Transcriptomic analysis of the host response to an iridovirus infection in Chinese giant salamander, *Andrias davidianus*

Yuding Fan, Ming Xian Chang, Jie Ma, Scott E. Lapatra, Yi Wei Hu, Lili Huang, Pin Nie, Lingbing Zeng

To cite this version:

Yuding Fan, Ming Xian Chang, Jie Ma, Scott E. Lapatra, Yi Wei Hu, et al.. Transcriptomic analysis of the host response to an iridovirus infection in Chinese giant salamander, *Andrias davidianus*. Veterinary Research, BioMed Central, 2015, 46 (1), pp.136. 10.1186/s13567-015-0279-8. hal-01341445

HAL Id: hal-01341445
https://hal.archives-ouvertes.fr/hal-01341445
Submitted on 4 Jul 2016

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Transcriptomic analysis of the host response to an iridovirus infection in Chinese giant salamander, *Andrias davidianus*

Yuding Fan1,4*, Ming Xian Chang2†, Jie Ma1, Scott E. LaPatra3, Yi Wei Hu2, Lili Huang1, Pin Nie2* and Lingbing Zeng1*

**Abstract**

The emergence of an infectious viral disease caused by the Chinese giant salamander iridovirus (GSIV) has led to substantial economic losses. However, no more molecular information is available for the understanding of the mechanisms associated with virus–host interaction. In this study, de novo sequencing was used to obtain abundant high-quality ESTs and investigate differentially-expressed genes in the spleen of Chinese giant salamanders that were either infected or mock infected with GSIV. Comparative expression analysis indicated that 293 genes were down-regulated and 220 genes were up-regulated. Further enrichment analysis showed that the most enriched pathway is “complement and coagulation cascades”, and significantly enriched diseases include “inherited thrombophilia”, “immune system diseases”, “primary immunodeficiency”, “complement regulatory protein defects”, and “disorders of nucleotide excision repair”. Additionally, 30,678 simple sequence repeats (SSRs) from all spleen samples, 26,355 single nucleotide polymorphisms (SNPs) from the spleens of uninfected animals and 36,070 SNPs from the spleens of infected animals were detected. The large amount of variation was specific for the Chinese giant salamanders that were infected with GSIV. The results reported herein provided significant and new EST information that could contribute greatly in investigations into the molecular functions of immune genes in the Chinese giant salamander.

© 2015 Fan et al. This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

**Introduction**

Amphibians are an important evolutionary bridge between aquatic and terrestrial vertebrates [1]. The Chinese giant salamander, *Andrias davidianus*, is the largest extant amphibian species and is considered a living fossil because it has existed for more than 350 million years [2]. The phylogenetic position of the Chinese giant salamander makes it an invaluable model for evolutionary and comparative studies. The Chinese giant salamander also has significant economic value as an edible delicacy and for medicinal purposes. However, in the past 50 years the Chinese giant salamander population has declined sharply due to deterioration of habitat, environmental pollution, climate change, infectious diseases and commercial trade [3, 4]. Currently, artificial breeding and culture is being used to protect the Chinese giant salamander. Approximately two million Chinese giant salamanders are bred annually in China.

With the rapid expansion of Chinese giant salamander farming, emerging infectious diseases have been increasing. Viral diseases, including the iridovirus, have caused major impacts to the Chinese giant salamander industry [5, 6]. The economic losses caused by the Chinese giant salamander iridovirus (GSIV) reached 300 million RMB (48 million USD) in 2010. There is an urgent need to understand the immune system of the Chinese giant salamander and the pathogenic mechanism(s) associated with a GSIV infection. In previous studies, the morphogenesis, pathological changes, rapid detection methods and virion-associated viral proteins of GSIV have been
reported [7–12]. Although a ranavirus-induced thymus cDNA library and two immune organs (skin and spleen) from the healthy Chinese giant salamanders were sequenced [13, 14], and several immune genes were reported [15–17], the molecular information available is still limited for the Chinese giant salamander and that has hindered the understanding of the molecular mechanisms associated with viral infection and virus–host interactions.

Next-generation sequencing technology, such as de novo transcriptome sequencing, can be used for large-scale efficient and economical production of ESTs, and has become an important method for studying non-model species [18, 19]. Transcriptome sequencing facilitates functional genomic studies, including global gene expression, novel gene discovery, assembly of full-length genes, simple sequence repeats (SSRs) and single nucleotide polymorphism (SNP) discovery [20].

This technology has also been used widely in comparative transcriptomics to identify differences in transcript abundance among different developmental stages and under different treatment conditions [21, 22]. In the present study, the spleen transcriptome of the Chinese giant salamander was sequenced by de novo sequencing technology, and a comparative analysis of transcriptome data was performed between a control and a group infected with GSIV. The results provided a significant amount of information on the genes in the Chinese giant salamander, and suggested the conservation and divergence of several important immune signaling pathways. The different genes expressed and enrichment analysis of pathways could contribute significant new information regarding the pathogenic mechanism(s) of the virus and the interaction(s) of the virus and the host.

Materials and methods

Chinese giant salamanders, viral challenge and sample collection

Chinese giant salamanders (average weight, ~180 g) were obtained from the research farm of the Yangtze River Fisheries Research Institute in Wuhan, China. Prior screening indicated that these animals were free of GSIV. All the salamanders were kept in aerated, tap water supplied tanks at 20 °C and fed with diced bighead carp (Hypophthalmichthys nobilis) for 2 weeks prior to the experiment. The virus suspension used to infect a portion of the salamanders was obtained from GSIV-infected EPC cells. The fish in infected group (GS_TS) were injected intramuscularly with 0.2 mL of the GSIV suspension (5 × 10^7 TCID50/mL), and fish in control group (GS_CS) were injected intramuscularly with equal volume of DPBS (Sigma). The spleens from three individual salamanders from each group were collected at 48 h post-injection.

RNA isolation and cDNA synthesis

Total RNA was extracted from the spleens using TRIzol® Reagent (Invitrogen, USA). Samples of the three individuals from each treatment group were pooled in equal amounts to generate one RNA sample per group. These two RNA samples were sent to Shanghai Majorbio Bio-pharm Biotechnology Co., Ltd. (Shanghai, China) for the cDNA library construction and Illumina deep sequencing.

cDNA library construction and Illumina deep sequencing

Two cDNA libraries were prepared using the TrueSeqTM RNA sample prep Kit (Illumina, San Diego, CA, USA) following the manufacturer’s instructions. Briefly, poly (A)+ RNA was purified from 5 μg of pooled total RNA using oligo (dT) magnetic beads, sheared into short fragments, and primed for cDNA library synthesis using the TrueSeq RNA sample preparation kit according to the manufacturer’s instructions (Illumina). After quantitation using a TBS-380 minifluorometer (PicoGreen), the samples were clustered (TrueSeq paired-end cluster kit v3-cBot-HS; Illumina) and sequenced on the HiSeq 2000 platform (100 bp, TrueSeq SBS kit v3-HS 200 cycles; Illumina).

Data analysis

The raw reads from the images were generated using Solexa GA pipeline 1.6. After the removal of low-quality reads, processed reads with an identity value of 95% and a coverage length of 100 bp were assembled using the Trinity de novo assembler [23]. The isogenes generated were compared with the NCBI non-redundant (nr) database using the BLASTx algorithm, with a cut-off E value of ≤10^{-5}. GO terms were extracted from the best hits obtained from the BLASTx against the nr database using Blast2GO [24]. These results were then sorted by GO categories using in-house Perl scripts. BLASTx was also used to align unique sequences to the Swiss-Prot database, Kyoto Encyclopedia of Genes and Genomes (KEGG) and Clusters of Orthologous Groups (COG) (with the e value of 10^{-6}) to predict possible functional classifications and molecular pathways [25].

Differential expression analysis

To identify differentially expressed genes/isogenes between infected and uninfected groups, genes/isogenes expression levels were measured by using numbers of fragments per kilobase of transcript per million fragments sequenced (FPKM) [26], similar to RPKM (reads
per kilobase of gene model exon per million mapped reads) measure used earlier [27]. The differential expression analysis was carried out using RSEM [28] and edgeR [29] softwares. For each gene/isogene, the \( p \) value was computed, and then Benjamini–Hochberg false discovery rate (FDR) was applied to correct the results for \( p \) value. The transcripts that were increased or decreased at an estimated absolute log2-fold change of >1 and FDR adjusted \( p \) value \( \leq 0.05 \) were considered to be differentially expressed.

Identification of EST-SSR motifs and EST-SNPs

MSATCOMMANDER V. 0.8.2 [30] was used to analyze the microsatellite (SSR) distribution. The minimum number of repeats for SSR detection was six for di-SSRs and four repeats for tri-, tetra-, penta-, and hexa-SSRs. The open reading frame (ORF) and untranslated region (UTR) within the isotig were identified using Trinity [23]. The location of SSRs was estimated based on ORFs and UTRs. SSR-containing isotigs were annotated based on BLAST similarity searches. SNPs were detected based on alignment using BWA V. 0.5.9 [31] and SAMtools V. 0.1.18 [32]. From the “pileup” output of SAMtools, VarScan V.2.2.7 filtered SNPs based on the following criteria including (1) the total coverage and the number of reads to cover a candidate SNP (>8 reads); (2) the base quality where base calls with low Phred quality (<25) were removed from the coverage; and (3) frequency of mutated bases higher than 30% among all reads covering the position.

Quantitative real-time PCR

Quantitative real time PCR was performed using iQ™ SYBR Green Supermix (Bio-Rad, Singapore) on a BIO-RAD CFX96 Real-Time System under the following conditions: 3 min at 95 °C, followed by 45 cycles of 15 s at 94 °C, 15 s at 55 °C and 30 s at 72 °C. Different genes including complement component C1R, C1S, C1S-like, C2, C3, C4, C5, C7, C8A and C9 were used for validation. An additional file shows the primer sequences used in this study (Additional file 1). The relative expression levels of the selected genes were normalized to β-Actin and calculated using \( 2^{-\Delta\Delta Ct} \) method.

**Results**

**De novo sequencing and assembly**

Two sequencing libraries were prepared from spleen samples obtained from control (GS_CS) and GSIV-infected (GS_TS) Chinese giant salamanders that were sequenced using an Illumina Hiseq 2000. In total, 122.48 million raw reads were generated from GS_CS and 154.75 million for GS_TS. The data was refined by discarding low-quality reads that contained unknown bases or whose length was lower than 20 nt after removal of the adaptors and low-quality bases. The resulting high-quality reads numbered 113.45 million and 143.78 million for the GS_CS and GS_TS samples, respectively. The total length of these reads was 9.6 \( \times \) 10^9 and 11.97 \( \times \) 10^9 bp for GS_CS and GS_TS samples, respectively and the Q20 percentage (the percentage of sequences with a sequencing error rate lower than 1%) was over 98% for both samples (Table 1). All high-quality reads were deposited in the National Central for Biotechnology Information (NCBI) and can be accessed under the accession number SRP047798.

De novo assembly was performed using Trinity that resulted in 80 367 genes and 123 440 isogenes. The total length was 182 916 518 bp, with an average length of 1481 bp (Table 2). Each isogene was longer than 351 bp, and 71 295 (57.76%) of the isogenes were 350–1000 bp. Additionally, 27 826 (22.5%) of the isogenes were longer than 2000 bp. The size distribution of isogenes is shown in Figure 1A.

### Table 1 Summary of sequencing results.

|                | GSIV-infected | Control |
|----------------|---------------|---------|
| Total raw reads | 154 748 118   | 122 483 450 |
| Total clean reads | 143 776 686   | 113 450 692  |
| Total clean nucleotides (bp) | 11 967 607 097 | 9 600 577 224 |
| Q20 percentage  | 98.44%        | 98.43%  |

### Table 2 Summary of assembly results.

| Type            | Number |
|-----------------|--------|
| Total gene      | 80 367 |
| Total isogenes  | 123 440 |
| Total residues  | 182 916 518 |
| Average length  | 1481.83 |
| Largest isogene | 18 965 |
| Smallest isogene| 351    |

**Functional annotation and classification**

All assembled high-quality isogenes had their ORFs predicted using Trinity. A total of 58 979 (47.8%) isogenes contained an ORF. The predicted protein sequences were blasted against NR (non-redundant protein sequences in NCBI), String and Gene databases using BLASTP with a cut-off E value of \( 10^{-5} \). There were 38 106 (64.6%) isogenes with homologous sequences in at least one of the databases. Among them, 38 106 (64.6%), 15 579 (26.4), and 20 352 (34.5%) isogenes were found in NR, String and Gene databases, respectively. A total of 9629 (16.3%)
isogenes were found in all three databases, while 20 873 (35.4%) isogenes were not identified (Figure 1B).

The ORFs of 64 461 (52.2%) isogenes could not be predicted, and the DNA sequences of 64 461 isogenes were blasted against the NR, String and Gene databases using BLASTX with a cut-off E value of $10^{-5}$. There were 10 138 (15.7%) isogenes with homologous sequences in at least one of above databases. Among them, 10 138 (15.7%), 959 (1.5%), and 3351 (5.2%) isogenes were found in the NR, String and Gene databases, respectively. A total of 516 (0.8%) isogenes were found in all three databases, while 54 323 (84.3%) isogenes were not identified (Figure 1C).

Based on NR annotations, the Gene Ontology (GO) classification system was used to classify the possible functions of the isogenes. A total of 31 356 (25.4%) isogenes were successfully assigned to at least one GO term annotation and were classified into three main categories including biological process, cellular component and molecular function (Figure 2A). For biological process, the top six largest categories were cellular process (23 645), single-organism process (19 125), metabolic process (17 849), biological regulation (15 665), regulation of biological process (14 954) and response to stimulus (11 342). For the cellular component category, the top three largest categories were cell (22 548), cell part (22 544) and organelle (16 663). Only a few isogenes belonged to the virion (29), and virion part (24).
Figure 2. GO functional annotation and COG function classification. A. GO functional annotation. Isogenes with the best BLAST hits were aligned to GO database. All 31,356 isogenes were assigned to at least one GO term and were grouped into three main GO categories and 61 subcategories. Right y-axis represents number of isogenes in a category. Left y-axis indicates percentage of a specific category of isogenes in each main category. B. COG classification of putative proteins. All 16,538 putative proteins showing significant homology to those in the COG database were functionally classified into 25 families. The y-axis indicates number of isogenes in a specific functional cluster.
sub-categories. Interestingly, for the molecular function category, 20,914 and 10,902 isogenes were classified into the sub-categories “binding” and “catalytic activity”, respectively.

Isogenes sequences in our transcriptome library were also analyzed by the Clusters of Orthologous Groups of proteins (COG). Out of 38,106 NR hits, 7,719 (20.3%) of the sequences showed a COG classification (Figure 2B). The largest category was “general function prediction only” (2,875 of 7,719 isogenes or 37.2%), followed by “signal transduction mechanisms” (1,224 isogenes, 15.9%), “transcription” (1,126 isogenes, 14.6%), “replication, recombination and repair” (1,105 isogenes, 14.3%) and “post-translational modification, protein turnover, chaperones” (706 isogenes, 9.1%). The sub-categories “extracellular structures” (0, 0), “nuclear structure” (4, 0.05%) and “cell motility” (8, 0.1%) had the fewest related genes. Additionally, 310 (4.02%) isogenes were annotated as “function unknown”.

The 38,106 annotated sequences were mapped to the reference canonical pathways in the Kyoto Encyclopedia of Genes and Genomes (KEGG). Among those, 23,712 isogenes were assigned to 317 KEGG pathways (Additional file 2). Of the 23,712 isogenes, 2,195 (9.26%) were related to metabolic pathways, 784 (3.31%) to pathways in cancer, 750 (3.16%) to PISK-Akt signaling pathway, 643 (2.71%) to MAPK signaling pathway, 614 (2.59%) to neuroactive ligand-receptor interaction, and 604 (2.55%) to HTLV-I infection.

**Differential expression analysis**

To identify differential expression changes between GS_CS and GS_TS samples, FPKM method was used to calculate the expression levels of genes and isogenes. The results showed that 293 genes were down-regulated and 220 genes were up-regulated with an FDR < 0.05 and ratios larger than 2 (Figures 3A and B). Among them, 102 down-regulated genes were detected only in the GS_CS samples (Additional file 3A), and 67 up-regulated genes specific for the GS_TS samples (Additional file 3B). For the isogenes, 2,888 down-regulated and 3,588 up-regulated isogenes were identified (Figures 3C and D).

---

**Figure 3** Identification of differentially expressed genes/isogenes between infected and uninfected groups. **A** The expression levels of differentially expressed genes. **B** The distribution of differentially expressed genes. **C** The expression levels of differentially expressed isogenes. **D** The distribution of differentially expressed isogenes. Differentially expressed genes or isogenes were determined using a threshold of FDR ≤ 0.001 and |log$_{2}$Ratio| > 1. Red spots represent differentially expressed genes or isogenes. Black spots represent genes or isogenes that didn’t show obvious changes in the GSV-infected Chinese giant salamander.
Enrichment analysis was conducted to help clarify the biological functions of all differentially expressed isogenes (DEGs) that were identified. All DEGs were mapped to each term of the GO database, and the GO terms with a corrected P value ≤0.05 were defined as significantly enriched in DEGs. The results indicated that 2493 DEGs were enriched in 433 GO terms (Additional file 4). Among these GO terms, “immune system process” (172 DEGs), “regulation of immune system process” (123 DEGs), “response to stimulus” (962 DEGs), “response to DNA damage stimulus” (78 DEGs), “lymphocyte activation” (45 DEGs), “T cell activation” (32 DEGs), “leukocyte migration” (30 DEGs) and “chemotaxis” (81 DEGs) were significantly enriched among DEGs compared with the whole transcriptome background.

We also performed an enrichment analysis of the KEGG pathways and diseases. These DEGs were significantly enriched in 7 pathways including “complement and coagulation cascades”, “hematopoietic cell lineage”, “Staphylococcus aureus infection”, “tight junction”, “viral myocarditis”, “transcriptional misregulation in cancer” and “allograft rejection” (Additional file 5). The most enriched pathway was “complement and coagulation cascades” (66 DEGs), with 7 DEGs up-regulated and 59 DEGs down-regulated (Figure 4). The significantly enriched disease terms included inherited thrombophilia (11 DEGs), immune system diseases (94 DEGs), primary immunodeficiency (71 DEGs), complement regulatory protein defects (9 DEGs), afibrinogenemia (5 DEGs), congenital disorders of the DNA repair systems (15 DEGs), agammaglobulinemias (9 DEGs), disorders of nucleotide excision repair (13 DEGs), congenital disorders of metabolism (112 DEGs), cockayne syndrome (7 DEGs), H00821 (15 DEGs), familial combined hyperlipidemia (5 DEGs), H00978 (8 DEGs) and epidermolysis bullosa, simplex (7 DEGs) (Additional file 5). Among those DEGs involved in immune system diseases, 41 DEGs were up-regulated and 53 DEGs were down-regulated (Figure 5A). Those DEGs associated with inherited thrombophilia, complement regulatory protein defects, afibrinogenemia, H00821, epidermolysis bullosa, and simplex diseases were strongly suppressed in the spleen of GSIV-infected Chinese giant salamanders (Figure 5).

SSR and SNP discovery

Simple sequence repeats (SSRs) have been shown to be an efficient tool for performing quantitative trait loci (QTL) analysis and constructing genetic linkage(s) due to their high diversity and abundance. In the present study, 30 678 SSRs were obtained from the transcriptomic information from the Chinese giant salamander. Among them, the most frequent repeat motifs were mononucleotide SSR motifs (76.2%), followed by trinucleotide (11.95%), dinucleotide (10.06%), tetranucleotide (1.6%), pentanucleotide (0.14%) and hexanucleotide (0.02%) (Figure 6A). Based on the distribution of SSR motifs, (A)^n, (G)^n and (T)^n (n ≥ 10) were the three predominant types among mononucleotide SSR motifs, with frequencies of 30.93, 30.14 and 27.77%, respectively (Figure 6B). In the six types of dinucleotide repeat motifs, GT (27.93%) was the most common motif, followed by AT (26.34%) and AC (22.2%) (Figure 6C). Among 20 types of trinucleotide repeats, AGC (13.91%), GCT (13.64%) and AGG (11.86%) were the three predominant types (Figure 6D). Tetranucleotide and pentanucleotide SSR motifs contained 35 and 23 types of repeats, respectively (Figures 6E and F).

SNPs were identified from alignments of multiple sequences used for contig assembly. A total of 26 355 and 36 070 SNPs were obtained from the GS_CS and GS_TS samples, respectively. In the GS_CS samples, 16 524 were putative transitions and 9831 were putative transversions (Figure 7A). In the GS_TS samples, the numbers of various SNP types were higher than that in GS_CS samples (Figures 7A and B), with 23 030 SNPs for transitions and 13 040 SNPs for transversions (Figure 7B). No insertion–deletion polymorphisms (indels) were found in either the GS_CS or GS_TS samples.

DEGs, SSRs and SNPs analysis related to disorders of nucleotide excision repair

Among the sequencing libraries, we identified 75 sequences related to nucleotide excision repair. The members of the DNA repair gene family including ERCC1-6, ERCC8, XPA, POLH, BLM, TFIIH1-4, RBX1, CUL4, DDB1, DDB2, XPC, HR23, CETN2, CDK7, MNA1, CCNH, TTDA, RPA, PCNA, RFC and LIG1 were all expressed in the spleen of the Chinese giant salamander. However, 10 genes including BLM, DDB2, ERCC4, ERCC5, ERCC6, ERCC8, POLH, RPA1, TFIIH3 and XPA were DEGs, with 11 isogenes up-regulated and 6 isogenes down-regulated (Table 3).

Further analysis of SSRs and SNPs involved in disorders of nucleotide excision repair indicated a total of 12 genes that were related to disorders of nucleotide excision repair that contained 21 SSRs motifs (Table 4). Among them, DEGs including BLM, ERCC6 and ERCC8 contained 1 mononucleotide SRR and 4 trinucleotide SRR motifs. Thirteen genes including ERCC1, ERCC2, CCNH, CUL4, DDB1, LIG1, PCNA, RBX1, RPA, RFC, TFIIH2, TTDA and XPA contained 76 SNP sites (Table 5). Up to 15 SNPs existed in the isogene of PCNA (comp90786_c1_seq5). Among 76 SNPs, 33 (43.42%) and 29 (38.16%) of the SNPs were specific for the uninfected and infected Chinese giant salamanders, respectively. There were 14 SNPs, that were identical for the GS_CS and GS_TS samples but different from the reference sequences.
The gene cluster for "complement and coagulation cascades" pathway. 66 differentially expressed genes (DEGs) were enriched in the pathway, with 7 DEGs up-regulated and 59 DEGs down-regulated.

| CS | TS | Transcript ID | Gene Symbol |
|----|----|---------------|-------------|
|    |    | comp95383_c0_seq1 | KNG         |
|    |    | comp94398_c2_seq1 | A2M         |
|    |    | comp611755_c0_seq1 | A2M         |
|    |    | comp94469_c0_seq6 | A2M         |
|    |    | comp94905_c0_seq1 | A2M         |
|    |    | comp94905_c0_seq3 | A2M         |
|    |    | comp94905_c0_seq2 | A2M         |
|    |    | comp94905_c0_seq4 | A2M         |
|    |    | comp94469_c0_seq5 | A2M         |
|    |    | comp81391_c0_seq2 | C1R         |
|    |    | comp89723_c0_seq1 | C1S         |
|    |    | comp72777_c0_seq1 | C1S         |
|    |    | comp86649_c0_seq3 | C2          |
|    |    | comp94356_c0_seq1 | C3          |
|    |    | comp84612_c0_seq1 | C3          |
|    |    | comp90690_c0_seq1 | C4          |
|    |    | comp89005_c0_seq1 | C5          |
|    |    | comp93876_c1_seq8 | C7          |
|    |    | comp84568_c0_seq1 | C8A         |
|    |    | comp83899_c0_seq1 | C8G         |
|    |    | comp33080_c0_seq1 | C9          |
|    |    | comp85174_c0_seq1 | CD142       |
|    |    | comp94068_c1_seq6 | CD46        |
|    |    | comp94068_c1_seq3 | CD46        |
|    |    | comp92123_c0_seq22 | CR1L       |
|    |    | comp92123_c0_seq16 | C4BPA     |
|    |    | comp92123_c0_seq19 | CR2L       |
|    |    | comp92123_c0_seq12 | CR1L       |
|    |    | comp92123_c0_seq10 | CR1L       |
|    |    | comp92123_c0_seq15 | C4BPA     |
|    |    | comp92123_c0_seq1 | CR1L       |
|    |    | comp92123_c0_seq6 | CR1L       |
|    |    | comp92123_c0_seq4 | CR2L       |
|    |    | comp77916_c0_seq1 | CPB2       |
|    |    | comp93584_c0_seq1 | F10         |
|    |    | comp58404_c0_seq1 | F11         |
|    |    | comp92422_c0_seq10 | F13B      |
|    |    | comp93987_c1_seq15 | F13B      |
|    |    | comp58233_c0_seq1 | F2          |
|    |    | comp94479_c0_seq1 | F2R        |
|    |    | comp86862_c0_seq1 | F7          |
|    |    | comp98857_c0_seq1 | F9          |
|    |    | comp85051_c0_seq1 | FGA         |
|    |    | comp76358_c0_seq1 | FGB         |
|    |    | comp85137_c0_seq2 | FGG         |
|    |    | comp85137_c0_seq1 | FGG         |
|    |    | comp85137_c0_seq3 | FGG         |
|    |    | comp92421_c0_seq9 | HF1         |
|    |    | comp92421_c0_seq14 | HF1        |
|    |    | comp92421_c0_seq4 | HF1         |
|    |    | comp92421_c0_seq6 | HF1         |
|    |    | comp92421_c0_seq16 | HF1        |
|    |    | comp58072_c0_seq1 | KLKB1       |
|    |    | comp98432_c0_seq1 | KNG         |
|    |    | comp33808_c0_seq1 | MASP2       |
|    |    | comp92722_c0_seq6 | MBL         |
|    |    | comp80308_c0_seq2 | PLG         |
|    |    | comp80308_c0_seq1 | PLG         |
|    |    | comp72838_c0_seq1 | PROC        |
|    |    | comp75671_c1_seq1 | SERPINA1, AAT |
|    |    | comp95543_c0_seq1 | SERPINA1, AAT |
|    |    | comp33154_c0_seq1 | SERPINA1, AAT |
|    |    | comp75671_c0_seq1 | SERPINA1, AAT |
|    |    | comp84313_c0_seq1 | SERPINC1, AT3 |
|    |    | comp58285_c0_seq1 | SERPIND1, HCF2 |
|    |    | comp78063_c0_seq1 | VWF         |
Figure 5 Number and gene clusters of differentially expressed isogenes in KEGG diseases. A KEGG enrichment analysis showed significantly enriched disease. The y axis indicates numbers of DEGs enriched in KEGG diseases. The x axis represents disease terms. The black bar indicates down-regulated DEGs and the white bar represents up-regulated DEGs. B The gene cluster for inherited thrombophilia. C The gene cluster for complement regulatory protein defects. D The gene cluster for Epidermolysis bullosa, simplex. Those DEGs associated with inherited thrombophilia, complement regulatory protein defects and Epidermolysis bullosa, simplex diseases appeared strongly suppressed in the spleen of GSIV-infected Chinese giant salamanders.
Figure 6 Distribution of simple sequence repeats (SSR) among different nucleotide types in the spleen transcriptome. 

A Distribution of repeated nucleotide types. B Distribution of repeated mononucleotide. C Distribution of repeated dinucleotide. D Distribution of repeated trinucleotide. E Distribution of repeated tetranucleotide. F Distribution of repeated pentanucleotide.
Figure 7 Distribution of putative single nucleotide polymorphisms (SNP) in the spleen transcriptome. A Distribution of putative SNP in uninfected (CS type) Chinese giant salamanders. B Distribution of putative SNP in GSIV-infected (TS type) Chinese giant salamanders.

Table 3 DEGs related to disorders of nucleotide excision repair.

| Transcript_id      | KEGG GENE NAME | CS fpkm | TS fpkm | log2FC1 (TS/CS) | Significant |
|--------------------|----------------|--------|--------|-----------------|-------------|
| comp90201_c0_seq1  | BLM            | 1.18   | 0      | −8.73           | Yes         |
| comp90201_c0_seq2  | BLM            | 0      | 0.99   | 8.29            | Yes         |
| comp90773_c0_seq2  | DDB2           | 0      | 1.18   | 6.51            | Yes         |
| comp93741_c0_seq2  | ERCC4, XPF     | 0      | 0.25   | 6.27            | Yes         |
| comp92742_c0_seq18 | ERCC5, XPG     | 0.54   | 0      | −7.86           | Yes         |
| comp91446_c0_seq12 | ERCC6, CSB     | 0      | 0.35   | 6.94            | Yes         |
| comp91446_c0_seq15 | ERCC6, CSB     | 0.45   | 0      | −7.57           | Yes         |
| comp91446_c0_seq3  | ERCC6, CSB     | 0.57   | 0      | −7.89           | Yes         |
| comp91446_c0_seq4  | ERCC6, CSB     | 0      | 0.61   | 7.82            | Yes         |
| comp91446_c0_seq8  | ERCC6, CSB     | 0      | 0.63   | 7.85            | Yes         |
| comp81303_c0_seq1  | ERCC8, CSA     | 0      | 0.59   | 6.14            | Yes         |
| comp81303_c0_seq2  | ERCC8, CSA     | 0.82   | 0      | −6.78           | Yes         |
| comp92141_c0_seq1  | POLH           | 0      | 0.18   | 5.53            | Yes         |
| comp92141_c0_seq3  | POLH           | 0      | 0.19   | 5.53            | Yes         |
| comp88203_c0_seq1  | RPA1, POLR1A   | 0      | 3.78   | 7.76            | Yes         |
| comp96879_c0_seq2  | TFIIH3, TFB4   | 1.61   | 0      | −7.53           | Yes         |
| comp91081_c0_seq4  | XPA            | 0      | 1.46   | 7.99            | Yes         |

DEGs was filtered using threshold of false discovery rate (FDR) ≤0.05 and absolute value of log2Ratio ≥1.
DEGs, SSRs and SNPs analysis related to antiviral activity

RIG-I-like receptor and Toll-like receptor signaling pathways are two pivotal pathways involved in antiviral immune responses. In the Chinese giant salamander, 143 isogenes involved in the RIG-I-like receptor signaling pathway and 201 isogenes involved in the Toll-like receptor signaling pathway were identified. However, only 20 and 18 isogenes were DEGs for the RIG-I-like receptor and Toll-like receptor signaling pathways, respectively (Additional file 6). Interestingly, the expression of antiviral pattern recognition receptors such as RIG-I, MDA5, LGP2, TLR3, TLR8, TLR7 and TLR9 did not change in response to the GSIV infection. The expression of NLRX1 was significantly increased, whereas TRIM25 decreased (Additional file 6).

For the RIG-I-like receptor signaling pathway, 21 genes contained 38 SSR motifs. Of these, IL8 contained the most SSRs (6 motifs), followed by IRF7 (5 motifs) and RIG-I (4 motifs). For the Toll-like receptor signaling pathway, 21 genes that included AKT, CCL3, ERK1, FADD, IKBKE, IKBKG, IL8, IRF7, JUN, MAP2K7, MAP3K7IP2, MYD88, NFKB1, NFKB1A, P38, PIK3C, PIK3R, TBK1, TBK1, TLR1 and TLR5 contained 59 SSR motifs. Among them, AKT contained the most mononucleotide SSRs, and PIK3C had the most trinucleotide SSRs (Additional file 7).

For SNPs analysis, 25 genes involved in the RIG-I-like receptor signaling pathway contained 272 motifs. Up to 75 and 52 SNPs existed in the genes of IKBKG and OTUD5, respectively. Among 272 SNPs, 207 (76.1%) SNPs were specific for the GSIV-infected Chinese giant salamanders. The antiviral PRRs MDA5, LGP2 and RIG-I also contained SNP motifs. For the Toll-like receptor signaling pathway, 41 genes contained 445 SNPs that consisted of 141 SNPs (31.69%) specific for the GS_CS samples and 228 SNPs (51.24%) specific for the GS_TS samples, and 73 common SNPs (16.4%) for both the GS_CS and GS_TS samples (Additional file 8). The genes involved in both the RIG-I-like receptor and Toll-like receptor signaling pathways, which included IRF3, IRF7, NFKBIA, FADD, JNK, RELA, IL12B, IL8, CASP8, P38, IKBKA, IKBKE and IKBKG contained a total of 141 SNPs (Additional files 8 and 9).

Experimental validation

Since the most enriched pathway was the “complement and coagulation cascades”, 10 genes belonging to the complement components were selected and used for qPCR validation. PCR amplification showed that all qPCR primers produced single fragments of the expected lengths (141–354 bp). Except for complement components C1R (comp81391_c0_seq2), C1S (comp72777_c0_seq1) and C7 (comp93878_c1_seq8), the expression

### Table 4 SSRs related to disorders of nucleotide excision repair.

| Transcript_id | KEGG GENE NAME | Start BP | Repeat | End BP | Type       |
|---------------|----------------|----------|--------|--------|------------|
| comp90201_c0_seq1 | BLM          | 5420     | (GCT)^4 | 5432   | Trinucleotide |
| comp90201_c0_seq1 | BLM          | 3386     | (ATC)^4 | 3398   | Trinucleotide |
| comp57681_c0_seq1 | CDK7        | 43       | (AT)^6  | 55     | Dinucleotide |
| comp71102_c0_seq1 | CUL4        | 944      | (A)^10  | 954    | Mononucleotide |
| comp107286_c0_seq1 | ERCC3, XPB  | 2478     | (G)^10  | 2488   | Mononucleotide |
| comp107286_c0_seq1 | ERCC3, XPB  | 2204     | (AGC)^4 | 2216   | Trinucleotide |
| comp93741_c0_seq1 | ERCC4, XPF  | 3807     | (T)^10  | 3817   | Mononucleotide |
| comp91446_c0_seq12 | ERCC6, CSB  | 3738     | (AGG)^6 | 3756   | Trinucleotide |
| comp91446_c0_seq15 | ERCC6, CSB  | 3727     | (AGG)^6 | 3745   | Trinucleotide |
| comp81303_c0_seq1 | ERCC8, CSA  | 66       | (T)^14  | 80     | Mononucleotide |
| comp33205_c0_seq1 | RAD23, HR23 | 1046     | (AGG)^4 | 1058   | Trinucleotide |
| comp33042_c0_seq1 | RBX1, ROC1  | 1        | (T)^11  | 12     | Mononucleotide |
| comp89984_c0_seq1 | rpa          | 577      | (G)^10  | 587    | Mononucleotide |
| comp33075_c0_seq1 | RPA2        | 191      | (GGT)^4 | 203    | Trinucleotide |
| comp57325_c0_seq1 | RPA12       | 1        | (G)^21  | 22     | Mononucleotide |
| comp72565_c0_seq1 | RPA3        | 59       | (A)^12  | 71     | Mononucleotide |
| comp56791_c0_seq1 | RFC          | 1        | (T)^13  | 14     | Mononucleotide |
| comp36422_c0_seq1 | RFC          | 1        | (T)^11  | 12     | Mononucleotide |
| comp37382_c0_seq1 | RFC          | 192      | (GGT)^4 | 204    | Trinucleotide |
| comp53798_c0_seq1 | TFIIH1, TFBI| 2301     | (AC)^7  | 2315   | Dinucleotide |
| comp82958_c1_seq1 | TFIIH2, SSL1| 3        | (A)^19  | 22     | Mononucleotide |
Table 5 List of SNPs related to disorders of nucleotide excision repair.

| Transcript_id     | KEGG GENE NAME | Site  | Ref | CS_type | TS_type |
|-------------------|----------------|-------|-----|---------|---------|
| comp74141_c0_seq1 | ERCC1          | 2223  | G   | –       | A       |
| comp74141_c0_seq2 |                | 2070  | G   | –       | A       |
| comp74141_c0_seq2 |                | 847   | G   | T       | T       |
| comp88233_c0_seq1 | ERCC2          | 128   | G   | A       | –       |
| comp88233_c0_seq1 |                | 2140  | T   | –       | C       |
| comp88233_c0_seq1 |                | 2450  | G   | –       | T       |
| comp108677_c0_seq1 | CCNH           | 190   | C   | T       | –       |
| comp61358_c0_seq1 | Cul4A          | 1710  | A   | C       | –       |
| comp61358_c0_seq1 |                | 871   | C   | –       | A       |
| comp71102_c0_seq1 | Cul4B          | 1738  | A   | –       | G       |
| comp71102_c0_seq1 |                | 811   | A   | –       | G       |
| comp32714_c0_seq1 | DDB1           | 129   | G   | –       | A       |
| comp32717_c0_seq1 |                | 1429  | G   | –       | A       |
| comp32717_c0_seq1 |                | 433   | T   | C       | –       |
| comp32741_c0_seq1 |                | 1157  | G   | C       | C       |
| comp80349_c0_seq1 | LIG1           | 3623  | C   | –       | T       |
| comp90786_c1_seq5 | PCNA           | 105   | T   | C       | C       |
| comp90786_c1_seq5 |                | 13    | G   | –       | A       |
| comp90786_c1_seq5 |                | 1314  | G   | –       | A       |
| comp90786_c1_seq5 |                | 1683  | A   | G       | –       |
| comp90786_c1_seq5 |                | 1934  | G   | A       | A       |
| comp90786_c1_seq5 |                | 2038  | A   | G       | –       |
| comp90786_c1_seq5 |                | 2068  | A   | G       | G       |
| comp90786_c1_seq5 |                | 22    | A   | G       | –       |
| comp90786_c1_seq5 |                | 2395  | A   | G       | –       |
| comp90786_c1_seq5 |                | 2465  | G   | –       | A       |
| comp90786_c1_seq5 |                | 2487  | G   | A       | A       |
| comp90786_c1_seq5 |                | 2501  | A   | G       | –       |
| comp90786_c1_seq5 |                | 2543  | G   | –       | A       |
| comp90786_c1_seq5 |                | 2562  | A   | G       | –       |
| comp90786_c1_seq5 |                | 93    | G   | C       | –       |
| comp90786_c1_seq7 | PCNA           | 105   | T   | C       | C       |
| comp90786_c1_seq7 |                | 1240  | T   | C       | C       |
| comp90786_c1_seq7 |                | 1244  | C   | A       | A       |
| comp90786_c1_seq7 |                | 13    | G   | –       | A       |
| comp90786_c1_seq7 |                | 1818  | G   | A       | A       |
| comp90786_c1_seq7 |                | 2077  | A   | C       | C       |
| comp90786_c1_seq7 |                | 22    | A   | G       | –       |
| comp90786_c1_seq7 |                | 2401  | A   | –       | G       |
| comp90786_c1_seq7 |                | 2615  | C   | –       | G       |
| comp90786_c1_seq7 |                | 93    | G   | C       | –       |
| comp33042_c0_seq1 | RBX1           | 89    | G   | T       | –       |
| comp89984_c0_seq1 | RPA            | 1679  | C   | –       | T       |
| comp89984_c0_seq1 |                | 1829  | G   | A       | –       |
| comp89984_c0_seq1 |                | 2071  | G   | –       | A       |
| comp89984_c0_seq1 |                | 2600  | C   | G       | –       |
| comp89984_c0_seq1 |                | 2645  | G   | T       | –       |
| comp89984_c0_seq1 |                | 3312  | A   | G       | –       |
| comp89984_c0_seq1 |                | 799   | C   | T       | –       |
| comp89984_c0_seq1 |                | 815   | G   | T       | –       |
of other 7 complement components including C1S-like, C2, C3, C4, C5, C8A and C9 was in agreement with their transcript abundance changes determined by RNA-seq (Figure 8).

Discussion

Transcriptome sequencing of spleen samples from uninfected and infected Chinese giant salamanders

The complete genome of GSIV has been sequenced [12]. However, the available molecular information is limited for the Chinese giant salamander, and not sufficient for the investigation of the mechanisms of immune reactivity against pathogen infection. In recent years, a number of reports reveal that the transcriptome sequencing of cDNA is an efficient method for generating large sequences that represent expressed genes [33]. Given that the spleen is one of the most important immune organs and is also the main target organ for the iridovirus GSIV [10, 22, 34, 35], the transcriptome sequencing of spleen samples from uninfected and infected Chinese giant salamanders was expected to provide abundant ESTs for amphibian immune genes and contribute to the understanding of GSIV-host interactions. In the study reported herein, using the recently developed Solexa sequencing technology and Trinity RNA-Seq assembly, the results of transcriptome sequencing are reported for the Chinese giant salamanders.

TLR, NLR and the RLR systems that are involved in immune responses

To begin this line of research, we first checked a set of sequences that encode components of TLR, NLR and the RLR systems that are involved in immune responses. In the spleen transcriptome of the Chinese giant salamanders, a total of 201, 206 and 143 isogenes were identified to be involved in the TLR, NLR and RLR pathways, respectively.

Table 5 continued

| Transcript_id | KEGG GENE NAME | Site | Ref | CS_type | TS_type |
|---------------|----------------|------|-----|---------|---------|
| comp33075_c0_seq1 | RPA2 | 100 | T | C | C |
| comp57325_c0_seq1 | RPA12 | 385 | G | – | C |
| comp59041_c0_seq1 | RPA49 | 757 | G | A | – |
| comp59041_c0_seq1 | 864 | A | – | G |
| comp59044_c0_seq1 | 380 | G | – | C |
| comp72565_c0_seq1 | RPA3 | 475 | A | – | G |
| comp72565_c0_seq1 | 802 | G | A | – |
| comp8203_c0_seq2 | RPA | 4663 | G | C | – |
| comp81979_c0_seq1 | RPA43 | 1011 | C | – | T |
| comp81979_c0_seq1 | 1032 | C | T | – |
| comp81979_c0_seq1 | 1098 | G | – | A |
| comp56791_c0_seq1 | RFC1 | 3527 | T | C | C |
| comp56791_c0_seq1 | 3757 | A | – | G |
| comp36422_c0_seq1 | RFC2 | 1188 | C | – | T |
| comp85228_c0_seq1 | RFC2 | 1319 | T | G | – |
| comp85228_c0_seq1 | 1744 | A | C | – |
| comp85228_c0_seq1 | 1844 | C | T | – |
| comp36017_c0_seq1 | RFC3 | 335 | A | C | C |
| comp7382_c0_seq1 | RFC3 | 49 | C | T | – |
| comp82958_c1_seq1 | TFIIH2 | 122 | G | – | A |
| comp7951_c0_seq1 | TTDA | 513 | G | T | – |
| comp91081_c0_seq1 | XPA | 1705 | T | C | – |
| comp91081_c0_seq3 | 1633 | T | G | – |
| comp91081_c0_seq3 | 2827 | A | T | – |
| comp91081_c0_seq4 | 2669 | A | T | – |
| comp91081_c0_seq4 | 2893 | C | A | A |

*The identical bases with reference sequences were indicated with “–”.*
conservation and divergence of the TLR, NLR and RLR families are summarized below and in the figure legends.

**The TLR system**

The Toll-like Receptors (TLRs) are present throughout virtually the entire animal kingdom and have important functions in initiating inflammatory responses and shaping adaptive immunity [36]. A typical TLR is a type I transmembrane protein with many extracellular leucine-rich repeat (LRR) motifs for ligand recognition, and a cytoplasmic TIR domain for signal transduction. In invertebrates, only one or two TLR genes exist in sea squirt *Ciona intestinalis* and the nematode *Caenorhabditis elegans* [37, 38], however the sea urchin *Strongylocentrotus purpuratus* and amphioxus *Branchiostoma lanceolatum* possess a large number of TLRs [39, 40]. In mammals, the human genome contains 10 functional TLRs whereas the mouse genome contains 12 TLRs, with TLR10 being a pseudogene, and TLR11, TLR12 and TLR13 being mouse-specific genes [41]. Most fish species possess a higher number of TLR genes than mammalian species due to the presence of duplicated TLRs and fish-specific TLRs [42, 43].

Six major TLR families were identified in all vertebrate taxa designated as TLR1, TLR3, TLR4, TLR5, TLR7 and TLR11 [44]. In the Chinese giant salamanders, the members of TLR3, TLR5 and TLR7 have clear orthologs with fish and mammals that was supported by the phylogenetic analysis (Figure 9A). For TLR1 family, clear orthologous relationships were found among fish, amphibians and mammals for TLR2 gene, and two sequences were found to group with mammalian TLR10, TLR1 and TLR6. Interestingly, the ortholog of the mouse-specific TLR13 and fish-specific TLR21 and 22 appear to exist in the Chinese giant salamander. Similar to fish, the Chinese giant salamander has duplicated copies of some TLRs, such as TLR2, TLR5 and TLR21. These results suggest that amphibians may have a greater TLR repertoire than fish or mammals as a result of living in both aquatic and terrestrial environments.

**The NLR system**

A large family of cytoplasmic NACHT-LRR receptors (NLRs), characterized by the presence of a nucleotide-binding protein (NACHT domain), have important functions in apoptosis, inflammation and innate immune signaling [45–48]. There are 20–30 NLR genes in vertebrates, which are divided into three distinct subfamilies with regard to their phylogenetic relationships including the NODs (NOD1-2, NOD3/NLRC3, NOD4/NLRC5, NOD5/NLRX1, CIITA), the NLRPs (NLRP1–14, also called NALPs) and the IPAF subfamily, consisting of IPAF (NLRC4) and NAIP [49]. In the Chinese giant salamander, the canonical NOD proteins (NOD1, NOD2, NOD3, NOD5, CIITA), IPAF, NAIP, as well as APAF1 (apoptotic protease activating factor 1) exhibit clear orthologous relationships (Figure 9B). However, no homologous sequences were found for NOD4 in the spleen transcriptome of the Chinese giant salamander.
Figure 9  Phylogenetic tree of Toll-like receptors (A) and NACHT-domain family (B). Phylogenetic relationships were based on amino acid alignments. Bootstrap values based on 10,000 replicates are indicated on each branch. The evolutionary history was inferred using the neighbor-joining method. All positions containing gaps and missing data were eliminated from the dataset (pairwise deletion). Accession numbers of sequences used to build the tree are presented in Additional file 11.
The expansion of NLR-encoding genes has been described in the sea urchin, amphioxus and zebrafish [39, 40, 50, 51]. In their genome, at least 92 (amphioxus) and 200 (sea urchin and zebrafish) NLR genes were predicted. The large majority of the sea urchin NLR proteins consist of a central NACHT domain, an N-terminal DEATH domain and C-terminal LRRs [39]. However, the large groups of fish-specific NLR proteins do not contain any the amino-terminal effector domains [51]. Additionally, NLRP proteins with a PYD domain in the N-terminal region were not found in the sea urchin, amphioxus and zebrafish [39, 40, 51]. Unlike NLRs in the species mentioned above, the expansion of NLR proteins were not found in the spleen transcriptome of the Chinese giant salamander; however many annotated sequences grouped with the mammalian NLRPs in the phylogenetic tree (Figure 9B).

In the Chinese giant salamander, many NLRs such as APAF1, IPAF, NOD2, NOD3, CIITA and NALP3/NALP12 existed as multiple isoforms. Our previous studies have shown that the isoforms of immune genes regulated positively or negatively antibacterial and antiviral immunity [52–54]. The multiple isoforms of NLR genes in the present study suggested that the complexity and diversity of the innate immunity may be achieved in Chinese giant salamander through the use of alternative splicing or gene duplication.

The RLR system

Besides NLRs, the RIG-I-like receptors (RLRs) represent another crucial family of intracellular pattern recognition receptors, which use C-terminal RNA helicases to recognize viral RNA and N-terminal CARD domains for signaling [55]. The RLR family members include retinoic acid inducible gene-1 (RIG-I), melanoma differentiation gene-5 (MDA5) and laboratory of genetics and physiology-2 (LGP2), which have been identified in teleost species [52, 56–59]. These three RLRs were present in the Chinese giant salamander. As for adapters, the CARD-containing IPS-1 gene wasn’t identified in the spleen transcriptome of the Chinese giant salamander (Additional file 10A).

Comparative transcriptome analysis revealed the immune energy of TLR, NLR and the RLR systems in the spleen

Iridoviruses are large double-stranded DNA (dsDNA) viruses that can infect invertebrates and poikilothermic vertebrates, including insects, fish, amphibians and reptiles. A real-time polymerase chain reaction (PCR) assay for a marine fish iridovirus showed that the spleen and kidney contained the largest number of viral particles while no viral DNA was detected in the muscle tissue [34]. A comparison of the genes expressed during a red seabream iridovirus (RSIV) infection in the spleen and kidney suggested that RSIV preferentially targets the spleen [36]. In the Chinese giant salamander, our previous study showed tissue necrosis and the existence of GSIV viral particles in the spleen, liver and kidney [10]. Many studies have shown that the spleen and kidney are the target organs of iridovirus infections.

Using 454 pyrosequencing, 755 up-regulated genes and 695 down-regulated genes were identified in the two spleen-complementary DNA libraries, that were constructed from Singapore grouper iridovirus (SGIV) infected and control orange-spotted grouper [22]. Although 80 367 genes were identified in the spleen transcriptome of the Chinese giant salamander using Solexa sequencing technology, comparative transcriptome analysis indicated that only 293 genes were down-regulated and 220 genes were up-regulated in response to the GSIV infection. In addition, a large number of genes were involved in TLR (72 hits), NLR (63 hits) and RLR (44 hits) pathways, however only 11 (TLR), 12 (NLR) and 10 (RLR) genes in these pathways showed significant changes in their transcripts after GSIV infection. The non-significant changes for the majority of the genes including PRRs and the non-significant enrich for these pathways suggested that TLR, NLR and RLR systems in the spleen of Chinese giant salamanders were immune anergic during GSIV infection.

Comparative transcriptome analysis revealed the primary immunodeficiency of the complement system in the spleen

The vertebrate complement system is a humoral and proteolytic system that is composed of approximately 40 soluble and membrane-bound proteins. It is an integral part of the innate immune system protecting the host against invasion and proliferation of various pathogens. The mammalian complement system has three different activation pathways which include classical, alternative and lectin. Activation of the classical pathway is triggered by the binding of C1q proteins to immune complexes or aggregates containing IgG or IgM [60]. The lectin pathway parallels the classical pathway, the difference being at the initial step of target recognition and subsequent activation [61]. Activation of the lectin pathway occurs through the binding of the mannose-binding lectins (MBL) to their target, which results in the activation of the MBL-associated serine proteases (MASPs) [62]. Different from the classical and lectin pathways, the alternative pathway of complement activation is triggered spontaneously, and primarily makes use of the recognition of host-associated molecular patterns (HAMPs), not pathogen-associated molecular patterns (PAMPs) [63].

In the study reported herein, the majority of genes involved in the complement and coagulation cascades (53 of 59 total genes or 90%) were identified, and the
alternative, lectin and classical pathways appeared conserved in the Chinese giant salamander (Additional file 10B). The KEGG enrichment analysis indicated that the most enriched pathway is the "complement and coagulation cascades" and significantly enriched diseases include "primary immunodeficiency" and "complement regulatory protein defects". In mammals, primary immunodeficiencies (PIDs) are severe defects in the capacity of the host to mount a proper immune response and are characterized by an increased susceptibility to infections. Common PIDs include disorders of humoral immunity, T-cell defects, combined B- and T-cell defects, phagocytic disorders, and complement deficiencies [64].

In this study, the different genes that were expressed in the “primary immunodeficiency” were mainly the components of complement system, with down-regulated expression of C1s-like, C2, C3, C4, c8g.1, C9, complement component (3b/4b) receptor 1-like, complement receptor type 2-like, complement factor H and hemolytic complement. The data from the qRT-PCR analysis also validated the down-regulated expression of the complement components C1s-like, C2, C3, C4, C5, C8A and C9 in response to the GSIV infection, however the weak up-regulated expression of C1r, C1s and C7 in the transcriptome analysis could not be validated by qRT-PCR. The results supported that GSIV infection can trigger primary immunodeficiency of the complement system in the spleen of the Chinese giant salamander.

Additional files

Additional file 1. Oligonucleotide primers used for expression analysis. The table listed the sequences of all primers used in Quantitative real-time PCR for gene expression validation.

Additional file 2. KEGG summary of Chinese giant salamander isoforms. A total of 23 712 isoforms were assigned to 317 KEGG pathways. For each pathway, the pathway definition, number of sequences and isoform ID were listed in the table.

Additional file 3. Differentially expressed genes specific for uninfected Chinese giant salamander (A) and GSIV-infected Chinese giant salamander (B). The table provided a comprehensive list of differentially expressed genes specific for uninfected Chinese giant salamander (GS_CS) and GSIV-infected Chinese giant salamander (GS_TS). At estimated absolute log2-fold change of $>1$ and FDR $<0.05$. A total of 102 down-regulated genes were detected only in the GS_CS samples (A), and 67 up-regulated genes specific for the GS_TS samples (B).

Additional file 4. GO functional enrichment analysis of differentially expressed genes between uninfected and GSIV-infected Chinese giant salamander. This additional file showed that 2493 differentially expressed genes (DEGs) were enriched in 433 GO terms. The GO terms showed significant enrichment of "immune system process" and "response to stimulus" in addition to the other cellular processes. The GO terms with a corrected $P$ value $<0.05$ were defined as significantly enriched in DEGs.

Additional file 5. Enrichment analysis of KEGG PATHWAY and KEGG DISEASE. This additional file showed that differentially expressed genes were enriched in 7 KEGG pathways and 14 KEGG diseases.

Additional file 6. DEGs related to antiviral signaling pathway. This file showed that 20 and 18 isogenes were differentially expressed genes for the RIG-I-like receptor and Toll-like receptor signaling pathways, respectively.

Additional file 7. SSRs related to antiviral signaling pathway. The file provided a list of SSRs related to the RIG-I-like receptor and the Toll-like receptor signaling pathways. The SSRs including repeat, type, start and end positions were listed in the table.

Additional file 8. SNPs analysis related to antiviral signaling pathway. This additional file provided a detail of SNPs related to RIG-I-like receptor and the Toll-like receptor signaling pathways. The number of SNPs, which appeared in GSIV-infected (TS), uninfected (CS) and both of TS and CS samples, were also listed.

Additional file 9. A list of SNPs related to the RIG-I-like receptor signaling pathway (A) and the Toll-like receptor signaling pathway (B). 272 and 442 SNPs related to the RIG-I-like receptor and the Toll-like receptor signaling pathways, respectively, were listed in this additional file. The identical bases with reference sequences were indicated with "-".

Additional file 10. ESTs in the spleen library of Chinese giant salamanders. The majority of genes involved in the complement and coagulation cascades (53 of 59 total genes or 90%) were identified, and the alternative, lectin and classical pathways appeared conserved in the Chinese giant salamanders, which hit to the RIG-I-like receptor signaling pathway (A). The majority of genes involved in the complement and coagulation cascades (53 of 59 total genes or 90%) were identified, and the alternative, lectin and classical pathways appeared conserved in the Chinese giant salamander (B).

Additional file 11. Accession numbers of sequences used to build the phylogenetic tree. A total of 78 genes, which were used to build the phylogenetic tree, and their accession numbers were listed in this additional file.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

LZ and YF designed the experiments. YF executed the experiments, submitted data and drafted the manuscript. MXC performed primer design, the bioinformatic analysis and drafted the manuscript. YWH performed the bioinformatics analysis and qRT-PCR validation. JM and LH participated in sample collection and RNA extraction. SEL and PN contributed to manuscript editing. LZ conceived the study and revised the manuscript. All authors read and approved the final manuscript.

Acknowledgements

The work was supported by Special Fund for Agro-scientific Research in the Public Interest (201203086).

Author details

1 Yangtze River Fisheries Research Institute, Chinese Academy of Fishery Sciences, Wuhan, Hubei 430023, China. 2 State Key Laboratory of Freshwater Ecology and Biotechnology, Institute of Hydrobiology, Chinese Academy of Sciences, Wuhan, Hubei 430072, China. 3 Research Division, Clear Springs Foods, Inc., PO Box 712, Buhl, ID 83316, USA. 4 Freshwater Aquaculture Collaborative Innovation Center of Hubei Province, Huazhong Agricultural University, Wuhan, Hubei 430023, China.

Received: 16 July 2015 Accepted: 26 October 2015

Published online: 20 November 2015

References

1. Vogel G (1999) Frog is a prince of a new model organism. Science 285:25
2. Gao KQ, Shubin NH (2003) Earliest known crown-group salamanders. Nature 422:424–428
3. Zhang KJ, Wang XM, Wu W, Wang ZH, Huang S (2002) Advances in conservation biology of Chinese giant salamander. Biodivers Sci 10:291–297

4. Wang XM, Zhang KJ, Wang ZH, Ding YZ, Wu W, Huang S (2004) The decline of the Chinese giant salamander Andrias davidus and implications for its conservation. ORXY 38:197–202

5. Dong W, Zhang Y, Yang C, An J, Qin J, Song F, Zeng W (2011) Iridovirus infection in Chinese giant salamanders. China. Emerg Infect Dis 17:2388–2389

6. Geng Y, Wang KY, Zhou ZY, Li CW, Wang J, He M, Yin ZQ, Lai WM (2011) First report of a Ranavirus associated with morbidity and mortality in farmed Chinese giant salamanders (Andrias davidus). J Comp Pathol 145:95–102

7. Zhou Y, Zeng LB, Meng Y, Zhou QL, Zhang H, Gao ZY, Xiao Y, Sun JB (2012) Establishment of a TaqMan real-time PCR assay for detecting the giant salamander iridovirus. J Fish Chin 36:772–778

8. Zhou ZY, Geng Y, Ren SY, Wang KY, Huang XL, Chen DF, Liu XX, Lai WM (2012) Ranavirus (family Iridoviridae) detection by polymerase chain reaction (PCR) in Chinese giant salamander (Andrias davidus). Blanchard, 1871). China. Afr J Biotechnol 11:1530–15134

9. Meng Y, Zhang H, Liang H, Zeng L, Xiao H, Xie C (2013) Development of a loop–mediated isothermal amplification assay for rapid detection of iridovirus in the Chinese giant salamander. J Virol Methods 194:211–216

10. Meng Y, Ma J, Jiang N, Zeng LB, Xiao HB (2014) Pathological and microbiological findings from mortality of the Chinese giant salamander (Andrias davidus). Arch Virol 159:1403–1412

11. Ma J, Zeng L, Zhou Y, Jiang N, Zeng H, Fan YD, Meng Y, Xu J (2014) Ultrastructural morphogenesis of an amphibian iridovirus isolated from Chinese giant salamander (Andrias davidus). J Comp Pathol 150:325–331

12. Li W, Zhang X, Weng S, Zhao G, He J, Dong C (2014) Virion-associated viral proteins of a Chinese giant salamander (Andrias davidus) iridovirus (genus Ranavirus) and functional study of the major capsid protein (MCP). Vet Microbiol 172:129–139

13. Zhu R, Chen ZY, Wang J, Yuan JD, Liao XY, Gui JF, Zeng QY (2014) Thymus cDNA library survey uncovers novel features of immune molecules in Chinese giant salamander Andrias davidus. Dev Comp Immunol 46:413–422

14. Li FG, Wang LX, Lan QJ, Yang H, Li Y, Liu XL, Yang ZX (2015) RNA-Seq analysis and gene discovery of Andrias davidus using Illumina short read sequencing. PLoS One 10:e0123730

15. Zhu R, Chen ZY, Wang J, Yuan JD, Liao XY, Gui JF, Zeng QY (2014) Extensive diversification of MHC in Chinese giant salamanders Andrias davidus (Anda-MHC) reveals novel splice variants. Dev Comp Immunol 42:311–322

16. Wang L, Yang H, Li F, Zhang Y, Yang Z, Li Y, Liu X (2013) Molecular characterization, tissue distribution and functional analysis of macrophage migration inhibitory factor protein (MIF) in Chinese giant salamanders Andrias davidus. Dev Comp Immunol 39:161–168

17. Huang LL, Fan YD, Zhou Y, Jiang N, Liu WZ, Meng Y, Zeng LB (2015) Cloning, sequence analysis and expression profiles of Toll-like receptor 7 from Chinese giant salamander Andrias davidus. Comp Biochem Phys A 184:52–57

18. Alagna F, D’Agostino N, Torchia L, Servili M, Rao R, Pietrella M, Perrotta G, Alagna F, D’Agostino N, Torchia L, Servili M, Rao R, Pietrella M, Perrotta G (2010) The developmental dynamics of the maize leaf transcriptome. Genome Res 18:1112–1126

19. Ariffin JK, Sweet MJ (2013) Differences in the repertoire, regulation and functionalities of vertebrate Toll-like receptors. J Biol Chem 284:27336–27343

20. Satake H, Sekiguchi T, Kusumoto S, Satake H (2009) Toll-like receptors of deuterostome invertebrates. Front Immunol 3:34

21. Hibino T, Loza-Coll M, Messier C, Majeske AJ, Cohen AH, Terwilliger DP, C, Chen S, Xu A (2008) Genomic analysis of the immune gene repertoire encoded in the purple sea urchin genome. Dev Biol 300:349–365

22. Fan et al. Vet Res (2015) 46:136

23. Kanneganti TD, Lamkanfi M, Núñez G (2007) Intracellular NOD-like receptors in host defense and disease. Immunity 27:549–559

24. Conesa A, Gotz S, García-Gómez JM, Terol J, Talón M, Robbins M (2005) Blast2GO: a universal tool for annotation, visualization and analysis in functional genomics research. Bioinformatics 21:3674–3676

25. Kaneshia M, Araki M, Goto S, Hattori M, Hirakawa M, Itoh M, Katayama T, Kawashima S, Okuda S, Tomikatsu T, Yamashiri Y (2008) KEGG for linking genomes to life and the environment. Nucleic Acids Res 36:D480–D484

26. Trapnell C, Williams BA, Pertea G, Mortazavi A, Kwan G, van Baren MJ, Salzberg SL, Wold BJ, Pachter L (2010) Transcript assembly and quantification using RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. Nat Biotechnol 28:511–515

27. Mortazavi A, Williams BA, McCue K, Schaeffer L, Wold B (2008) Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat Methods 5:621–628

28. Li B, Dewey CN (2011) RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. BMC Bioinformatics 12:323

29. Robinson MD, McCarthy DJ, Smyth GK (2010) edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26:139–140

30. Faircloth BC (2008) msatcommander: detection of microsatellite repeat arrays and automated, locus-specific primer design. Mol Ecol Resour 8:92–94

31. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, 1000 Genome Project Data Processing Subgroup (2009) The Sequence Alignment/Map format and SAMtools. Bioinformatics 25:2078–2079

32. Morozova O, Hirst M, Marra MA (2009) Applications of new sequencing technologies for transcriptome analysis. Annu Rev Genomics Hum Genet 10:135–151

33. Wang XW, Ao JQ, Li QG, Chen XH (2006) Quantitative detection of a marine fish iridovirus isolated from large yellow croaker, Pseudosciaenacanthus, using a molecular beacon. J Virol Methods 133:754–760

34. Dang TL, Yasuke K, Hirono I, Kondo H, Aoki T (2007) Transcriptional profile of red seabream iridovirus in a fish model as revealed by viral DNA microarrays. Virus Genes 35:449–461

35. Iwasaki A, Medzhitov R (2004) Toll-like receptor control of the adaptive immune responses. Nat Immunol 5:987–995

36. Sáská N, Ogasawara M, Sekiguchi T, Kusumoto S, Satake H (2009) Toll-like receptors of the ascidian Ciona intestinalis: prototypes with hybrid functionalities of vertebrate Toll-like receptors. J Biol Chem 284:27336–27343

37. Satake H, Sekiguchi T (2012) Toll-like receptors of deuterostome invertebrates. Front Immunol 3:34

38. Hibino T, Loza-Coll M, Messier C, Majeske AJ, Cohen AH, Tenwilliger DP, Buckley KB, Brockton V, Naed SV, Berneye K, Fugmann SD, Anderson MK, Pancer Z, Cameron RA, Smith LC, Rast JF (2006) The immune gene repertoire encoded in the purple sea urchin genome. Dev Biol 300:349–365

39. Huang S, Yuan S, Guo L, Yu Y, Li J, Wu J, Ge J, Yu Y, Huang H, Dong M, Yu C, Chen S, Xu A (2008) Genomic analysis of the immune gene repertoire of amphioxus reveals extraordinary innate complexity and diversity. Genome Res 18:1112–1126

40. Aruffo AK, Sweet MJ (2013) Differences in the repertoire, regulation and function of Toll-like Receptors and inflammasome-forming Nod-like Receptors between human and mouse. Curr Opin Microbiol 16:303–310

41. Quiniou SM, Boudinot P, Bengtén E (2013) Comprehensive survey and genomic characterization of Toll-like receptors (TLRs) in channel catfish, Ictalurus punctatus: identification of novel fish TLRs. Immunogenetics 65:511–530

42. Kanwal Z, Wiegertjes GF, Veneman WJ, Meijer AH, Spaink HP (2014) Comparative studies of Toll-like receptor signalling using zebrafish. Dev Comp Immunol 65:511–530

43. Roach JC, Glusman G, Rowen L, Kaur A, Purcell MK, Smith KD, Hood LE, Aderem A (2005) The evolution of vertebrate Toll-like receptors. Proc Natl Acad Sci U S A 102:957–962

44. Fritz JH, Ferrero RL, Philpott DJ, Girardin SE (2006) Nod-like proteins in immunity, inflammation and disease. Nat Immunol 7:1250–1257

45. Kanneganti TD, Lamkanfi M, Núñez G (2007) Intracellular NOD-like receptors in host defense and disease. Immunity 27:549–559
47. Franchi L, Park JH, Shaw MH, Marina-Garcia N, Chen G, Kim YG, Núñez G (2008) Intracellular NOD-like receptors in innate immunity, infection and disease. Cell Microbiol 10:1–8
48. Chen G, Shaw MH, Kim YG, Núñez G (2009) NOD-like receptors: role in innate immunity and inflammatory disease. Ann Rev Pathol 4:365–398
49. Schroder K, Tschopp J (2010) The inflammasomes. Cell 140:821–832
50. Rast JP, Smith LC, Loza-Coll M, Hibino T, Litman GW (2006) Genomic insights into the immune system of the sea urchin. Science 314:952–956
51. Stein C, Caccamo M, Laird G, Leptin M (2007) Conservation and divergence of gene families encoding components of innate immune response systems in zebrafish. Genome Biol 8:R251
52. Chang M, Collet B, Nie P, Lester K, Campbell S, Secombes CJ, Zou J (2011) Expression and functional characterization of the RIG-I-like receptors MDAS and LGP2 in Rainbow trout (Oncorhynchus mykiss). J Virol 85:8403–8412
53. Chang MX, Zou J, Nie P, Huang B, Yu Z, Collet B, Secombes CJ (2013) Intracellular interferons in fish: a unique means to combat viral infection. PLoS Pathog 9:1003736
54. Yu ZL, Li JH, Xue NN, Nie P, Chang MX (2014) Expression and functional characterization of PGRP6 splice variants in grass carp Ctenopharyngodon idella. Dev Comp Immunol 47:264–274
55. Bruns AM, Horvath CM (2012) Activation of RIG-I-like receptor signal transduction. Crit Rev Biochem Mol Biol 47:194–206
56. Su J, Huang T, Dong J, Heng J, Zhang R, Peng L (2010) Molecular cloning and immune responsive expression of MDAS gene, a pivotal member of the RLR gene family from grass carp Ctenopharyngodon idella. Fish Shellfish Immunol 28:712–718
57. Ohtani M, Hikima J, Kondo H, Hirono I, Jung TS, Aoki T (2011) Characterization and antiviral function of a cytoplasmic sensor gene, MDAS, in Japanese flounder, Paralichthys olivaceus. Dev Comp Immunol 35:554–562
58. Rajendran KV, Zhang J, Liu S, Peatman E, Kucuktas H, Wang X, Liu H, Wooda T, Terhunea J, Liu ZZ (2012) Pathogen recognition receptors in channel catfish: II Identification, phylogeny and expression of retinoic acid-inducible gene I (RIG-I)-like receptors (RLRs). Dev Comp Immunol 37:381–389
59. Zou PF, Chang MX, Xue NN, Liu XQ, Li JH, Fu JP, Chen SN, Nie P (2014) Melanoma differentiation-associated gene 5 in zebrafish provoking higher interferon-promoter activity through signalling enhancing of its shorter splicing variant. Immunology 141:192–202
60. Gaboriaud C, Juanhuix J, Gruen A, Lacroix M, Darnault C, Pignol D, Verger D, Fontecilla-Camps JC, Arlaud GJ (2003) The crystal structure of the globular head of complement protein C1q provides a basis for its versatile recognition properties. J Biol Chem 278:46974–46982
61. Desn SE, Thiel S (2013) Humoral pattern recognition and the complement system. Scand J Immunol 78:181–193
62. Fujita T (2002) Evolution of the lectin-complement pathway and its role in innate immunity. Nat Rev Immunol 2:346–353
63. Pangburn MK, Ferreira VP, Cortes C (2008) Discrimination between host and pathogens by the complement system. Vaccine 8:115–121
64. Cooper MA, Pommerring TL, Korányi K (2003) Primary immunodeficiencies. Ann Fam Physician 68:2001–2008