	Up-regulation	Down-regulation
T. rubripes genome	Total Number of DEGs	0	0	3	1	2	7	5	8
	Number of known DEGs	0	0	2	1	1	6	3	6
Contigs	Total Number of DEGs	14	7	31	33	48	45	50	60
	Number of known DEGs	3	1	5	9	14	17	18	16

**Table 4** Genes that were up- and down-regulate in nasal region (olfactory epithelium and skin) of TTX-sensed *Takifugu rubripes* juveniles analyzed by mapping to contigs (FDR-corrected *p*-value <0.05)

Contig ID	Carra	Expression in RPKM	Expression in RPKM $^a$ (mean $\pm$ SD, n=3)		
Contig ID	Gene	Control	TTX	change	
c58136_g1_i1	Mitogen-activated protein 4 kinase 4-like isoform x2	$21,\!892 \pm 20,\!896$	$135,\!241 \pm 65,\!074$	6.18	
c61343_g1_i2	Calpain-1 catalytic subunit-like	$265,\!065 \pm 78,\!263$	$442,\!603 \pm 8,\!410$	1.67	
c73927_g1_i2	Long-chain-fatty-acidCoA ligase 5-like	$\mathrm{ND^b}$	$120,\!695 \pm 38,\!720$	$NA^c$	
c81231_g1_i3	Bromodomain-containing protein 3-like isoform x1	$319,\!451 \pm 48,\!324$	$65,\!555 \pm 27,\!773$	-4.87	

<sup>&</sup>lt;sup>a</sup> RPKM: reads per kilobase of exon model per million mapped reads

<sup>&</sup>lt;sup>b</sup> ND: not detected

<sup>°</sup> NA: not applicable

**Table 5** Genes that were up- and down-regulated in brain of TTX-sensed *Takifugu rubripes* juveniles analyzed by mapping to *T. rubripes* genome and contigs (FDR-corrected *p*-value <0.05)

ID.	Com	Expression in RPKM	Fold	
ID	Gene -	Control	TTX	change
Differential expression ge (Ensembl ID)	nes (DEGs) detected by mapping to T. rubripes genome			
ENSTRUT00000043560	Spermine synthase	$9.3\pm1.4$	$39.8 \pm 11.6$	4.28
ENSTRUT00000046894	Centromere protein N	$ND^b$	$9.2 \pm 4.7$	NAc
ENSTRUT00000047470	Peripherin	$69.6\pm30.9$	$11.2\pm3.6$	-6.19
Differential expression ge (Contig ID)	nes (DEGs) detected by mapping to contigs			
c73927_g1_i2	Long-chain-fatty-acid CoA ligase 5-like	$233.5\pm404.4$	$18,\!221.6 \pm 8,\!607.3$	78.04
c51638_g1_i1	Hemoglobin embryonic subunit alpha-like	$426.0 \pm 737.9$	$26,\!383.3 \pm 22,\!612.7$	61.93
c71980_g1_i2	NADH dehydrogenase	$14,\!183.9 \pm 6,\!567.4$	$41{,}755.4 \pm 22{,}042.0$	2.94
c95098_g2_i1	Zinc finger protein Eos	$48,\!870.1 \pm 27,\!875.8$	$139,\!364.0\pm11,\!130.5$	2.85
c80686_g1_i1	General transcription factor IIF subunit 1-like	$19,\!245.8 \pm 2,\!027.7$	$47,\!290.7 \pm 11,\!002.6$	2.46
c70981_g1_i1	Peptide yy-like	$8,\!298.1 \pm 3,\!566.7$	$575.3 \pm 996.5$	-14.42
c78080_g1_i1	Urotensin ii-related peptide precursor	$24,\!257.1 \pm 8,\!487.6$	$3,352.4 \pm 2,607.7$	-7.24
c87450_g1_i1	Vip peptides-like	$25,\!643.6 \pm 16,\!773.2$	$4,403.0 \pm 2,267.9$	-5.82
c47673_g1_i1	Ras-related protein rab-8b-like	$10,\!444.2 \pm 4,\!183.4$	$2,\!020.7 \pm 2,\!096.5$	-5.17
c61457_g1_i1	Fibroblast growth factor receptor substrate 2-like	$189,\!326.3 \pm 94,\!323.7$	$77,\!067.8 \pm 20,\!944.3$	-2.46
c79330_g1_i1	Thy-1 membrane glycoprotein	$242,\!220.3 \pm 71,\!740.0$	$131,\!042.2\pm26,\!743.4$	-1.85
c90268_g1_i1	TPA_inf: tachykinin 1	$97,141.8 \pm 17,982.3$	$69,\!015.0 \pm 5,\!749.7$	-1.41
c90791_g1_i1	Neurobeachin-like isoform x3	$59,\!884.0 \pm 35,\!206.2$	$59,\!172.9 \pm 22,\!897.8$	-1.01
c75435_g1_i1	Growth hormone	$114,\!851.9 \pm 198,\!929.4$	ND	NA

<sup>&</sup>lt;sup>a</sup> RPKM: reads per kilobase of exon model per million mapped reads

<sup>&</sup>lt;sup>b</sup> ND: not detected

<sup>&</sup>lt;sup>c</sup> NA: not applicable

**Table 6** Genes that were up- and down-regulated in nasal region (olfactory epithelium and skin) of TTX-administered *Takifugu rubripes* juveniles analyzed by mapping to *T. rubripes* genome and contigs (FDR-corrected *p*-value <0.05)

ID	Com	Expression in RPKM	Fold	
ID	Gene	Control	TTX	change
	nes (DEGs) detected by mapping to T. rubripes genome			
(Ensembl ID) ENSTRUT00000016957	Receptor (chemosensory) transporter protein 4	$1.1 \pm 1.1$	$13.7 \pm 2.8$	12.20
ENSTRUT00000016937 ENSTRUT00000046106	Podocalyxin-like	$20.3 \pm 16.7$	$0.8 \pm 1.4$	-24.90
ENSTRUT00000040100	Period homolog 3 (Drosophila)	$42.4 \pm 21.3$	$2.5 \pm 1.7$	-16.71
ENSTRUT00000007590	Period homolog 3 (Drosophila)	$28.9 \pm 6.2$	$3.7 \pm 2.8$	-7.74
ENSTRUT00000003686	Complement component 8, gamma polypeptide	$1,914.5 \pm 723.9$	$463.4 \pm 148.9$	-4.13
ENSTRUT00000023711	Tubulin, alpha 2	$112.8 \pm 29.6$	$29.4 \pm 2.3$	-3.83
ENSTRUT00000038281	Circadian associated repressor of transcription	$14.4 \pm 10.5$	ND <sup>b</sup>	NA°
E116111610000030201	Cheudain associated repressor of manoripuon	11.12.10.3	T(D	1171
Differential expression ge (Contig ID)	nes (DEGs) detected by mapping to contigs			
c63197_g1_i1	Extracellular superoxide dismutase	$60.6\pm105.0$	$2,\!229.8 \pm 1,\!997.3$	36.79
c87942_g1_i5	Envelope polyprotein	$108.8\pm127.7$	$1,\!655.2 \pm 145.6$	15.22
c77064_g1_i2	Cyclin-dependent kinase inhibitor 1-like isoform x1	$761.1 \pm 397.1$	$5,517.0 \pm 2,584.5$	7.25
c79561_g1_i2	Cytoplasmic dynein 1 intermediate chain 2-like isoform x3	$1,283.2 \pm 89.2$	$7,521.1 \pm 6,305.5$	5.86
c62397_g1_i1	Protein inscuteable homolog	$481.7 \pm 261.8$	$2{,}723.7 \pm 816.7$	5.65
c62397_g1_i1	Diamine acetyltransferase 1-like	$10,\!569.0 \pm 693.0$	$29{,}130.7 \pm 6{,}054.7$	2.76
c81702_g1_i1	Double stranded rna-activated protein kinase 2	$1,\!551.9 \pm 1,\!704.3$	$3,931.7 \pm 1,071.3$	2.53
c85001_g1_i7	Tumor necrosis factor receptor superfamily member 4-like	$5,\!622.7 \pm 1,\!286.2$	$13,\!152.5\pm3,\!530.8$	2.34
c77678_g1_i2	Mannose-specific lectin-like	$194,\!004.1 \pm 15,\!127.9$	$362,\!323.5 \pm 41,\!810.5$	1.87
c77678_g1_i1	Lily-type lectin	$104,\!210.6 \pm 7,\!187.5$	$194,\!205.6 \pm 10,\!327.1$	1.86
c77678_g1_i5	Mannose-specific lectin-like	$91,\!364.8 \pm 12,\!267.0$	$166,\!608.0 \pm 6,\!407.8$	1.82
c90536_g2_i1	Ribonucleoside-diphosphate reductase subunit m2-like isoform x1	$4,\!292.3 \pm 1,\!508.3$	$7,\!823.3 \pm 2,\!087.8$	1.82
c91998_g1_i4	Apoptosis facilitator bcl-2-like protein 14	$2,\!139.7 \pm 586.8$	$3{,}790.4 \pm 1{,}082.3$	1.77
c62456_g1_i1	Serine threonine-protein kinase psk2	ND	$3,\!232.8 \pm 4,\!607.5$	NA
c60649_g1_i1	tRNA-splicing endonuclease subunit sen15-like	$1,\!649.6 \pm 862.9$	$78.2\pm135.4$	-21.11
c59514_g1_i1	Nuclear fragile x mental retardation-interacting protein 1-like	$2,\!010.7 \pm 1,\!146.2$	$99.6\pm172.5$	-20.19
c89300_g1_i3	Integrin alpha-3-like	$1,409.7 \pm 255.1$	$110.3\pm191.0$	-12.78
c55712_g1_i1	Isoleucinetrna cytoplasmic-like	$1,\!610.1 \pm 250.9$	$172.2\pm149.5$	-9.35
c63419_g1_i3	Lysyl oxidase	$2,\!177.1 \pm 310.6$	$233.7 \pm 202.9$	-9.31
c89139_g2_i1	Protein capicua homolog isoform x3	$1,\!901.6 \pm 920.7$	$216.6 \pm 192.3$	-8.78
c91950_g1_i1	Period circadian protein homolog 2-like	$9,\!901.5 \pm 860.0$	$1,\!685.4 \pm 919.8$	-5.88
c15805_g1_i1	Immunoglobulin superfamily member 8-like	$11,\!099.7 \pm 8,\!539.1$	$2,\!290.3 \pm 573.9$	-4.85
c59867_g1_i1	Salivary glue protein	$7,\!503.1 \pm 2,\!447.5$	$1{,}745.9 \pm 395.5$	-4.30
c52572_g1_i1	Polyhomeotic-like protein 3-like isoform x3	$2,979.2 \pm 585.1$	$749.2 \pm 408.3$	-3.98
c84319_g1_i3	SEC14-like protein 2-like	$3,\!540.0 \pm 959.1$	$981.0 \pm 719.3$	-3.61
c95794_g1_i2	C-terminal binding protein 1	$4{,}728.5 \pm 1{,}069.8$	$1,\!509.4 \pm 581.8$	-3.13
c80106_g2_i1	Lipocalin precursor	$179{,}726.2 \pm 53{,}330.8$	$62,\!486.5 \pm 20,\!641.1$	-2.88
c94326_g5_i1	Elongation of very long chain fatty acids protein 6-like	$2,\!132.0 \pm 984.2$	$749.9 \pm 303.0$	-2.84
c85958_g2_i1	Cytochrome c oxidase subunit ii	$773,\!930.0 \pm 36,\!745.0$	$641,\!615.3 \pm 51,\!394.4$	-1.21
c62014_g1_i1	LIM domain and actin-binding protein 1-like	$7,\!473.6 \pm 6,\!187.9$	$6,\!769.8 \pm 2,\!706.2$	-1.10

<sup>a</sup> RPKM: reads per kilobase of exon model per million mapped reads

<sup>b</sup> ND: not detected

<sup>c</sup> NA: not applicable

**Table 7** Genes that were up- and down-regulated in brain of TTX-administered *Takifugu rubripes* juveniles analyzed by mapping to *T. rubripes* genome and contigs (FDR-corrected *p*-value <0.05)

ID	Com	Expression in RPK	Fold	
ID	Gene	Control	TTX	change
	nes (DEGs) detected by mapping to T. rubripes genome			
(Ensembl ID) ENSTRUT00000043847	LIM domain only 2 (rhombotin-like 1)	$7.9 \pm 3.1$	$43.4 \pm 15.4$	5.48
ENSTRUT00000039938	Clathrin, light chain (Lca)	$6.8 \pm 2.3$	$35.7 \pm 13.3$	5.24
ENSTRUT000000039938	Family with sequence similarity 192, member A	ND <sup>b</sup>	$7.2 \pm 1.2$	NA°
ENSTRUT00000007585	Period homolog 3 (Drosophila)	$27.4 \pm 4.3$	$7.2 \pm 1.2$ $2.7 \pm 1.3$	-10.26
	NHP2 non-histone chromosome protein 2-like 1b			
ENSTRUT00000043060	(Saccharomyces cerevisiae)	$25.4 \pm 19.0$	$2.9 \pm 1.1$	-8.65
ENSTRUT00000022965	Pterin-4 alpha-carbinolamine dehydratase/dimerization	$28.6\pm19.6$	$3.5\pm2.3$	-8.07
ENSTRUT00000008208	cofactor of hepatocyte nuclear factor 1 alpha (TCF1) 2 Immunoglobulin heavy variable 1-4	$40.8 \pm 24.0$	$7.9 \pm 2.2$	-5.14
ENSTRUT00000044170	Cytochrome P450, family 27, subfamily C, polypeptide 1	$29.2 \pm 4.9$	$8.2 \pm 3.3$	-3.56
ENSTRUT000000044170	Vasoactive intestinal peptide	$16.6 \pm 19.5$	8.2 ± 3.3 ND	NA
LIVSTRC 100000002071	v asouetive intestinai peptide	10.0 ± 17.5	ND	1471
Differential expression ge	nes (DEGs) detected by mapping to contigs			
(Contig ID)				
c33428_g1_i1	Potassium voltage-gated channel subfamily b member 2-like	$91.3 \pm 31.2$	$1,081.1 \pm 598.5$	11.85
c50125_g1_i1	Sorbin and sh3 domain-containing protein 2-like	$92.2\pm80.6$	$930.5 \pm 105.7$	10.09
c61302_g1_i1	Transmembrane protein 119-like	$100.1 \pm 109.9$	$918.0 \pm 464.2$	9.17
c33376_g1_i1	Ubiquitin-conjugating enzyme e2 o-like	$148.6 \pm 156.4$	$1,238.8 \pm 442.8$	8.34
c87942_g1_i5	Envelope polyprotein	$289.5 \pm 107.9$	$2,169.1 \pm 76.3$	7.49
c10845_g1_i1	Ankyrin repeat and sterile alpha motif domain containing 1b	$231.3 \pm 205.0$	$1,\!700.9 \pm 1015.0$	7.36
c81025_g1_i1	Protein nynrin-like	$279.8 \pm 28.8$	$1,876.9 \pm 462.9$	6.70
c50509_g1_i1	Ubiquitin-conjugating enzyme e2 r1-like	$153.0\pm41.6$	$1,\!020.9 \pm 409.7$	6.67
c64715_g1_i1	Protein phosphatase 1B-like	$613.6 \pm 367.0$	$2,\!661.5 \pm 865.0$	4.34
c78791_g3_i1	Apoptogenic protein mitochondrial-like	$480.3 \pm 402.1$	$1,\!849.8 \pm 1,\!000.8$	3.85
c88645_g4_i3	SRC kinase signaling inhibitor 1-like	$1,\!280.5 \pm 621.3$	$2,\!083.3 \pm 136.7$	1.63
c82175_g1_i2	Serine threonine-protein kinase 38-like	$2,\!230.3 \pm 898.8$	$3,\!210.7 \pm 470.3$	1.44
c86443_g1_i1	PAX3- and PAX7-binding protein 1	$35{,}715.3 \pm 6{,}540.5$	$37,\!981.7 \pm 1,\!870.0$	1.06
c92296_g2_i2	Tubulin alpha-1A chain-like	$273,\!304.4\pm3,\!334.7$	$285,\!511.9 \pm 2,\!173.3$	1.04
c13692_g1_i1	Ribosomal protein L29	$45,\!614.5 \pm 4,\!756.2$	$47,\!495.1 \pm 1,\!940.2$	1.04
c113887_g1_i1	Supervillin-like isoform x5	ND	$1,\!359.3 \pm 1,\!989.9$	NA
c149_g1_i1	Star-related lipid transfer protein 13-like	ND	$1,\!263.5 \pm 1,\!570.3$	NA
c57536_g1_i4	Neuronal pas domain-containing protein 2-like	ND	$1,141.5 \pm 500.8$	NA
c79100_g1_i2	Chymotrypsin-like elastase family member 2a-like	$1,171.3 \pm 1,374.8$	$189.6 \pm 126.0$	-6.20
c92921_g2_i1	Period 1	$1{,}780.3 \pm 178.6$	$290.9 \pm 229.9$	-6.10
c57364_g1_i1	Pterin-4-alpha-carbinolamine dehydratase 2-like	$1,\!400.4 \pm 468.0$	$329.0 \pm 91.2$	-4.26
c91950_g1_i1	Period circadian protein homolog 2-like	$2,653.6 \pm 152.5$	$894.3\pm80.5$	-2.97
c85532_g1_i2	Immunoglobulin mu heavy chain	$2,\!279.3 \pm 909.3$	$814.9 \pm 421.1$	-2.80
c51266_g1_i1	Rho GTPase-activating protein 23-like isoform x9	$6,\!700.8 \pm 3,\!078.6$	$2,\!570.6 \pm 1,\!353.0$	-2.61
c55701_g1_i1	Synaptotagmin-c-like	$1,\!958.7 \pm 246.0$	$782.5\pm395.3$	-2.50
c76601_g1_i1	Period circadian protein homolog 1-like isoform x1	$7,\!336.8 \pm 672.7$	$3,601.1 \pm 1,293.5$	-2.04
c77744_g1_i1	Period circadian protein homolog 1-like isoform x1	$4,\!053.6 \pm 474.7$	$1,\!990.0 \pm 376.7$	-2.04

c85980_g1_i1	Nuclear receptor subfamily 1 group d member 2-like	$3,\!015.9 \pm 803.3$	$1,\!641.8 \pm 347.3$	-1.84
c96661_g4_i1	Polyadenylate-binding protein 2-like isoform x3	$5,\!808.0 \pm 822.8$	$3,673.1 \pm 277.3$	-1.58
c80106_g2_i1	Lipocalin precursor	$66,744.1 \pm 8,471.7$	$52,\!467.3 \pm 4,\!148.2$	-1.27
c77464_g1_i1	Protein FAM107B-like	$9,727.2 \pm 1,094.3$	$7,\!685.5 \pm 863.4$	-1.27
c32430_g1_i1	60S acidic ribosomal protein P2	$54,877.1 \pm 2,565.7$	$50,\!338.9 \pm 3,\!830.0$	-1.09
c140829_g1_i1	Unconventional myosin-xviiib-like	$1,\!257.2 \pm 1,\!929.3$	ND	NA
c126916_g1_i1	Polycystin-1-like	$1,127.9 \pm 1,685.0$	ND	NA

<sup>&</sup>lt;sup>a</sup> RPKM: reads per kilobase of exon model per million mapped reads

<sup>&</sup>lt;sup>b</sup> ND: not detected

c NA: not applicable

#### Figure legend

Fig. 1 Hierarchical clustering dendrograms from the RNA-seq analyzed by mapping to *Takifugu rubripes* genome (a) and contigs (b). The numbers represent independent samples. The vertical scale represents between-cluster distance

