A Novel Cold-Regulated Cold Shock Domain Containing Protein from Scallop *Chlamys farreri* with Nucleic Acid-Binding Activity

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Abstract

**Background:** The cold shock domain (CSD) containing proteins (CSDPs) are one group of the evolutionarily conserved nucleic acid-binding proteins widely distributed in bacteria, plants, animals, and involved in various cellular processes, including adaptation to low temperature, cellular growth, nutrient stress and stationary phase.

**Methodology:** The cDNA of a novel CSDP was cloned from Zhikong scallop *Chlamys farreri* (designated as CfCSP) by expressed sequence tag (EST) analysis and rapid amplification of cDNA ends (RACE) approach. The full length cDNA of CfCSP was of 1735 bp containing a 927 bp open reading frame which encoded an N-terminal CSD with conserved nucleic acids binding motif and a C-terminal domain with four Arg-Gly-Gly (RGG) repeats. The CSD of CfCSP shared high homology with the CSDs from other CSDPs in vertebrate, invertebrate and bacteria. The mRNA transcripts of CfCSP were mainly detected in the tissue of adductor and also marginally detectable in gill, hepatopancreas, hemocytes, kidney, mantle and gonad of healthy scallop. The relative expression level of CfCSP was up-regulated significantly in adductor and hemocytes at 1 h and 24 h respectively after low temperature treatment (*P*<0.05). The recombinant CfCSP protein (rCfCSP) could bind ssDNA and in vitro transcribed mRNA, but it could not bind dsDNA. BX04, a cold sensitive *Escherichia coli* CSP quadruple-deletion mutant, was used to examine the cold adaptation ability of CfCSP. After incubation at 17°C for 120 h, the strain of BX04 containing the vector pIII showed growth defect and failed to form colonies, while strain containing pIII-CSPA or pIII-CSPG grew vigorously, indicating that CfCSP shared a similar function with *E. coli* CSPs for the cold adaptation.

**Conclusions:** These results suggest that CfCSP is a novel eukaryotic cold-regulated nucleic acid-binding protein and may function as an RNA chaperone in vivo during the cold adaptation process.

Introduction

All living organisms must adapt to changes in the environment, such as cold shock. Increasing evidence has confirmed the importance of cold-induced proteins as molecular chaperones involved in the cold adaptation [1–3]. Among these proteins, cold shock domain (CSD) containing proteins (CSDPs) are one group of the evolutionarily conserved nucleic acid-binding proteins and they are widely distributed in bacteria, plants, and animals [4–6]. These CSDPs are involved in various cellular processes, including adaptation to low temperatures, cellular growth, nutrient stress and stationary phase [7].

In prokaryotes, the members of CSDPs are called cold shock proteins (CSPs) and they have been extensively studied in *Escherichia coli*. These proteins consist of a cold shock domain (CSD) with two consensus RNA-binding motifs (RNP1 and RNP2) [8–10] and they are responsible for cold shock response. There are nine CSP genes identified from *E. coli*, and four (CSPA, CSPB, CSPG and CSPI) of them can be induced by cold shock [3]. CSPG, the most predominant CSP, may accumulate up to 10% of total proteins after low temperature exposure [11]. BX04, an *E. coli* strain with quadruple-deletion of CSPA, CSPB, CSPG and CSPE, can not grow at low temperature [12,13]. Under low temperature, bacterial CSPs can bind to RNA and destabilize the secondary structures of RNA to prevent the premature transcription termination. CSPA, CSPG, and CSPE have been confirmed to possess *in vivo* and *in vitro* transcription antitermination activity [14]. CSPA mRNA is able to sense the temperature downshifts, and adopt functionally distinct structures at different temperature, even without the aid of trans-acting factors [15].

In eukaryotes, the CSDPs display multiple functions with the structural features of variable N-terminal sequences [16], diverse auxiliary C-terminal domains and a highly conserved CSD [7,17]. Based on the C-terminal domain, eukaryotic CSDPs are divided
into three classes. The most extensively studied eukaryotic CSDPs are the Y-box (YB) proteins with C-terminal basic/aromatic islands as transcription factors to regulate gene expression [18–25]. For example, both the human YB-1 [19,20] and the Xenopus FRGY2 [21–24] function as components of the messenger ribonucleoprotein complex (mRNP) to regulate translation. Another class of eukaryotic CSDPs includes LIN-20 from Caenorhabditis elegans [25] and a group of glycinergic-rich plant proteins [26] with C-terminal retroviral-type zinc fingers. LIN28 are involved in the enhancement of translation [27], biogenesis of miRNA [28] and generation of induced pluripotent stem cells [29]. Most of the plant CSDPs, such as AtCSP2 [30,31], AtCSP3 [32] and WCSPI [33,34], act as RNA chaperones in response to low temperature. The third class of eukaryotic CSDPs are identified in a range of invertebrates [35–39] with the typical feature of RGG motif, which is defined as closely spaced Arg-Gly-Gly repeats separated by other, often aromatic amino acids [40]. However, the information about the function of invertebrate CSDPs is very limited. ApY1 from Aplysia californica [35] and YPS from Trypanosoma brucei is involved in kinetoplastid RNA editing and/or translation [41,42].

Compared with those in prokaryotes, the study on eukaryotic CSDPs, especially on those of invertebrate is still at the beginning, and there is no report about the involvement of the invertebrate CSDPs in the cold shock response. Zhikong scallop (Chlamys farrei), one of the most important cultured scallop species in North China, has developed protective mechanisms in response to the cold stress in winter. In the present paper, a cold shock domain containing protein (CfCSP) was identified from Chlamys farrei. The mRNA transcripts of CfCSP in different tissues and its temporal expression in adductor and hemocytes after acute cold shock treatment were investigated. The in vitro nuclear acids binding activity of the recombinant protein and the in vivo functional complementation of bacterial mutants were also examined to characterize its roles in the cold shock response of scallop.

**Materials and Methods**

**Ethics statement**

The scallops used in the present study are marine cultured animals, and all the experiments are conducted according to the regulations of local and central government.

**Scallop, cold shock treatment and tissue collection**

Adults of scallop C. farrei with an average 55 mm of shell length were collected from a farm in Qingdao, Shandong Province, China, and maintained in the aerated seawater at 16°C for a week before processing.

For the tissue distribution analysis of CfCSP mRNA, six tissues, including gill, hepatopancreas, kidney, mantle, gonad and muscle from five healthy adult scallops were collected. Hemolymph from these five scallops was collected from the adductor muscle and then immediately centrifuged at 800 × g for 10 min to harvest the hemocytes. All these tissue samples were stored at –80°C after addition of 1 mL TRIzol reagent (Invitrogen) for subsequent RNA extraction.

Forty scallops were employed in the acute cold shock treatment experiment. Thirty five scallops were cultured in 24 L tanks containing aerated seawater at 4°C, and other 5 scallops were still kept in 24 L tanks containing aerated seawater at 16°C as the blank group. Five individuals were randomly collected from the experimental group at 1, 3, 6, 12, 24 and 26 h after they were cultivated at 4°C. Muscle and hemocytes from the scallops were collected and stored as described above.

**RNA isolation and cDNA synthesis**

Total RNA was isolated from the tissues of scallops using Trizol reagent (Invitrogen). The first strand cDNA synthesis was carried out based on Promega M-MLV RT Usage information using the DNase I (Promega)-treated total RNA as template and oligo (dT)-adaptor primer P1 (Table 1). The reaction mixtures were performed at 42°C for 1 h, terminated by heating at 95°C for 5 min, and subsequently stored at –20°C.

**Cloning of the full-length CfCSP cDNA**

BLAST analysis of all the EST sequences from the C. farrei cDNA library [43] revealed that one EST (no. nscag0_004862; 1373 bp) was homologous to previously identified CSDPs in other animals and a gene specific sense primer P2 (Table 1) was designed to clone the full sequence cDNA of CiCSP by rapid amplification of cDNA ends (RACE) approach. The 3’ end of CiCSP cDNA was obtained using primers P2 and P3 (Table 1). The PCR product was cloned into the pMD18-T simple vector (TaKaRa) and sequenced with primers P4 and P5 (Table 1). The resulting sequences were verified and subjected to cluster analysis.

**Sequence analysis**

The cDNA sequence and deduced amino acid sequence of CiCSP were analyzed using the BLAST algorithm (http://www.ncbi.nlm.nih.gov/blast) and the Expert Protein Analysis System (http://www.expasy.org/). The protein domains were predicted with the simple modular architecture research tool (SMART) version 4.0 (http://www.smart.embl-heidelberg.de/). The ClustalW

### Table 1. Primers used in this study.

| Primer          | Sequence (5’-3’)                  |
|-----------------|-----------------------------------|
| Clone primers   |                                   |
| P1 (oligo (dT)-adaptor) | GCCACCGCTGCACTAGTACT17 |
| P2 (CiCSP primer) | TAGAAAGAACAACCACTACTAGCG |
| P3 (T7)         | GATAATCAGACTCCTATAGGGC |
| Sequencing primers |                               |
| P4 (M13-47)    | CGCCAGGAAAACTTTCCAGTCAGAC |
| P5 (Rv-M)      | GAGGCGTAAACAAATTTCACACAGG |
| RT primers     |                                   |
| P6 (CiCSP-RTF) | GACGTCCATTACAAGAACACCC |
| P7 (CiCSP-RTR) | GCCAGGTCCTCCGTTAGTAA |
| P8 (β-actin-RTF)| CAAACGACGGCTCCCTCGTCAT |
| P9 (β-actin-RTR)| CTGCGGACCCTGAACCTTGGT |
| Transcription primers |                             |
| P10 (Luc-mRNAF)| GCCGTTATTGACTATAGGATGAGAGCACG |
| P11 (Luc-mRNAR)| TTACCACGCAGTTTCCCACG|
| Recombination primers |                         |
| P12 (CiCSP-Ref1) | TTGAGTGCTGCAAGAAATGTGACACCTTATA |
| P13 (CiCSP-ReR1) | AGAGAGCTCGCAGGAGTACCTACT |
| P14 (CiCSP-Ref2) | TTACATAGGCAAAATAGGTACACCTTTATA |
| P15 (CiCSP-ReR2) | AGAGAGCTCCAGGAGTACCTACT |
| P16 (CSPA-Ref)  | TTACATAGGTCGGTAAAGATTGAGTACCTTAT |
| P17 (CSPA-ReR)  | AGAGAGCTCCAGGAGTACCTACT |

<sup>doi:10.1371/journal.pone.0032012.t001</sup>
Multiple Alignment program (http://www.ebi.ac.uk/clustalw/) was used to create the multiple sequence alignment.

Real-time PCR analysis of CfCSP mRNA expression

The cDNA mix was diluted to 1:100 and stored at −80°C for subsequent SYBR Green fluorescent quantitative real-time PCR (RT-PCR). Two CiCSP-specific primers, sense primer P6 and reverse primer P7 (Table 1), were used to amplify the corresponding product of 199 bp. The scallop β-actin, amplified with primers P8 and P9 (Table 1), was chosen as reference gene for internal standardization. DEPC-water for the replacement of cDNA template was used as negative control.

The SYBR Green RT-PCR assay was carried out in an ABI PRISM 7300 Sequence Detection System. The amplifications were conducted in triplicates in a total volume of 25 μL. Dissociation curve analysis of amplification products was performed at the end of each PCR reaction to confirm that only one PCR product was amplified and detected. After the PCR program, data were analyzed with SDS 2.0 software (Applied Biosystems). To maintain consistency, the baseline was set automatically by the software. The comparative average cycle threshold method was used to analyze the mRNA expression level of CiCSP, and the value stood for an n-fold difference relative to the calibrator [44]. All data were given in terms of relative mRNA expression as mean ± S.E. (N = 5). Differences were considered extremely significant at P<0.01 and significant at P<0.05.

Recombinant expression of CF CSP and purification of the fusion protein

The cDNA fragment encoding the mature peptide of CiCSP was amplified with the primers P12 and P13 (Table 1). A Sac I site was added to the 5' end of primer P12 and a Hind III site was added to the 5’ end of primer P13 after the stop codon. The PCR fragment was digested with restriction enzymes Sac I and Hind III (NEB), and ligated into predigested expression vector pET-30a (Novagen). The recombinant plasmid (pET-30a-CiCSP) was transformed into E. coli BL21 (DE3)-pLysS (Novagen). The recombinant CiCSP protein was purified with a Ni²⁺ chelating Sepharose column, and the purified protein was refolded as previously described [45]. Then the resultant protein was separated by reducing 15% SDS-polyacrylamide gel electrophoresis (SDS-PAGE), and visualized with Coomassie Brilliant Blue R250. The concentration of purified rCiCSP was quantified by BCA method [46].

Nucleic acid binding analysis

Gel retardation analysis with ss/dsDNA substrates was performed as previously described [33]. Totally 150 ng of either single-stranded (M13mp8) or double-stranded (M13mp8 RFI) DNA (NEB) was incubated with the his-tagged rCiCSP proteins in amounts ranging from 0, 20, 100, to 300 pmol. Nucleotide and protein were incubated in 15 μL of binding buffer (10 mM Tris-HCl, pH 7.5) at 0°C for 30 min.

Luciferase (lac) mRNA was in vitro transcribed from a sequence-confirmed firefly luc plasmid with a Ribomax kit (Promega). For in vitro transcription, two luc gene specific primers, sense primer P10 and antisense P11 (Table 1) were designed and the phage T7 promoter was added to the 5’ end of primer P10. The DNA template generated by PCR was purified and transcribed with T7 RNA polymerase using the Ribomax kit (Promega). The in vitro transcribed luc mRNA and the his-tagged rCiCSP protein were incubated in 15 μL of binding buffer (10 mM Tris-HCl, pH 7.5) at 0°C for 30 min. The purified rCiCSP proteins were added to binding reactions in amounts of 0, 20, 60, 120, and 240 pmol. The binding reactions were subjected to agarose gel electrophoresis and visualized by ethidium bromide staining [47].

Bacterial Complementation

For bacterial complementation studies, the cDNA fragment encoding the mature peptide of CiCSP was amplified with the primers P14 and P15 (Table 1), while the cDNA fragment encoding the E. coli CSPA was amplified from the strain ‘BL21-Gold (DE3) pLysS AG’ with the primers P16 and P17 (Table 1). An Nde I site was added to the 5’ end of primer P14 and P16, while an EcoRI site was added to the 5’ end of primer P15 and P17. The PCR fragments were digested with restriction enzymes Nde I and EcoRI (NEB), and ligated into predigested expression vectors pINIII. These pINIII constructs (pINIII-CiCSP and pINIII-CSPA) were transformed into E. coli BX04 (ΔcspAΔcspBΔcspCΔcspD) cells [12]. The E. coli strain JM109 with pINIII plasmid was selected as positive control and the E. coli strain BX04 with pINIII plasmid was selected as negative control. Overnight cultures of E. coli with respective plasmid were streaked on LB-ampiciline plates and grown at either 37 or 17°C.

Results

The sequence characters of CiCSP cDNA

A PCR product of 565 bp was amplified from cDNA template by 3’ RACE. By overlapping this segment with EST no. clag0_004862, a full-length CiCSP cDNA of 1735 bp was obtained (Fig. 1A) and deposited in GenBank under accession JN869460. It included a 5’-untranslated regions (UTRs) of 16 bp, a 3’-UTRs of 792 bp with a classical polyadenylation signal (AATAAA) and one A+U destabilizing elements (ATTTA), and an open reading frame (ORF) of 927 bp encoding a polyepitope of 308 amino acids with a predicted molecular weight of 34.57 kDa and theoretical isoelectric point (pl) of 9.69 (Fig. 1A).

Several distinct functional domains were identified in the deduced amino acid sequence of CiCSP (Fig. 1). A CSD with two consensus RNA-binding motifs (RNP1 and RNP2) and four DNA binding sites (Trp39, Phe48, Phe59 and Lys92) were located at the N terminus that contributed to the binding of nucleic acids (Fig. 1). Subsequent to the CSD, there were four RGG repeats in the C-terminal domains which were important for invertebrate CSDPs to bind nucleic acids (Fig. 1).

Homologous analysis of CiCSP

Sequences of CSDPs from bacteria, invertebrate and vertebrate were downloaded and used for Ident and Sim Analysis (Table 2). The CSD sequence of CiCSP shared significant sequence similarity with that of the CSDPs in invertebrates (96.9% identities to that of L. stagnalis and 95.4% to that of A. californica) and YB proteins in vertebrates (87.7% identities to YB3 of X. laevis and YB-1 of Human). However, the similarity between the CSD sequence of CiCSP and E. coli CSPA was relatively low (Percentage identities: 43.1%). The signature sequences of nucleic acid binding motif were identified in CiCSP by multiple sequences alignment (Fig. 2). The amino acid residues among the RNA-binding motif (RNP1 and RNP2) and the DNA binding sites (Trp39, Phe48, Phe59 and Lys92 in CiCSP) were well conserved except that three residues were substituted in the CSPA of E. coli (Fig. 2). An unrooted phylogenetic tree with four branches was constructed using neighbor-joining method with 1000 boot-strap test (Fig. 3). The CSPs containing one CSD from bacteria (including CSPA and CSPB of E. coli, LICSPB and BcCSPB) formed one branch. The plant CSDPs containing retroviral-type zinc fingers (including
AtCSP2, AtCSP3, WCSP1 and NsGRP2) gathered as another branch. CSDPs from invertebrate (including CfCSP, LsYB and ApY1) and vertebrate (including YB-1 and XlYB3) formed the third one, and the last branch was composed of LIN-28 from *C. elegans* (Fig. 3).

The distribution of CfCSP mRNA in different tissues

Quantitative real-time RT-PCR was employed to investigate the distribution of CfCSP mRNA in different tissues with β-actin as internal control. For CfCSP and β-actin genes, there was only one peak at the corresponding melting temperature in the dissociation curve analysis, indicating that the PCR was specifically amplified. The CfCSP transcript was ubiquitously detectable in all the tested tissues including gill, hepatopancreas, hemocytes, kidney, mantle, gonad and adductor. The highest CfCSP expression level was found in adductor, which was 31.62-fold of that in hemocytes (P<0.01). By contrast, the CfCSP mRNA transcript in gonad, kidney, hepatopancreas and gill was in low level, which was approximately 7.74-, 4.39-, 3.68- and 3.62-fold of that in hemocytes, respectively, and they are significantly higher than that in adductor.

**Figure 1.** Nucleotide and deduced amino acid sequences of CfCSP. The nucleotides and amino acids are numbered along the left margin. The putative CSD is boxed. The two consensus RNA binding domains are shaded in gray and four DNA binding sites are marked in italic and bold. The RGG repeat motifs are underlined. The classical polyadenylation signals and the A+U destabilizing elements in the 3′ UTR are marked in bold.

doi:10.1371/journal.pone.0032012.g001
Temporal expression of CfCSP mRNA in adductor and hemocytes after acute cold shock treatment

After acute cold shock treatment, the level of CfCSP transcript in adductor was up-regulated quickly and reached the maximum level of 5.98-fold \((P<0.05)\) within 26 h, indicating that CfCSP probably functions as a cold response gene in this species. In addition, the level of CfCSP transcript in hemocytes dropped back to normal level at 6 h \((1.49-\text{fold})\) and 12 h \((1.38-\text{fold})\), but no significant difference was observed between the treatment and the blank group \((P>0.05)\) (Fig. 5A).

After acute cold shock treatment, the mRNA expression level of CfCSP in hemocytes, was up-regulated and reached the maximum level of 3.11-fold \((P<0.05)\) at 26 h compared to the blank group. There was no significant difference \((P>0.05)\) between the mRNA expression level of CfCSP in treatment and control groups during the first 12 h after treatment (Fig. 5B).

Recombinant expression of CfCSP in E. coli

The plasmid pET-30a-CfCSP was transformed into E. coli BL21(DE3)-pLyS\(\phi\). After IPTG induction, the whole cell lysates was analyzed by SDS-PAGE, and a distinct band with molecular weight of \(\sim53\ \text{kDa}\) was revealed, which was consistent with the predicted molecular mass (Fig. 6). The concentration of the purified rCfCSP was 210 \(\mu\text{g/mL}\) measured by the BCA assay.

The nuclear acids binding activities of CfCSP

Gel mobility shift assays utilizing single- or double-stranded phage DNAs as substrates were performed to test the nucleic acid-binding activities of rCfCSP. Shifts of ssDNA bands were detected when 20 pmol rCfCSP was added to the binding reaction (Fig. 7A), and more clear shifts were observed when higher amount \((100\ \text{pmol}\ or\ 300\ \text{pmol})\) of rCfCSP was added. However, no shift was detected when dsDNA was incubated with 20 pmol, 100 pmol or 300 pmol rCfCSP proteins (Fig. 7B).

The RNA gel mobility shift assay was performed using in vitro transcribed luc mRNA as a substrate. The luc mRNA was incubated with rCfCSP at 0°C in the presence of an RNase inhibitor, and the binding reactions were subjected to 0.8% agarose gel electrophoresis (Fig. 7C). Shifts of luc mRNA bands appeared when the concentration of rCfCSP reached 60 pmol (Fig. 7C, lane 3) and it became more obvious when the amount of rCfCSP increased to 120 or 240 pmol (Fig. 7C, lane 4 and 5).

CfCSP complements a cold sensitive E. coli mutant

The pNII-CSPA/pNII-CfCSP construction was in BX04 cells confirmed by PCR, and the protein products were also detected by SDS-PAGE (data not shown). After incubation at 37°C for 12 h, the JM109 cells with pNII and BX04 cells with pNII, pNII-CSPA or pNII-CfCSP all formed large numbers of clones with no difference (Fig. 8A). However, when incubated at 17°C for 120 h, BX04 cells containing the vector pNII showed growth defect and failed to form colonies, while BX04 cells containing pNII-CSPA or pNII-CfCSP showed vigorous growth (Fig. 8B). Though the BX04 cells containing pNII-CfCSP or pNII-CSPA also grew evidently at 17°C, the clone was much smaller as compared with the positive clone JM109 with pNII (Fig. 8B).

Discussion

CSDPs are nucleic acid-binding proteins involved in various cellular processes, such as cold adaptation [7]. In invertebrates, CSDPs have been identified from several species [35–39], but their functions are still not well understood. In the present paper, a CSDP was identified from scallop C. farreri. The ORF of CfCSP cDNA was of 927 bp encoding a polypeptide of 308 amino acids. The deduced amino acids of CfCSP consisted of an N-terminal CSD and a C-terminal domain with four RGG repeats. Compared to the bacteria and plant CSDPs, the CSD of the CfCSP displayed higher identity to that of the human YB-1. The presence of CfCSP in CSDP suggested that CfCSP might function as a RNA chaperone [14]. In addition to the CSD, there were four additional C-terminal RGG motifs in CfCSP [39]. RGG motif is usually defined as closely spaced Arg-Gly-Gly repeats separated by other amino acids [40], but the four RGG repeats in CfCSP are not closely distributed. The RGG motif is thought to combine with other RNA-binding motifs and functions by increasing the overall RNA affinity of proteins containing additional RNA-binding domain [40]. The difference between the RGG motif from CfCSP and other invertebrate CSDPs might induce some functional changes. These structural information indicated that CfCSP was a new member of invertebrate CSDPs, and it might play a similar role as RNA chaperone at low temperature.

The mRNA expression pattern of CfCSP in different tissue and its temporal expression in adductor and hemocytes after acute cold shock treatment were examined in the present study to explore its possible function. The CfCSP mRNA was constitutively expressed in all the examined scallop tissues, where higher expression level was found in adductor and gonad. However, to our knowledge, the immune or stress related genes are most actively expressed in the immune or stress related genes are most actively expressed in hepatopancreas or gill, the gene expression pattern of CfCSP gene might have some particular indication for physiological functions. For example, the high expression of cystatin gene in muscle [48,49] suggested that these proteins might function in muscle remodeling. Because the adductor is the main part related to the movement of scallop, the high expression of CfCSP in adductor suggested that it might play similar functions in muscle cells remodeling. After the treatment of an acute cold shock, the CfCSP mRNA expression in both adductor and hemocytes increased significantly \((P<0.05)\) within 26 h, indicating that CfCSP probably involved in the acute cold shock response of Zhihong scallops. Considering that the CSDPs from other species always acted as nucleic acid binding proteins [4–6], the present results indicated that CfCSP could be effectively induced by acute cold shock and...
Figure 2. Multiple alignment of the amino acid sequences for CSD containing proteins. Cold shock domains are boxed, consensus RNA binding domains (RNP1 and RNP2) and RGG repeats are shaded in gray and consensus DNA binding sites are shaded in dark. Perfectly matched residues, conserved residues, and less conserved residues are indicated by an asterisk (*), a colon (:), and a period (.), respectively. Accession numbers of the CSD proteins are: CfCSP (JN869460), LsYB (AAT97092), ApY1 (NP_001191560), E. coli CSPA (P15277), human YB-1 (P67808) and XlYB3 (CAA42778).

doi:10.1371/journal.pone.0032012.g002
Figure 3. Phylogenetic tree of CSD containing protein sequences from diverse organisms. The tree is constructed by the neighbor-joining (NJ) algorithm and the scale bar corresponds to 0.2 estimated amino-acid substitution per site. Accession numbers of the CSD containing proteins are: E. coli CSP (P15277), E. coli CSP (P36995), Lactococcus lactis CSPB (CAA76695), Bacillus subtilis CSPB (P32081), C. elegans LIN-28 (AAC47476), Human YB-1 (I39382), CfCSP (JN869460), Xenopus laevis YB3 (CAA42778), Lymnaea stagnalis YB (AAC42622), Aplysia californica Y1 (NP_001191560), WCSP1 (AA42622), AtCSP3 (NM_127341), AtCSP2 (NP_001191560), NsGRP2 (CAA42622).

doi:10.1371/journal.pone.0032012.g003

Figure 4. CfCSP mRNA expression in different tissues of C. farreri detected by RT-PCR. CfCSP transcripts level in adductor, gonad, kidney, mantle, hepatopancreas and gill were normalized to that of hemocytes. Vertical bars represented the mean ± S.E. (N = 5), and bars with different letters were significantly different (P < 0.05).

doi:10.1371/journal.pone.0032012.g004
might play an important role in cold adaptation as an RNA chaperone.

The bacterial CSPs could bind RNA and destabilize the secondary structures of RNA to prevent the premature transcription termination under low temperatures [7,14,50]. Though many eukaryotic CSDPs displayed DNA/RNA binding activities [4,19,23,25], only RBM3 in Human Cells had been reported in response to cold shock [51]. In CfCSP, a conserved CSD was identified, suggesting that it would exhibit RNA chaperon activity under low temperature. The in vitro nucleic acid-binding assay revealed that rCfCSP protein could bind ssDNA and in vitro transcribed mRNA, but it could not bind dsDNA. This was different from the Y-box proteins which could bind both ssDNA and dsDNA [4]. It was fascinating that CSPA of *E. coli* acquired dsDNA binding activity by replacing QNDGYK with the Y-box consensus sequence KKNPRKYL, suggesting that the length and basicity of the loop region is a determinant of dsDNA binding.

**Figure 5. Temporal expression of the CfCSP mRNA after acute cold shock treatment.** The mRNA level of CfCSP relative to β-actin in adductor (A) and hemocytes (B) were measured by RT-PCR. Vertical bars represent the mean ± S.E. (N = 5). (*: P<0.05). doi:10.1371/journal.pone.0032012.g005

**Figure 6. SDS–PAGE analysis of rCfCSP.** Lane M: protein molecular standard (kDa); lane 1: negative control for rCfCSP (without induction); lane 2: induced rCfCSP; lane 3: purified rCfCSP. doi:10.1371/journal.pone.0032012.g006
activity [52]. However, the sequence KKNNPRKYLR in Y-box of human was replaced by TKNNPRKYLR in CfCSP of scallop, and the substitution of the first amino acid from K to T was suspected to lead to the loss of the dsDNA binding activity. Furthermore, the difference of the C-terminal domain with other CSDPs might also result in the loss of the dsDNA binding activity. There were four RGG motifs in C-terminal domain of CfCSP, which was different from other CSDPs in vertebrates and plants. A mutant WCSP1 lacking C-terminal zinc fingers lost dsDNA binding activity, indicating that the C-terminal domain was important for the dsDNA binding.

Similar to the bacterial CSPs, some eukaryotic CSDPs are also suggested to be involved in the process of cold adaptation. In Arabidopsis, GRP7 complemented successfully the cold sensitivity of E. coli BX04 mutant and the over-expression of GRP7 increased the cold tolerances of plants [33]. However, GRP4 failed to complement the growth of E. coli BX04 mutant during cold stress and over-expression of GRP4 did not increase the cold tolerance of plants [34]. It seemed that there was a common structure-function relationship between bacterial and eukaryotic CSDPs during the process of cold adaptation [33]. In the present study, vigorous growth was observed for BX04 containing pLNIII-CfCSP at 17°C, indicating that CfCSP can partially complement the function of E. coli CSPs for cold adaptation. Together with the up-regulated expression of CfCSP after acute cold shock, it is reasonable to propose that the abundant transcripts of CfCSP might increase the cold tolerances of scallops. Because of the differences in the process of transcription termination between prokaryotic and eukaryotic systems, it seems that CfCSP can’t function as a transcription antiterminator to regulate the cold adaptation in scallops. Meanwhile, wheat WCSP1 has been reported to be involved in the translation process for its ER localization [34], it is suspected that eukaryotic CSDPs might regulate the cold adaptation at translation level.

In conclusion, the present work provided novel information on the roles of mollusk CSDPs in response to acute cold shock. The presence of a conserved CSD and four RGG motifs, the ubiquitous tissue expression and the cold-dependent induction pattern suggested that CfCSP was a primitive member of the invertebrate CSDPs. The nonspecific nuclear acids binding activity and the efficient complementation of the BX04 cold sensitivity suggested that CfCSP offers RNA chaperon activities during the cold adaptation process.

Acknowledgments

We are grateful to Dr. M. Inouye and Dr. S. Phadtare for BX04 mutant cell and pLNIII vector and to Dr. R. Imai for his helpful advice.

Author Contributions

Conceived and designed the experiments: LS CY LW QJ. Performed the experiments: CY VS XS JW. Analyzed the data: CY VS XS QJ. Contributed reagents/materials/analysis tools: CY LW HZ LS. Wrote the paper: CY LW LS.

Figure 7. Analysis of nuclear acids binding activity of CfCSP by gel shift assay. The purified rCfCSP proteins were incubated with ssDNA (A), M13mp8 dsDNA RFI DNA (dsDNA) (B) and the in vitro transcribed luciferase mRNA (C) to analyze the effect of CfCSP on the formation of nucleotide-protein complexes. A range of CfCSP fusion proteins from 0 to 300 pmol were used for analysis. Arrow shows nucleotide-protein complex.

doii:10.1371/journal.pone.0032012.g007

Figure 8. Complementation of cold sensitive growth of BX04 (ΔcspAΔcspBΔcspEΔcspG) with CfCSP and E. coli CSPs. Overnight cultures of JM109/pLNIII, BX04/pLNIII, BX04/CfCSP and BX04/CSPA were streaked on LB-ampiciline plates and incubated at 37°C for 12 h (A) or 17°C for 120 h (B).

doii:10.1371/journal.pone.0032012.g008
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