Expansion of C1Q Genes in Zhikong Scallop and Their Expression Profiling After Exposure to the Toxic Dinoflagellates

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C1Q (Complement 1Q) is an important recognition molecule in the immunological complement system, which could also be putatively involved in the stress responses induced by endotoxins or exotoxins, potentially through detoxification processes. Marine bivalves are well adapted to highly complex aquatic environments with various stressors. As filter feeders, they have to cope with highly potent microalgae-derived neurotoxins, such as paralytic shellfish toxin (PSTs). Zhikong scallops, \textit{Chlamys farreri}, are commercially important bivalve with the remarkable ability to accumulate PSTs. Exploring the C1Qs related to PST accumulation in \textit{C. farreri} could benefit our understanding of the adaptations of bivalve species. In the present study, we systematically analyzed C1Q genes in \textit{C. farreri}. In total, 97 \textit{CfC1Q} genes mainly from the expanded C1Q-B subfamily were identified, from which the \textit{CfC1QL}, \textit{CfC1QTNF}, and \textit{CfC1QDC1} members in \textit{C. farreri} were revealed to be under positive selection. Spatiotemporal expression analysis revealed that most \textit{CfC1QL} and \textit{CfC1QDC1} were highly expressed during the post-umbo stage and in hepatopancreas, while most \textit{CfC1QTNF} members were highly expressed after the creeping larva stage and in mantle. The hepatopancreas and kidney in \textit{C. farreri} are two toxin-rich organs with the highest concentrations of PSTs, acting as major “centers” for toxin accumulation and transformation, respectively. Therefore, after feeding the scallops with PST-producing dinoflagellates \textit{Alexandrium minutum} and \textit{Alexandrium catenella}, we determined the expression patterns of \textit{CfC1Qs} in these two organs. In kidney, more than 85% of \textit{CfC1QLs} and \textit{CfC1QDC1s} showed drastic up-regulation with both diets. However, among these members with significant induction, a different response manner was detected after feeding with \textit{A. minutum} and \textit{A. catenella}, respectively as acute and chronic response patterns. In comparison, far fewer \textit{CfC1Qs} showing
The C1Q proteins, also known as C1Q-domain-containing (C1QDC) proteins, widely exist in various species, from bacteria to mammals. Besides the conserved globular C1Q domain, they usually possess a leading signal peptide and a collagen-like region (CLR) (Liu et al., 2013). Previously, the vertebrate C1Q proteins were revealed to play a key role during immune and inflammation regulation (Kishore and Reid, 2000; Tom Tang et al., 2005; Huang et al., 2008; Yuzaki, 2008). As the first subcomponent of the complement C1 complex, C1Q is the target recognition protein of the classical complement pathway (Gestal et al., 2010). It plays a key role in initiating the complement activation through binding to antibody–antigen complexes, pathogen surfaces, apoptotic cells, or polyanionic structures (Gestal et al., 2010). At present, based on sequence homology, structure similarity, and functional relatedness, in-depth studies of C1Q proteins have been carried out in human (Homo sapiens) and zebrafish (Danio rerio). C1Q proteins have been classified into three major subfamilies: C1Q-A, -B, and -C (Tom Tang et al., 2005; Mei and Gui, 2008). In addition to the immunological functions, several C1Q proteins were also found to act as important neuronal transduction factors, mediating CNS (Central Nervous System) synapse development and in response to neurotoxicity as well as a variety of neuronal injuries (Stevens et al., 2007; Ghebrehiwet et al., 2012; Benoit et al., 2013). For example, C1Q was found to protect immature and mature primary neurons against fibrillar amyloid-β toxicity in mammals (Li et al., 2008; Benoit et al., 2013) and the precerebellin (CBLN) proteins (C1Q-B subfamily) were found to act as a new class of transneuronal regulatory factors, regulating synaptic development and synaptic plasticity in many regions of the brain (Yuzaki, 2008; Boulanger, 2009).

Marine bivalves are filter-feeding species that feed mainly on microalgae, and they can bioaccumulate various microalgae derived toxins, posing a health hazard to humans and other animals through food chains (Anderson et al., 2012; Borcier et al., 2017). Among these toxins, paralytic shellfish toxins (PSTs) are one of the most potent neurotoxins. PSTs attack the sodium channels (Na+) on nerve cell membranes through cross-pore binding at multi-polar active sites, inducing Na+-flux disruption and interfering transmission of an action potential (Bricelj et al., 2005; Borcier et al., 2017). The accumulation of PSTs can not only affect the biological function of bivalves, such as filtration/feeding activities, immune response, antioxidant response, and even reproduction, but also become a major concern in terms of their sanitary quality (Li et al., 2002; Fabioux et al., 2015). In invertebrate mollusks, the conservative functions of activating the complement system have been found in bivalve C1Q proteins. In Bay scallop (Argopecten irradians), AIC1QDC-1 and AIC1QDC-2 molecules exhibited significant yeast agglutination activity, and they could function as pattern-recognition receptors (PRRs) to recognize different pathogen-associated molecular patterns (PAMPs), triggering immune defense (Kong et al., 2010; Wang et al., 2012). In Zhikong scallop (Chlamys farreri), the CfC1Q (C1Q in C. farreri) proteins were found to combine multiple PAMPs to assist with pathogenic bacteria elimination (Zhang et al., 2008). They were also found to serve as pattern recognition receptors and bind with human heat-aggregated IgG in a dose-dependent manner (Zhang et al., 2008; Wang et al., 2012). Further, several lines of evidence have revealed that C1Q genes could be involved in toxin metabolism in bivalves, not only for PSTs, but also for amnesic shellfish poisoning (ASP) and environmental toxins (heavy metals, for instance). For example, several C1Qs have been identified as responsive genes during transcriptional research on mussel (Mytilus galloprovincialis) when exposed to PST-producing Alexandrium (Gerdol et al., 2014). ASP is caused by a species of the diatom genus Pseudo-nitzschia, which produces the toxin domoic acid. According to the differential gene expression analysis in mussels (M. galloprovincialis) naturally exposed to Pseudo-nitzschia, C1Qs were found to be significantly enriched and researchers pointed out that C1Qs could be involved in detoxification processes, in response to oxidative stress and/or in immunological processes (Pazos et al., 2017). In mussel (M. coruscus), time-dependent responsive pattern of a novel C1Q was revealed after copper and cadmium exposure, which might be developed as a potential indicator for monitoring heavy metals’ pollution (Liu et al., 2014). In particular, the number of C1Q genes was found to be enormously expanded in the genome of bivalves, such as Pinctada fucata and Crassostrea gigas (Takeuchi et al., 2016), which might be related to their adaptive functions. All these findings suggest that, besides the microbial recognition capability, the C1Q proteins could function in a

Keywords: PSTs challenge, bivalve mollusk, expression profiling, complement 1Q, genomic analysis

INTRODUCTION

The C1Q proteins, also known as C1Q-domain-containing (C1QDC) proteins, widely exist in various species, from bacteria to mammals. Besides the conserved globular C1Q domain, they usually possess a leading signal peptide and a collagen-like region (CLR) (Liu et al., 2013). Previously, the vertebrate C1Q proteins were revealed to play a key role during immune and inflammation regulation (Kishore and Reid, 2000; Tom Tang et al., 2005; Huang et al., 2008; Yuzaki, 2008). As the first subcomponent of the complement C1 complex, C1Q is the target recognition protein of the classical complement pathway (Gestal et al., 2010). It plays a key role in initiating the complement activation through binding to antibody–antigen complexes, pathogen surfaces, apoptotic cells, or polyanionic structures (Gestal et al., 2010). At present, based on sequence homology, structure similarity, and functional relatedness, in-depth studies of C1Q proteins have been carried out in human (Homo sapiens) and zebrafish (Danio rerio). C1Q proteins have been classified into three major subfamilies: C1Q-A, -B, and -C (Tom Tang et al., 2005; Mei and Gui, 2008). In addition to the immunological functions, several C1Q proteins were also found to act as important neuronal transduction factors, mediating CNS (Central Nervous System) synapse development and in response to neurotoxicity as well as a variety of neuronal injuries (Stevens et al., 2007; Ghebrehiwet et al., 2012; Benoit et al., 2013). For example, C1Q was found to protect immature and mature primary neurons against fibrillar amyloid-β toxicity in mammals (Li et al., 2008; Benoit et al., 2013) and the precerebellin (CBLN) proteins (C1Q-B subfamily) were found to act as a new class of transneuronal regulatory factors, regulating synaptic development and synaptic plasticity in many regions of the brain (Yuzaki, 2008; Boulanger, 2009). Marine bivalves are filter-feeding species that feed mainly on microalgae, and they can bioaccumulate various microalgae derived toxins, posing a health hazard to humans and other animals through food chains (Anderson et al., 2012; Borcier et al., 2017). Among these toxins, paralytic shellfish toxins (PSTs) are one of the most potent neurotoxins. PSTs attack the sodium channels (Na+) on nerve cell membranes through cross-pore binding at multi-polar active sites, inducing Na+-flux disruption and interfering transmission of an action potential (Bricelj et al., 2005; Borcier et al., 2017). 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According to the differential gene expression analysis in mussels (M. galloprovincialis) naturally exposed to Pseudo-nitzschia, C1Qs were found to be significantly enriched and researchers pointed out that C1Qs could be involved in detoxification processes, in response to oxidative stress and/or in immunological processes (Pazos et al., 2017). In mussel (M. coruscus), time-dependent responsive pattern of a novel C1Q was revealed after copper and cadmium exposure, which might be developed as a potential indicator for monitoring heavy metals’ pollution (Liu et al., 2014). In particular, the number of C1Q genes was found to be enormously expanded in the genome of bivalves, such as Pinctada fucata and Crassostrea gigas (Takeuchi et al., 2016), which might be related to their adaptive functions. All these findings suggest that, besides the microbial recognition capability, the C1Q proteins could function in a
toxin-responsive manner in bivalves, which may contribute to the adaptation of scallops for dealing with microalgal toxin challenge and merits further study.

Zhikong scallops (Hu et al., 2010) are commercially important shellfish, which thrive in environments from intertidal zones to deep oceans as semi-sessile filter feeders, with the outstanding ability to accumulate PSTs (Li et al., 2017). C. farreri are naturally distributed along the coasts of Northern China, Korea, Japan, and Eastern Russia. Most previous studies on bivalve C1Q proteins usually focus on a certain member that participated in the classical complement activation. In this study, we genome-wide identified the C1Q genes in C. farreri. The C1QL, C1QTNF, and C1QDC1 members from the expanded subfamily C1Q-B were revealed to be under positive selection. To explore these C1Qs’ involvement during PSTs accumulation, we chose two major strains of the PST-producer Alexandrium (A. catenella and A. minutum) to feed the scallops. In C. farreri, hepatopancreas and kidney are the main repository of PSTs during uptake and depuration, and dynamic PST metabolites were detected, contributing to PSTs detoxification (Hu et al., 2019; Lou et al., 2020). We systematically assessed the C1Q responses in hepatopancreas and kidney after feeding with toxic A. catenella and A. minutum, revealing the variable regulation of scallop C1Qs between organs and between different toxic microalgal species. Our findings imply the functional diversity of scallop C1Q genes in coping with microalgal-derived toxins, providing a better understanding on their adaptive evolution.

MATERIALS AND METHODS

Identification of C1Q Genes in Chlamys farreri

Based on our previous work (Li et al., 2017), the whole genome of a 2-years-old C. farreri was sequenced using the Illumina HiSeq 2000 platform through the construction and sequencing of both short-insert (180, 300, and 500 bp) and long-insert (2, 5, 10, 20, and 30 kb) DNA libraries. Thirteen adult tissues/organs of the scallop were chosen for transcriptome sequencing, including striated muscle, smooth muscle, foot, hepatopancreas, kidney, female gonad, male gonad, gill, eyes, mantle, cerebral ganglion, and visceral ganglion. All these data have been deposited on NCBI database. The available C1Q protein sequences of representative species, including humans (Homo sapiens), chickens (Gallus gallus), frogs (Xenopus laevis), zebrafish (Danio rerio), and mouse (Mus musculus) were downloaded from NCBI and UNIPROT databases. The downloaded orthologous C1Q sequences were used as query sequences for transcriptome and whole-genome blast searching in C. farreri, with the E value threshold as 1E-05. To confirm the completeness of C1Q gene family and to identify all possible C1Q gene loci in the genome, TBLASTN was used to verify the cDNA sequences by comparing the transcriptome sequences to the whole genome sequences. Further, sequence analysis was performed through C1Q HMM searching method to ensure the completeness of C1Q proteins, and the position of conserved C1Q Pfam domain (PF00386) was illustrated for C. farreri in Supplementary Table 1.

Sequence Analysis of CfC1Q Genes

To confirm the predicted amino acid sequences, the candidate C1Q genes sequences of C. farreri were submitted to the ORF Finder program to predict the open reading frame (ORF), then the ORFs were translated into amino acid sequences. The translated sequences were submitted to the SMART program to identify conserved domains. Compute pl/Mw tool was used to predict the information of isoelectric point (pI) and molecular weight. The protein structures of all the identified C1Q proteins were drawn with IBS1.0.3 software (Adema et al., 2017) and Geneious 7.1.7 was used to predict protein secondary structure. A web server GSDS (Gene Structure Display Server) was used to draw gene structure schematic diagrams. Based on the whole genome sequence, we analyzed the distribution of C1Q genes in genome and their gene linkage relationship.

Phylogenetic Analysis of C1Q Genes

To determine which subfamily the C1Q genes belong to, the phylogenetic analysis was performed using MEGA7 (Kumar et al., 2016). All C1Q proteins as well as C1Q proteins from human (Homo sapiens), chicken (Gallus gallus), frog (Xenopus laevis), zebrafish (Danio rerio), and mouse (Mus musculus) were employed to construct a phylogenetic tree. The C1Q protein sequences from these reference species were obtained from public databases such as NCBI and UNIPROT (see footnote). Multi-sequence alignment was performed using ClustalW (Thompson et al., 2003), and then was edited by Genedoc software (Nicholas, 1997). Poisson model and uniform rates were selected as the best fit model from MEGA 7.0. The tree was constructed based on neighbor-joining (NJ) method with a bootstrap of 1,000 replicates. Under the least stringent parameters, the poorly aligned amino acid positions were excluded using Gblocks, as implemented in the online version of the program (Castresana, 2000). In total, this analysis involved 235 amino acids across six animals.

Molecular Evolutionary Analysis of CfC1Q Genes

In order to test for positive selection at individual amino acid codons, the specific site model was implemented using the program of the PAML (Yang, 2007; Xun et al., 2020) under maximum-likelihood (ML) and Nei and Gojobori (NG) approach. Site model supposes that different branches of the phylogenetic tree are subject to the same selection pressure, but different amino acid positions experience different selection pressures (Gao et al., 2019). To measure the selective pressure at the protein level, this model allows the ratio to vary among

1https://www.ncbi.nlm.nih.gov/
2https://www.uniprot.org/
sites of the expanded C1Qs to estimate Non-synonymous (dN) and Synonymous (dS) substitution ratios (dN/dS or ω). Different positions of the same sequence have different ω values. In the situation of a ω = 1, <1, or >1 indicates neutral evolution, purifying selection/negative selection, or positive selection, respectively. In total, eight different hypothetical models were tested. Model M0 (single ratio) assumes that all sites have the same value of ω, whereas model M3 (discrete) assumes that all sites have a simple discrete distribution trend. Model M1a (neutral) assumes that there are two classes of sites in proteins with 0 < ω < 1 (conservative sites) and ω = 1 (neutral sites), and the ratio of these two types of sites is p0 and p1, respectively. M2a (selection) adds a proportion (p2) to account for a class of sites where ω 2 is estimated from the data and can be >1. Model M7 (beta) assumes that ω at all sites belongs to the matrix (0, 1) and is in beta distribution, whereas model M8 (beta and ω) additionally contains ω value (ω > 1). If the data fitted by M8 (M2a) is significantly better than M7 (M1a), we will apply the Bayesian Empirical Bayes (BEB) estimation method to determine the site category estimated to be positively selected.

**Spatiotemporal Expression of C1Q Genes in C. farreri**

The expression profiles of Zhikong scallop across all developmental stages and in adult organs/tissues were analyzed based on the RNA-seq data from our previous research (Li et al., 2017). Heat map of expression patterns of CfC1Q genes was drawn using custom R scripts (Robinson et al., 2010).

**Feeding the Scallops With Two Different Toxic Dinoflagellates: A. minutum and A. catenella**

Three hundred 2-years-old adult C. farreri were collected from the hatchery of Xunshan Group Co. Ltd. (Shandong, China). The scallops without shell damage were transported on ice to the laboratory and acclimated for 1 week in a recirculated aquarium system equipped with a filtering system. For the next 3 weeks, the scallops were fed with non-toxic microalgae Isochrysis galbana for depuration. After depuration, scallops were separated into six ventilated tanks independently as six different experimental group (50 scallops/each group). Three groups were fed with toxic A. minutum (AM-1 strain) and the other three groups were fed with toxic A. catenella (ACDH strain). Each scallop was fed once a day with 2,500 cells/mL for microalgae in a 3L volume and it could filter nearly all the microalgal cells within the same day (Navarro et al., 2006; García-Lagunas et al., 2013). Before feeding with toxic microalgae, samples were taken as control (0 day). After feeding with A. minutum or A. catenella, samples were taken at different test time points (1, 3, 5, 10, and 15 days) as treatment group. In detail, the specimens that were sampled at 1 and 3 days were taken as the acute response group and the specimens that were sampled at 5, 10, and 15 days were taken as chronic response group. At each test time point, three random scallops were selected and the hepatopancreas and kidney sample from each scallop were dissected and stored in liquid nitrogen directly for further RNA-seq analysis.

**Expression Analysis of CfC1Q Genes After Toxic A. minutum and A. catenella Feeding**

According to the phenol–chloroform method of RNA extraction (Shafer et al., 1997; Fanson et al., 2000), total RNA was extracted from 1mg sampled tissue specimen obtained from section “Feeding the Scallops With Two Different Toxic Dinoflagellates: A. minutum and A. catenella.” RNA-seq library was constructed using NEB Next mRNA Library Prep Kit, following the manufacturer's instructions, and was subjected to PE125 sequencing on the Illumina HiSeq 2000 platform (Park et al., 2019). After the RNA-seq data were pre-processed, quality filtering was carried out to screen out the sequences. The raw sequencing sequence was first processed with low quality pass rate, and then the sequence errors with NN were removed after the adapter sequences were removed. More than 80% of the sequences with a site quality value greater than 30 points were regarded as high-quality sequences and reserved for subsequent analysis. The RNA-seq sequences were aligned to the genome with TopHat (ver2.0.9). Expression of all C1Q genes was normalized and represented in the form of RPKM (Reads Per Kilobase of exon model per Million mapped reads) based on R software, following the formula total exon reads/mapped reads (millions)/exon length (Kb) (Hu et al., 2019). RPKM of the C1Q genes obtained from RNA-seq data were used to analyze their expression profile in kidney and hepatopancreas with different toxic dinoflagellates diets. Using the R package edgeR, the significant differences between the test group and the control group and the differentially expressed genes (DEGs) were detected with | log 2 FC | > 2 and p < 0.05. Using MeV 4.90 software, a heat map was drawn.

**RESULTS**

**Identification and Characterization of C1Q Genes in C. farreri**

In this study, a total of 97 C1Q genes were identified based on the genome and transcriptome of C. farreri. There were 77 members of C1QL, 7 members of C1QTNF, and 12 members of C1QDC1. The basic information (gene name, genomic location, number of exons, protein length, number of C1Q domains, isoelectric point, and molecular weight) of these C1Q genes was summarized in **Supplementary Table 1**. The coding sequence (CDS) of the longest C1Q gene was 2,331 bp, while the CDS of the shortest C1Q gene was only 97 bp. The exon number of cfC1Q genes varies from 1 to 10 (Figure 1), with encoding protein lengths ranging from 49 to 758 amino acids (Figure 2). The predicted molecular weights of CfC1Q proteins ranged from 5.1–84.9 kDa, with PI values from 3.97 to 9.49. At the N-terminal of all 97 CfC1Q proteins, the conserved C1Q domain could be detected, from

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http://www.r-project.org/

http://www.tm4.org
which 89 members contained one C1Q domain and eight members contained two C1Q domains. In addition, there were 21 CfC1Q proteins containing 1-4 coiled coil structures at the C-terminus. Further, we found most CfC1Qs contained one or more α-helix structures at the N-terminus or C-terminus, and there were multiple β-sheets, corners, and coiled coil structures in the middle (Supplementary Figure 1). According to the results of multiple sequence alignment of C1Q domain amino acids, most of the aromatic amino acids of C. farreri were relatively conservative, such as phenylalanine (F), tyrosine (Y), and tryptophan (W) (Supplementary Figure 2). It was consistent with previous findings that those aromatic amino acid residues participated in architecting the hydrophobic core of C1Q center (Liu et al., 2013). Genomic distribution analysis showed that C1Q gene linkage phenomenon was widespread in C. farreri (Supplementary Table 2). There were 66 C1Q genes found unequally distributed on 44 scaffolds, and among them, 35 genes were found located in terms of gene clusters. Of note, scaffold26237, scaffold64279, and scaffold56669 contained the biggest C1Q gene cluster, consisting of four CfC1QL members. Scaffold48207 and scaffold49013 were found to consist of three C1Q and CfC1QDC1 members, respectively. In addition, out of the 18 chromosomes that C1Q genes located, chromosome 5 contained the largest number of C1Q genes, ten of which were all members of CfC1QL, followed by chromosomes 18 and 19, both of which contained seven C1Q genes.

**Phylogenetic Analysis**

A phylogenetic tree was constructed using the Neighbor-Joining (NJ) method (Figure 3). We noticed that subfamily C1Q-B (red) and C1Q-C (pink) were clustered together first. In comparison, members of C1Q-A subfamily from vertebrate were found clustered together as a branch (blue). All CfC1Qs except one protein, CFA.C1QTNF.5198.10 (green), were found clustered into C1Q-B subfamily, revealing the significant expansion of this subfamily in C. farreri. To put it further, members of the C1Q-B subfamily were subdivided into four subcategories, namely C1QDC1, CfC1QL, CBLN, CfC1QTNF, and, respectively, represented as yellow, light blue, dark purple, and rose red. Of note, the CBLN members were found lost in C. farreri. The expansion of CfC1QLs was most obvious, in which 77 members were identified, accounting for 80% of CfC1Q members.

**Evolutionary Analysis**

In order to explore whether or not the expanded CfC1Q genes were impacted under positive selective pressure, we assess their non-synonymous and synonymous substitution divergences (dN/dS, ω) using the codon models implemented in PAML. As shown in Table 1, for all three major subcategories of expanded CfC1Q-B subfamily, M0 (one-ratio) was rejected in favor of the alternative model M3 (discrete) (M3 vs. M0, p < 0.05), revealing their different ω ratios. Meanwhile, in comparison between the M7 and M8 models, the p-value of the chi-square test for CfC1QLs, CfC1QTNFs, and CfC1QDC1s, respectively, reached 1e-09, 9e-04, and 0.027, demonstrating that they all allow for the positive selection. Furthermore, one positive selection site in the CfC1QL subfamily and seven positive selection sites in the CfC1QTNF subfamily was worth noting, which further supports the existence of positive selection for the lineage specific expanded subfamilies.

**Spatiotemporal Expression Pattern of C1Q Genes**

Expression patterns of CfC1Q genes during different developmental stages (Figure 4) and in different adult tissues (Figure 5) were analyzed. For CfC1QLs and CfC1QDC1s, most genes began to express on a large scale until umbo-early larva
stage, which continue to elevate their expression level and reach peak expression either in umbo-post larva or after creeping larva stage (with a log₂RPKM ranging from 0 to 524.48). For most CfC1QTNFs, their period of high expression shifted later, usually during the creeping larva stage. However, one C1QTNF3 (CFA.8357.34) showed elevated expression during
FIGURE 3 | Phylogenetic trees of selected organisms’ C1Q amino acid sequences. The whole amino acid sequence of C1Q proteins were used to perform multiple alignment and phylogenetic analysis using NJ method computed using the poisson correction method. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown above the branches. In the figure, different symbols were used to represent different subfamilies. Black solid dots represent A and C subfamily, colored solid dots represent B subfamily (HSA; MMU; Mus musculus; GGA; Gallus gallus; XLA; Xenopus laevis; DRE; Danio rerio; CFA).

zygote cleavage and one C1QTNF6 (CFA.49845.1) showed elevated expression in blastula. Similarly, the tissue expression pattern of CfC1QLs and CfC1QDC1s is consistent, most of which showed hepatopancreas dominant expression. Of note, three CfC1QLs (CFA.64863.15, CFA.60489.44, and CFA.64279.84) and two CfC1QDC1s (CFA.49013.20 and CFA.62185.4) out of these 90 genes showed obvious different tissue-dominant expression, differentially in female gonad, gill, kidney, and smooth muscle. Meanwhile, diverse tissue expression could be detected for C1QTNF genes. Similar to the situation of
TABLE 1 | Site model analysis for the C1Q subfamilies.

| C1Q     | No. copies | LRT statistics (p-value) (No. Sites Pr > 95%) |
|---------|------------|-----------------------------------------------|
|         |            | M0 vs. M3          | M1a vs. M2a          | M7 vs. M8          |
| C1QL    | 713.86(0)  | 0(1)              | 41.51(0.000000001)(1) |
| C1QTNF  | 232.13(0)  | 0(1)              | 14.02(0.0009)(7)     |
| C1QDC1  | 166.24(0)  | 0(1)              | 7.28(0.027)          |

LRT is likelihood ratio test statistic for M0 vs. M3; M1a vs. M2a; M7 vs. M8 from PAML package. Significant results with p < 0.05 are highlighted in bold. Pr, posterior probability for sites under positive selection.

CfC1Qls and CfC1QDC1s, three C1QTNF6 genes (CFA.60489.5, CFA.45303.3, and CFA.49845.1) exhibited high mRNA levels, respectively, in kidney, female gonad, and smooth muscle. Two other members of C1QTNF6 and both C1QTNF3 members showed highest expression in mantle, suggesting their differential biological function in comparison with hepatic major expressed CfC1Qs.

Expression of CfC1Qs in Kidney After Feeding Toxic A. minutum and A. catenella

In kidney, the organ that has been proven to transform incoming PSTs into derivatives with higher toxicity (Hu et al., 2019), more than 85% of CfC1Qls and CfC1QDC1s showed drastic up-regulation with both diets (Figure 6 and Supplementary Table 3). After feeding A. minutum, 26 C1QL members and 2 C1QDC1 members were significantly up-regulated (|log2 FC| > 2). After feeding A. catenella, 18 C1QL members and 1 C1QDC1 member were significantly up-regulated (|log2 FC| > 2). However, among the members with significant induction, most of them showed different response manners after feeding with A. minutum or A. catenella. With A. minutum diets, 26 CfC1Qs showed acute responses, while with A. catenella diets, chronic responses could be detected in 13 CfC1Qs. After feeding with A. minutum or A. catenella, all C1QTNF6 genes showed down regulation tendencies on day 10 and day 15. With AM-1 diet, two members (CFA.59133.4 and CFA.49845.1) showed significantly reduced expression on day 5 and day 1, respectively. All C1QTNF6 showed significant downregulation with ACDH diet, most of which responded acutely on day 1 or day 3, while only one gene (CFA.59133.4) responded chronically on day 5.

Expression of CfC1Qs in Hepatopancreas After Feeding Toxic A. minutum and A. catenella

In hepatopancreas, the main organ for accumulating the incoming microalgae-derived PSTs, most CfC1Qs only showed mild regulation pattern with toxic dinoflagellates diets (Figure 6 and Supplementary Table 3). Together, a total of 29 gene members showed significant responses with two toxic diets, among which 11 and 18 members, respectively, showed up- and down-regulation patterns. Different from the situation in kidney, for C1QTNFs, C1QLs, and C1QDC1s, no obvious or consistent regulation pattern could be found between or within these categories. However, we did find several members which showed consistent regulation with both toxic diets. For example, after feeding with AM-1 and
AcDH diets, significantly declined expression of three C1QL members (CFA.64863.15, CFA.19561.21, CFA.52505.1) and one C1QTNF6 (CFA.59133.4) could be detected, in acute and chronic manners, respectively. Besides, expressions of one C1QTNF3 (CFA.8357.34) and one C1QL (CFA.54573.12) were both significantly and acutely enhanced after feeding with AM-1 and ACDH diets. Meanwhile, the dinoflagellate-type dependent response manner that was mentioned in kidney was also detected for several other genes. For example, with AM-1 and ACDH diet, C1QTNF6 (CFA.60489.5) and C1QL (CFA.37535.22) showed acute and chronic up-regulation, respectively, not to mention that four members (CFA.62173.12, CFA.6499.30, CFA.721399.1, and CFA.18207.1) only showed significant up-regulation with AM-1 diet.

**DISCUSSION**

Filter-feeding marine bivalves are able to cope with highly potent microalgae-derived neurotoxins, such as paralytic shellfish toxins (PSTs). Exploring the genes related to toxin resistance in bivalves could benefit our understanding of the adaptations of these species. C1Q proteins are active molecules with multiple biological functions (Mj and Mf, 2016). They can activate the classical pathway of complement, clear pathogens, phagocytose, and lyse bacteria, and regulate inflammation (Kishore and Reid, 2000). It was reported that C1Qs could be involved in the detoxification processes in bivalves, in response to the induced stresses and the affected physiological state (Pazos et al., 2017). Zhikong scallops possess the remarkable ability to accumulate PSTs. In this study, we systematically identified CfC1Q genes and analyzed their responsive pattern after feeding with PST-producing dinoflagellates. The candidate responsive CfC1Qs might be developed as potential molecular indicators for monitoring toxin accumulation in edible scallops.

The C1Q proteins that widely existed in many species are structurally conserved. The typical C1Q domain can form a jam roll shape composed of 10 β-strands. Eight conserved amino acid residues exist in the center of the C1Q domain, among which five aromatic amino acid residues constitute the central hydrophobic core, and these conserved amino acid residues play an important role in forming or stabilizing the hydrophobic core of the C1Q domain (Gaboriaud et al., 2012). As shown in Figure 4, most of the aromatic amino acids of C. farreri were relatively conservative, which may suggest a similar hydrophobic function (Liu et al., 2013). On a side note, the existence of the conserved C1Q domain also reflects its role as one kind of charge pattern recognition molecule in order to bind certain ligands (Gestal et al., 2010). In C. farreri, approximately 90% C1Q proteins contain 1-2 C1Q domains at the N-terminal, and 0-1 coiled coil structures at the C-terminal, consistent with the C1Q structural characteristics of D. rerio and H. sapiens. Of note, there are nine CfC1Q proteins that contain multiple curled helices, and one member possesses three C1Q domains, showing a distinguishable structure that might be specific in bivalve lineage.

Despite their wide existence, the retention or loss of C1Q genes occurred during the evolution of metazoans (Supplementary Table 4). Fungi and Plantae seem to be completely free of C1Q genes (Yuzaki, 2008). According to a previous study, a total of seven C1Q genes have been identified in sea urchin...
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**FIGURE 6** | Heatmap of C1Q genes expression in C. farreri hepatopancreas and kidneys after feeding with AM-1 and ACDH. The red branch represents the C1QTNF member, the blue branch represents the C1QL member, and the yellow branch represents the C1QDC1 member. * indicated the significantly regulated C1Q genes with $|\log_{2} \text{FC}| > 2$ and $p < 0.05$ (for interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article).

(Hibino et al., 2006), and only two CIQs were found in sea squirt *Ciona intestinalis* (Azumi et al., 2003). Then, the number of CIQs seems to increase during evolution, for instance, 50 CIQ genes were discovered in amphioxus *Branchiostoma floridae*, 52 genes in zebrafish *D. rerio*, and 31 in human (Tom Tang et al., 2005; Huang et al., 2008; Mei and Gui, 2008; Yuzaki, 2008). Further, we found that the CIQ genes had quite obvious gene expansion in mollusks. For example, in mussel *Mytilus californianus* and
M. galloprovincialis, respectively, 52 and 168 genes could be found, and in oyster C. gigas, 321 C1Q genes have been found (Gerdol et al., 2011; Zhang et al., 2015). In the present study, 97 C1Qs were identified in scallop C. farreri, from which the C1Q-B subfamily was found being mainly expanded. Phylogenetic analysis showed that proteins of C1QD1C and C1QL were first clustered together, suggesting they may share a common ancestor gene. Besides, the CBLN members were lost in C. farreri, and the C1C1QTNF members were isolated in a single cluster and separated from the C1QTNF members of vertebrates, indicating the independent gene replication events in the bivalve lineage. Tandem gene duplication and an unequal crossing over drove the expansion of the C1Q gene family in bivalves, which represented an important evolutionary advantage in these filter-feeding organisms (Gerdol et al., 2019). Under the circumstances of various stressors in the ocean, the expansion of bivalve C1Q genes may provide insights into lineage specific adaptations. Adaptive evolution is usually driven by positive selection, the acquisition of new genes combined with the modification of existing genes (Ellegren, 2014). Our results showed that, for all three major subcategories of expanded CfC1Q-B subfamily, they all allow for the positive selection, and one positive selection site in the C1C1QTNF subfamily and seven positive selection sites in the C1C1QTNF subfamily was revealed. These results also remind us that the C1Q gene family could be a good candidate gene family for lineage specific molecular evolution analysis.

Expression of C1Q genes in C. gigas was found to be tightly regulated during early development, most of which were not expressed at all until the larva settling (Gerdol et al., 2015). Researchers pointed out that, during the planktonic/benthic life transition, the cumulative expression of C1Q genes in the frame of a radical reorganization of body parts may serve to suit a sedentary existence. Consistent expression patterns during the development of C. farreri were found. Staggered expression peaks for different C1C1Q subcategories could be detected, which may suggest their functional differentiation during development. Most C1C1QDs and C1C1QDClCs showed obvious elevated expression in umbo larva, which might be involved with the formation of eye points and motor organs. However, most C1C1QTNFs began to express in a large scale during the metamorphosis development and may contribute to stress tolerance to assist with body reorganization.

The specificity of C1Q genes’ expression in a certain tissue may be related to the tissue specific functions (Yang et al., 2013). From Figure 6, we can see that C1Q genes were highly specifically expressed in hepatopancreas, consistent with previous study results in Pacific oyster, giant freshwater prawn, and Nile tilapia (Fabiox et al., 2015; Ye et al., 2015). Hepatopancreas is an important organ for digestion and absorption, and it is equivalent to the fat body of insects or the liver of mammals, serving as the first defense barrier against pathogen invasion (Gross et al., 2001). Therefore, we speculate that the C1C1Q genes play important roles in resisting external aggressions in the organism and maintaining the steady state of the internal environment. Besides, most C1QTNF members showed relative high expression level in mantle, the tissue that possess numerous tentacles and eyes to perceive the surrounding environment (Land, 1966). The enhancement of sense skills expands the range of exercise against predators and is conducive for food intake, and mantle-specific high level of C1QTNF expression might contribute to future studies.

Zhihong scallops could tolerate and accumulate potent neurotoxic PSTs. PSTs comprise several structural variants, such as saxitoxin (STX), sulfated gonyautoxins (GTX), and N-sulfocarbamoyl toxins (B- and C-toxins), of which STX is the most neurotoxic (Thottumkatara et al., 2014). Our previous study found that Zhihong scallop possess two sodium channel genes (Nav1 and Nav2), and Nav1 has a potentially toxin-resistant T mutation at position 1,425 (Li et al., 2017). This mutation has been proven to be able to yield a significant resistance to STX and TTX in rat (Jost et al., 2008). Alexandrium spp is one of the major PSTs-producing dinoflagellates. It has been reported that A. minutum (AM-1 strain) mainly produces gonyautoxins (mainly GTX1-4), while A. catenella (ACDH strain) mainly produces N-sulfocarbamoyl toxins (CJ1/2) (Tian et al., 2010; Li et al., 2017; Blossom et al., 2019). Hepatopancreas and kidney are two major bivalve toxic organs (Tian et al., 2010; Li et al., 2017), and diverse regulation of C1C1Qs presented in hepatopancreas and kidney after challenge with A. minutum or A. catenella. Compared with hepatopancreas, it is obvious that the responses of most C1C1Qs were more intense in kidney, which may attribute to the more toxic analogs’ transformation in kidney (Joerink et al., 2006; Tellez-Bañuelos et al., 2009; Xu et al., 2018). Besides, dinoflagellate type dependent gene regulation was also detected in kidney. For C1C1QDs and C1C1QDClCs, acute and chronic responses could be, respectively, detected after feeding with A. minutum and A. catenella. This result reinforced our inference that different toxin composition and virulence could be the causal factors for C1C1Q genes’ responsive pattern. Meanwhile, although not significant, the induction of the C1C1QTNF3 (CFA.41509.64) in kidney could be detected just like the regulation of C1C1QDs and C1C1QDClCs. However, the responses of five C1C1QTNF6 members in kidney are obviously regulated in the opposite way, demonstrating the functional differentiation for C1C1Q genes. In both kidney and hepatopancreas, two C1Qs (CFA.721399.1 and CFA.18207.1) were significantly up-regulated with AM-1 diet, and two members (CFA.59133.4 and CFA.49845.1) were significantly down-regulated with ACDH diet. Only one member (CFA.52505.1) showed consistent significant downregulation with both AM-1 and ACDH diet and in both kidney and hepatopancreas, and these five genes could be good candidates as tissue responsive indicator genes with PSTs challenge.

CONCLUSION

The present study demonstrated a systematic genomic and transcriptomic survey of C1Q genes in C. farreri. Phylogenetic analysis indicated a bivalve-specific gene duplication event may occur for C1Q genes, and expansion of C1C1Q-B subfamily was revealed. Diverse spatiotemporal expression patterns of C1Q genes were observed, and their response manners to the stresses of PST-producing A. minutum and A. catenella varied between scallop tissues or between dinoflagellate strains. These findings
may suggest the adaptive functional diversity of expanded C1Q genes in C. farreri, which could serve as a potential candidate gene family for lineage specific molecular evolution analysis and several members could be developed as potential indicator genes for PST challenges. We believe our study provides valuable information that would offer insights into the evolution and functional characteristics of C1Q genes in bivalves, and in the future, further in-depth research on the functional divergence for different members would be needed.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: The accession numbers of genomic data are SRR1305705, SRR2486272, SRR2486273, SRR2486281–SRR2486284, SRR2486300, and SRR2913253–SRR2913260. The accession numbers of transcriptomic data are SRRX244484–SRRX244876, SRRX2508197–SRRX2508199, SRRX244468–SRRX2444682, SRRX2445405–SRRX2445440, and SRR11840040–SRR11840053.

AUTHOR CONTRIBUTIONS

SL, SW, and ZB conceived and designed the study. KX and YW performed the experiments. XC and XD participated in data analysis. KX and SL wrote the manuscript. LZ and JH participated in the revision of the article. All authors contributed to the article and approved the submitted version.

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FUNDING

We acknowledge the grant support from the National Natural Science Foundation of China (32002373), the Fundamental Research Funds for the Central Universities (202064008), the Project of Sanya Yazhouwan Science and Technology City Management Foundation (SKJC-KJ-2019KY01), and the Taishan Scholar Project Fund of Shandong Province of China.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2021.640425/full#supplementary-material

Supplementary Figure 1 | The secondary structures of C1Q domain. The pink cylinder stands for alpha helices; the orange arrows stand for beta strands. The curved arrows stand for turns and the wavy lines stand for coils.

Supplementary Figure 2 | Alignment of deduced C1q amino acid sequences. The gray and black shadow, respectively, show the most shared amino acids and the conserved amino acid group (such as aromatic amino acids) in C. farreri.

Supplementary Table 1 | Summary of the C1Q gene family in Chlamys farreri genome.

Supplementary Table 2 | Distribution of C1q genes on chromosome of Chlamys farreri genome.

Supplementary Table 3 | Summary of C1q genes regulated expression in C. farreri kidney and hepatopancreas after feeding the scallops with PST-producing AM-1 and ACDH.

Supplementary Table 4 | Distribution of C1q genes in different species.
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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